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Integrated Science Assessment for Ozone and Related Photochemical Oxidants

(Third External Review Draft)

National Center for Environmental Assessment-RTP Division Office of Research and Development U.S. Environmental Protection Agency Research Triangle Park, NC

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TABLE OF CONTENTS

DI	SCLAIMER		ii
Т	ABLE OF CONTENTS		iii
0	ZONE PROJECT TEAM		xxvii
A	JTHORS, CONTRIBUTORS,	AND REVIEWERS	xxx
CI	LEAN AIR SCIENTIFIC ADVI	SORY COMMITTEE OZONE NAAQS REVIEW PANEL	xxxvi
A	CRONYMS AND ABBREVIA	LIONS	_xxxviii
PI	REAMBLE		liv
	Figure I	Illustration of the key steps in the process of the review of National	lv
	Figure II	Illustration of processes for literature search and study selection used for development of ISAs.	
	Figure III	Characterization of the general process of ISA development	
	Table I	Aspects to aid in judging causality	lxviii
	Table II	Weight of evidence for causal determination	lxxi
	References		lxxvii
LE	EGISLATIVE AND HISTORIC	AL BACKGROUND	_ lxxviii
	Table III	Summary of primary and secondary NAAQS promulgated for ozone during the period 1971-2008	lxxx
	References		lxxxiv
1	EXECUTIVE SUMMARY		1-1
	Introduction and Purpose		1-1
	Integration of Ozone Health Effects		
	Table 1-1	Summary of ozone causal determinations by exposure duration and health outcome	
	Respiratory Effects		
	Mortality Effects		1-7
	Populations Potentially at Inc	reased Risk	1-7
		and Ecosystems	
	Table 1-2	Summary of ozone causal determination for welfare effects	1-8
	Visible Foliar Injury	-	
	Growth, Productivity, Carbon	Storage and Agriculture	
	Below Ground Processes		1-10
		Relationships	
		Climate Change and UV-B Effects	
		e Change	
	UV-B Effects		1-12

			Table 1-3	Summary of ozone causal determination for climate change and UV-B effects.	1-13
	Cond	lusion _			1-13
2	INT	EGRA	TIVE SUMMARY_		2-1
	2.1	ISA D	evelopment and Scope	·	2-2
	2.2	Atmos	pheric Chemistry and	Ambient Concentrations	2-5
		2.2.1	Physical and Chemic	al Processes	2-5
			Figure 2-1	Schematic overview of photochemical processes influencing stratospheric and tropospheric ozone	2-6
		2.2.2	Atmospheric Modelin	g of Background Ozone Concentrations	2-7
			Table 2-1	Comparison of seasonal mean MDA8 ozone concentrations simulated by the GEOS-Chem and CAMx base case models for 2006, with measurements at CASTNET sites.	2-8
			Figure 2-2	Mean daily average 8-hour ozone concentrations in surface air, for spring and summer 2006.	2-9
			Monitoring		2-11
				ons	
	2.3				
	2.4			n	
	2.5			Effects	
				evious Ozone AQCDs	
		2.5.2		Determinations	2-19
			Table 2-2	Summary of evidence from epidemiologic, controlled human exposure, and animal toxicological studies on the health effects associated with short- and long-term exposure to ozone.	2-23
		2.5.3	Integrated Synthesis	of Evidence for Health Effects	2-26
		2.5.4	Policy Relevant Cons	iderations	2-32
				Potentially at Increased Risk	
				etrics in Epidemiologic Studies	
				e in Epidemiologic Studies	
				entration-Response Relationship	
	2.6	Integr		etation and Ecosystems	
	2.0	-	Visible Foliar Injury		2-38
		2.0.1	Figure 2-3	An illustrative diagram of the major pathway through which ozone enters leaves and the major endpoints that ozone may affect in plants and ecosystems.	2 00
			Table 2-3	Summary of ozone causal determinations for vegetation and ecosystem	2-39
				effects.	2-40
		2.6.2	Growth, Productivity,	Carbon Storage and Agriculture	2-41
			2.6.2.1 Natural Ecos	systems	2-41
			0	Crops	2-43
		2.6.3			2-44
		2.6.4		SS85	2-44
		2.6.5 2.6.6		tion	2-45 2-46
		2.0.0		ndices	2-40 2-46
			2.6.6.2 Exposure-Re	esponse	2-48
	2.7	The R	•	one in Climate Change and UV-B Effects	
		2.7.1		as a Greenhouse Gas	2-49
			Figure 2-4	Schematic illustrating the effects of tropospheric ozone on climate; including the relationship between precursor emissions, tropospheric ozone abundance, radiative forcing, climate response, and climate impacts.	2-50
		2.7.2	Tropospheric Ozone	and UV-B related effects	2-50
		2.7.3		and UV-B Related Effects	
	2.8			nations for Health Effects and Welfare Effects	

				Table 2-4	Summary of ozone causal determinations by exposure duration and health outcome.	
				Table 2-5	Summary of ozone causal determination for welfare effects.	_
				Table 2-6	Summary of ozone causal determination for climate and UV-B effects.	
I	Refe	rences				
3	ATN	IOSP	HERIC	CHEMIST	RY AND AMBIENT CONCENTRATIONS	
;	3.1	Introd	uction			
:	3.2				SSES	
				Figure 3-1	Schematic overview of photochemical processes influencing stratospheric and tropospheric ozone.	
		3.2.1	Sources	s of Precursor	s Involved in Ozone Formation	
				Figure 3-2		
		3.2.2	Gas Ph	ase Reactions	Leading to Ozone Formation	
		3.2.3	Multipha	ase Processes	8	
		3.2.4			emical Precursor Relationships	
				Figure 3-3	Measured concentrations of ozone and NO _z .	
:	3.3	Atmos	spheric M	odeling		
				Figure 3-4	Sample Community Multi-scale Air Quality (CMAQ) modeling domains.	
				Figure 3-5	Main components of a comprehensive atmospheric chemistry modeling system, such as the U.S. EPA's Community Multi-scale Air Quality (CMAQ) modeling system.	
		3.3.1	Clobal	Scale CTMs		
		0.0.1	Giobal	Figure 3-6	Comparison of global chemical-transport model (CTM) predictions of maximum daily 8-h avg ozone concentrations and multi-model mean with monthly averaged CASTNET observations in the Intermountain West and Southeast regions of the U.S.	
:	3.4	Backo	round Oz	zone Concentr	ations	
		200.03	,	Figure 3-7	Schematic overview of contributions to North American background concentrations of ozone.	
		3.4.1	Contrib	utions from Na	atural Sources	
			3.4.1.1	Contribution	s from the Stratosphere	
					s from Other Natural Sources	
		3.4.2			hthropogenic Emissions	
		0.4.2	Contrib	Figure 3-8	Time series of daily maximum 8-h avg (MDA8) ozone concentrations (ppm) measured at Trinidad Head, CA, from April 18, 2002 through December 31, 2009	
		3.4.3	Estimat	ing Backgrour	d Concentrations	
				0 0	OS-Chem Model Estimates of Background Concentrations	
					Mean MDA8 ozone concentrations in surface air for spring and summer 2006 calculated by GEOS-Chem for the base case (Base), U.S. background (USB), and NA background (NAB).	
				Figure 3-10	Spring and summer mean Canadian and Mexican (CM) contributions to MDA8 ozone determined as the difference between the U.S. background and NA background.	
				Figure 3-11	MDA8 ozone concentrations for spring (March-May) and summer (June- August) 2006 simulated by GEOS-Chem vs. measured by the ensemble of CASTNET sites in the Intermountain West, Northeast, Great Lakes, and Southeast.	
				Figure 3-12	Frequency distributions of MDA8 ozone concentrations in March- August 2006 for the ensemble of low-altitude (<1,500 meters) and high-altitude CASTNET sites (>1,500 meters) in the U.S.	
			3.4.3.2	Using Other	Models to Estimate Background Concentrations	
				Figure 3-13	Mean MDA8 ozone concentrations in surface air during spring and summer 2006 (top) calculated by GEOS-Chem/CAMx for the base case	
					(Base, top) and NA background (NAB, bottom).	

			Figure 3-14	Monthly average MDA8 ozone concentrations observed (Obs) and predicted for the base case and NA background (NAB) by GEOS-Chem (GC) and GEOS-Chem/CAMx (CX) at CASTNET sites above 1,500 meters elevation (upper panel) and CASTNET sites below 1,500 meters elevation (lower panel).	3-53
			Figure 3-15	Annual 4th highest MDA8 ozone predicted by GEOS-Chem $(0.5^{\circ} \times 0.667^{\circ})$ for the base case (Base) with corresponding U.S. background (USB) and NA background (NAB) MDA8 ozone for the same days in 2006.	3-58
			Figure 3-16	Annual 4th highest MDA8 ozone predicted by CAMx for the base case (Base) and corresponding NA background (NAB) MDA8 ozone for the same days in 2006.	3-59
			Table 3-1	Comparison of seasonal mean MDA8 ozone concentrations simulated by the GEOS-Chem and CAMx base case models for 2006, with measurements at CASTNET sites.	3-61
			Table 3-2	Comparison of annual 4th-highest MDA8 ozone concentrations measured at CASTNET sites in 2006 with MDA8 ozone concentrations simulated by the GEOS-Chem and CAMx base case models.	3-63
3.5	Monito	orina			3-64
	3.5.1	0		echniques	
			-		3-67
			Table 3-3	Summary of ozone monitors meeting 40 CFR Part 58, Appendix A Precision and Bias Goals	3-68
			0	Box plots of precision data by year (2005-2009) for all ozone monitors reporting single-point QC check data to AQS.	3-69
			-	Box plots of percent-difference data by year (2005-2009) for all ozone monitors reporting single-point QC check data to AQS.	3-69
		3.5.2.1		m Co-located UV Ozone Monitors in Missouri	3-70
			•	Box plots of RPD data by year for the co-located ozone monitors at two sites in Missouri from 2006-2009.	3-70
			Figure 3-20		0.74
	252	Dorform	onno Crosifia	point QC check data to AQS from 2005-2009	
	3.5.3	Fellom	Table 3-4	ations Performance specifications for ozone based in 40 CFR Part 53	
	3.5.4	Monitor			3-72 3-72
	3.5.5			hniques	
	0.0.0			Ozone Monitors	
				hemiluminescence Monitors	
				Sampling Devices and Sensors	
				Detical Absorption Spectrometry	
				note Sensing	
	3.5.6			ork Design	
				g Requirements	
				U.S. ozone sites reporting data to AQS in 2010.	3-79
				U.S. Rural NCore, CASTNET and NPS POMS ozone sites in 2010.	
		3.5.6.2	Probe/Inlet S	Siting Requirements	3-82
3.6	Ambie			• •	3-82
	3.6.1	Measur	ement Units.	Metrics, and Averaging Times	3-83
				Distribution in nation-wide year-round site-level correlations between daily ozone metrics including 24-h avg, 1-h daily max and 8-h daily max using	
		0	.,	AQS data, 2007-2009.	3-85
	3.6.2	•	•	117 - 1 - 199	3-86
		3.6.2.1		Required ozone monitoring time periods (ozone season) identified by	3-86
			Table 2 F	monitoring site Summary of ozone data sets originating from AQS	3-86
			Table 3-5 Figure 3-25	Location of the 457 ozone monitors meeting the year-round data set completeness criterion for all 3 years between 2007 and 2009.	3-87 3-88
			Figure 3-26	Location of the 1,064 ozone monitors meeting the warm-season data set completeness criteria for all 3 years between 2007 and 2009.	3-88

	Table 2.0	Nations data distributions of some some strations (with) from the	
	Table 3-6	Nationwide distributions of ozone concentrations (ppb) from the vear-round data set.	3-90
	Table 3-7	Nationwide distributions of ozone concentrations (ppb) from the warm- season data set.	3-91
	Table 3-8	Seasonally stratified distributions of 8-h daily max ozone concentrations (ppb) from the year-round data set (2007-2009).	3-93
	Figure 3-27	Highest monitor (by county) 3-year avg (2007-2009) of the 8-h daily max ozone concentration based on the year-round data set (top map) with seasonal stratification (bottom 4 maps).	3-94
	Figure 3-28	Highest monitor (by county) 3-year avg (2007-2009) of the 8-h daily max ozone concentration based on the warm-season data set (top map) with annual stratification (bottom 3 maps).	3-95
	Table 3-9	Focus cities used in this and previous assessments	3-98
	Table 3-10	City-specific distributions of 8-h daily max ozone concentrations (ppb) from the warm-season data set (2007-2009).	3-100
	Figure 3-29	Map of the Atlanta CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-101
	Figure 3-30	Map of the Boston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-101
	Figure 3-31	Map of the Los Angeles CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-102
	Figure 3-32	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Atlanta CSA	3-104
	Figure 3-33	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Boston CSA.	3-104
	Figure 3-34	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Los Angeles CSA.	3-105
	Figure 3-35	Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.	3-107
	Figure 3-36	Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.	3-108
	Figure 3-37	Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.	3-109
	Figure 3-38	Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.	3-110
	Figure 3-39	Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.	3-111
	Figure 3-40	Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.	
	Figure 3-41	Terrain map showing the location of two nearby AQS ozone monitoring sites (red dots) along the western edge of the Los Angeles CSA. Site AL is near shore, 3 meters above sea level, while Site AK is in an agricultural valley surrounded by mountains, 262 meters above sea level.	3-112
	Figure 3-42	Terrain map showing the location of four AQS ozone monitoring sites (red dots) located in or near the city limits in the center of the Boston CSA. Site characteristics range from Site A near downtown at 6 meters above sea level to Site D in a forested area on Blue Hill at 192 meters above sea level.	3-115
3.6.2.2	Rural-Focus	ed Variability and Ground-Level Vertical Gradients	
-	Table 3-11	Rural focus areas.	
	Figure 3-43	Rural focus area site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the rural focus areas.	3-119
	Figure 3-44	Terrain map showing the location of five AQS ozone monitoring sites (green/black stars) in Great Smoky Mountain National Park, NC-TN	
		(SMNP)	3-121

			Figure 3-45	Pair-wise monitor correlations (left) and coefficients of divergence (COD, right) expressed as a histogram (top), contour matrix (middle) and scatter plot vs. distance between monitors (bottom) for Great Smoky Mountain National Park, NC-TN (SMNP).	3-122
			Figure 3-46	Rocky Mountain National Park, CO (black/green star) and the Denver CSA (red dots) along with ozone monitoring sites used in the Brodin et al. (2010) study (blue circles)	3-123
			Figure 3-47	Terrain map showing the location of two AQS ozone monitoring sites (black/green stars) in Sequoia National Park, CA	3-124
	3.6.3	Tempor	al Variability _		
				ends	3-125
			Figure 3-48	National 8-h daily max ozone trend and distribution across 870 U.S. ozone monitors, 1998-2010 (annual 4th highest 8-h daily max ozone concentrations in ppm).	3-126
			Figure 3-49	National 1-h daily max ozone trend and distribution across 875 U.S. ozone monitors, 1998-2010 (annual second highest 1-h daily max ozone concentrations in ppm).	3-127
			Figure 3-50	Trend in 8-h daily max ozone by region, 1998-2010 (annual 4th highest 8- h daily max ozone concentrations in ppm).	3-128
			Figure 3-51	Trend in 1-h daily max ozone by region, 1998-2010 (annual second highest 1-h daily max ozone concentrations in ppm).	3-129
			Figure 3-52	Individual monitor 8-h daily max ozone design values displayed A) for the 2008-2010 period and B) as the change since the 2001-2003 period.	3-130
			Figure 3-53	Individual monitor 1-h daily max ozone design values displayed A) for the 2008-2010 period and B) as the change since the 2001-2003 period.	3-131
		3.6.3.2		tions	3-134
			Figure 3-54	Diel patterns in 1-h avg ozone for Atlanta, Boston and Los Angeles between 2007 and 2009.	3-135
			Figure 3-55	Diel patterns in 1-h avg ozone for six rural focus areas between 2007 and 2009.	3-138
	3.6.4	Associa	tions with Co-	pollutants	3-139
				Distribution of Pearson correlation coefficients for comparison of 8-h daily max ozone from the year-round data set with co-located 24-h avg CO, SO ₂ , NO ₂ , PM ₁₀ and PM _{2.5} from AQS, 2007-2009	3-141
			Figure 3-57	Distribution of Pearson correlation coefficients for comparison of 8-h daily max ozone from the warm-season (May-Sept) data set with co-located 24-h avg CO, SO ₂ , NO ₂ , PM ₁₀ and PM _{2.5} from AQS, 2007-2009.	3-142
3.7	Chant	or Summ	any	24 havg 00, 002, h02, h who and h with hom Add, 2007 2000.	3-143
5.7	3.7.1			al Processes	
	3.7.1	-			3-143
	3.7.3			ations	3-145
	3.7.4		ing		3-148
	3.7.5	Ambien	t Concentratio	ons	3-149
3.8	Supple	emental li	nformation on	Ozone Model Predictions	3-151
			Figure 3-58	Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Northeast with GEOS-Chem predictions for the base case and for the North American background case during March-August, 2006.	3-152
			Figure 3-59		
				American background case during March-August, 2006.	3-153
			Figure 3-60	Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Upper Midwest with GEOS-Chem predictions for the base case and for the North American background case during March-August, 2006.	3-153
			Figure 3-61	Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Intermountain West with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.	3-154
					0-104

	Figure 3-62	Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Intermountain West with GEOS-Chem predictions for the base case and	
	Figure 3-63	the North American background case during March-August, 2006 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the West with GEOS-Chem predictions for the base case and the North American	3-154
	Figure 3-64	background case during March-August, 2006 Comparison of time series of measurements of daily maximum 8-hour	3-155
		average ozone concentrations at three CASTNET sites and the Trinidad Head site in California with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.	3-155
	Figure 3-65	Comparison of daily maximum 8-h average ozone predicted using GEOS- Chem at $0.5^{\circ} \times 0.667^{\circ}$ (and $2^{\circ} \times 2.5^{\circ}$ resolution; left figure only) with measurements at Mount Bachelor, OR (left); and at Trinidad Head, CA (right) from March to August 2006.	3-156
	Figure 3-66	Comparison of monthly mean (± 1 standard deviation) ozone calculated GEOS-Chem (in red) with ozonesondes (in black) at Trinidad Head, CA	
	F igure 0.07	(top) and Boulder, CO (bottom) during April and August 2006.	3-156
	Figure 3-67	A deep stratospheric ozone intrusion over California on May 28-29, 2010.	3-157
	Figure 3-68 Figure 3-69	A deep stratospheric ozone intrusion over California on June 7-12, 2010 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Northeast and predictions from GEOS-Chem at ~50 x 50 km resolution (green) and CAMx at 12 x 12 km resolution (blue) for May-August 2006.	3-158 3-159
	Figure 3-70	Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Southeast and predictions from GEOS-Chem at ~50 x 50 km resolution (green) and CAMx at 12 x 12 km resolution (blue) for May-August 2006.	3-160
	Figure 3-71	Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Central U.S. and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.	3-161
	Figure 3-72	Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Northern Rockies and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.	3-162
	Figure 3-73	Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Southern Rockies and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.	3-163
	Figure 3-74	Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the West and predictions from GEOS-Chem at ~50 x 50 km resolution (green) and CAMx at 12 x 12 km resolution (blue) for May-August 2006.	3-164
	Figure 3-75	Daily maximum 8-hour average (MDA8) ozone in surface air at Gothic, CO for March through August 2006.	3-165
3.9	Supplemental Figures of Obs	erved Ambient Ozone Concentrations	_ 3-165
		for the Urban Focus Cities	3-165
		Map of the Atlanta CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-166
	Figure 3-77	gravity centers, urban areas, and major roadways.	3-166
	Figure 3-78	population gravity centers, urban areas, and major roadways.	3-167
	Figure 3-79	Map of the Boston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-167
	Figure 3-80 Figure 3-81	Map of the Chicago CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways Map of the Dallas CSA including ozone monitor locations, population	3-168
	0	gravity centers, urban areas, and major roadways.	3-168
	Figure 3-82	gravity centers, urban areas, and major roadways.	3-169
	Figure 3-83	Map of the Detroit CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-169

	Figure 3-84	Map of the Houston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-170
	Figure 3-85	Map of the Los Angeles CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-170
	Figure 3-86	Map of the Minneapolis CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-171
	Figure 3-87	Map of the New York CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-171
	Figure 3-88	Map of the Philadelphia CSA including ozone monitor locations,	
	Figure 3-89	population gravity centers, urban areas, and major roadways Map of the Phoenix CBSA including ozone monitor locations, population	3-172
	Eiguro 2.00	gravity centers, urban areas, and major roadways.	3-172
	Figure 3-90	Map of the Pittsburgh CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-173
	Figure 3-91	Map of the Salt Lake City CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-173
	Figure 3-92	Map of the San Antonio CBSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-174
	Figure 3-93	Map of the San Francisco CSA including ozone monitor locations,	3-174
	Figure 3-94	population gravity centers, urban areas, and major roadways Map of the Seattle CSA including ozone monitor locations, population	3-174
	Ũ	gravity centers, urban areas, and major roadways.	3-175
	Figure 3-95	Map of the St. Louis CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.	3-175
3.9.2	Ozone Concentration	Box Plots for the Urban Focus Cities	3-176
	Figure 3-96	Site information, statistics and box plots for 8-h daily max ozone from	
		AQS monitors meeting the warm-season data set inclusion criteria within the Atlanta CSA.	3-176
	Figure 3-97	Site information, statistics and box plots for 8-h daily max ozone from	
		AQS monitors meeting the warm-season data set inclusion criteria within the Baltimore CSA.	2 177
			3-177
	Figure 3-98	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Birmingham CSA	3-177
	Figure 3-99	Site information, statistics and box plots for 8-h daily max ozone from	0
		AQS monitors meeting the warm-season data set inclusion criteria within the Boston CSA.	3-178
	Figure 3-100	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within	
		the Chicago CSA	3-178
	Figure 3-101	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Dallas CSA.	3-179
	Figure 3-102	Site information, statistics and box plots for 8-h daily max ozone from	0170
		AQS monitors meeting the warm-season data set inclusion criteria within the Denver CSA.	3-179
	Figure 3-103	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within	
	F i 0.404	the Detroit CSA.	3-180
	Figure 3-104	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Houston CSA.	3-180
	Figure 3-105	Site information, statistics and box plots for 8-h daily max ozone from	
		AQS monitors meeting the warm-season data set inclusion criteria within the Los Angeles CSA	3-181
	Figure 3-106	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within	
		the Minneapolis CSA	3-182
	Figure 3-107	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the New York CSA.	3-182
	Figure 3-108	Site information, statistics and box plots for 8-h daily max ozone from	02102
		AQS monitors meeting the warm-season data set inclusion criteria within the Philadelphia CSA.	3-183

	Figure 3-109	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Phoenix CBSA.	3-183
	Figure 3-110	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Pittsburgh CSA.	3-184
	Figure 3-111	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Salt Lake City CSA	3-184
	Figure 3-112	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the San Antonio CBSA.	3-185
	Figure 3-113	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the San Francisco CSA	3-185
	Figure 3-114	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Seattle CSA	3-186
	Figure 3-115	Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the St. Louis CSA.	3-186
3.9.3	Ozone Concentration	Relationships for the Urban Focus Cities	3-187
	Figure 3-116	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.	3-187
	Figure 3-117	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Baltimore CSA.	3-188
	Figure 3-118	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Birmingham CSA.	3-189
	Figure 3-119	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.	3-190
	Figure 3-120	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Chicago CSA.	3-191
	Figure 3-121	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Dallas CSA.	3-192
	Figure 3-122	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Denver CSA.	3-193
	Figure 3-123	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Detroit CSA.	3-194
	Figure 3-124	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Houston CSA.	3-195
	Figure 3-125	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.	3-196
	Figure 3-126	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Minneapolis CSA.	3-197
	Figure 3-127	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the New York CSA.	3-198
	Figure 3-128	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Philadelphia CSA.	3-199
	Figure 3-129	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Phoenix CBSA.	0 100

Figure 3-130	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Pittsburgh CSA.	3-201
Figure 3-131	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Salt Lake City CSA.	3-202
Figure 3-132	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Antonio CBSA.	3-203
Figure 3-133	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Francisco CSA.	3-204
Figure 3-134	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Seattle CSA.	3-205
Figure 3-135	Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the St. Louis CSA.	3-206
Figure 3-136	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.	3-207
Figure 3-137	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Baltimore CSA.	3-208
Figure 3-138	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Birmingham CSA.	3-209
Figure 3-139	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA	3-210
Figure 3-140	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Chicago CSA.	3-211
Figure 3-141	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Dallas CSA	3-212
Figure 3-142	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Denver CSA.	3-213
Figure 3-143	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Detroit CSA.	3-214
Figure 3-144	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Houston CSA.	3-215
Figure 3-145	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.	3-216
Figure 3-146	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Minneapolis CSA.	3-217
Figure 3-147	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the New York CSA.	3-218
Figure 3-148	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Philadelphia CSA.	3-219
Figure 3-149	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Phoenix CBSA.	3-220
Figure 3-150	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Pittsburgh CSA.	3-221

				Figure 3-151	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance	
				F : 0.450	between monitors (bottom) for the Salt Lake City CSA.	3-222
				Figure 3-152	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Antonio CBSA	3-223
				Figure 3-153	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Francisco CSA.	3-224
				Figure 3-154	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Seattle CSA.	3-225
				Figure 3-155	Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the St. Louis CSA.	3-226
		3.9.4	Hourly	variations in O	zone for the Urban Focus Cities	3-227
			,		Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).	
				Figure 3-157	Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).	3-228
				Figure 3-158	Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).	0 220
				Figure 3-159	Diel patterns in 1-h avg ozone for select CSAs/CBSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the	2 220
				Figure 3-160	weekday/weekend comparison (right half)	3-230
	Defe				weekday/weekend comparison (right half).	
	Refer	rences _				_ 3-232
4	EXP	POSUF	RETO	AMBIENT (DZONE	4-1
•						
	4.1					
	4.2					
	4.3					
				-	echniques	
		4.3.2	Indoor-0		entration Relationships	4-5
		400	Danaan	Table 4-1	Relationships between indoor and outdoor ozone concentration	
		4.3.3	Persona	Figure 4-1	ncentration Relationships Variation in hourly personal and ambient concentrations of ozone and PM _{2.5} in various microenvironments during daytime hours	4-9 4-10
				Table 4-2	Correlations between personal and ambient ozone concentration.	4-13
				Table 4-3	Ratios of personal to ambient ozone concentration.	
		4.3.4	Co-exp	osure to Other	Pollutants and Environmental Stressors	
			4.3.4.1	Personal Exp	posure to Ozone and Copollutants	4-17
					xposure to Ozone and Copollutants	
				Figure 4-2	Correlations between 1-week concentrations of ozone and copollutants measured near roadways.	4-20
			4.3.4.3	Indoor Expos	sure to Ozone and Copollutants	4-20
	4.4	Expos	ure-Relat	ed Metrics		4-21
		4.4.1	Activity	Patterns		4-21
			2	Table 4-4	Mean fraction of time spent in outdoor locations by various age groups in the NHAPS study	4-22
				Table 4-5	Mean ventilation rates (L/min) at different activity levels for different age	

groups.

_4-23

			Figure 4-3	Distribution of time that NHAPS respondents spent in ten microenvironments based on smoothed 1-min diary data.	4-24
	4.4.2	Ozone /	Averting Beha	avior	4-25
	4.4.3	Populat	ion Proximity	to Fixed-Site Ozone Monitors	4-27
		·	Figure 4-4	Map of the Atlanta CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.	4-29
			Figure 4-5	Map of the Boston CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.	4-30
			Figure 4-6	Map of the Los Angeles CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.	4-3 ²
			Table 4-6	Fraction of the 2009 population living within a specified distance of an ozone monitor in selected U.S. cities.	4-34
1.5	Expos	ure Mode	eling		
.0	Expoo	are meae	Table 4-7		4-3
	4.5.1	Concen		e Modeling	
	4.5.2			ange Rate Modeling	4-3
	4.5.3			ased Models	4-4
.6				c Studies	
.0			-		
	4.6.1			Exposure	
	4.6.2	•		/	
				ability	
					4-4
	4.6.3				
				Exposure	
				Exposure	
	4.6.4			ants and Ozone Reaction Products	
	4.6.5	Averting			4-5
			Figure 4-7	Adjusted asthma hospital admissions by age on lagged ozone by alert status, ages 5-19	4-52
			Figure 4-8	Adjusted asthma hospital admissions by age on lagged ozone by alert status, ages 20-64	
				Mathada in Enidamialagia Studiag	
		•		Methods in Epidemiologic Studies	
.7		•			
	Summ	ary and (Conclusions_		4-54
Refe	Summ rences _	ary and (Conclusions_	· ·	4-54 4-58
DOS	Summ rences _ SIMET	RY AN	Conclusions_	DF ACTION	4-54 4-58 _5-^
Refe	Summ rences _ SIMET	ary and (Conclusions_	· ·	4-54 4-58 _5-^
Refe	Summ rences _ SIMET	RY AN	Conclusions_	DF ACTION	4-54 4-58 5- ^
Refe DOS 5.1	Summ rences SIMET Introdu	RY AN	D MODE C	DF ACTION	4-5 4-5 5- 5-
efe 00 \$	Summ rences SIMET Introdu	RY AN	D MODE C	DF ACTION	4-5 4-5 5- 5- 5-
efer)O \$.1	Summ rences SIMET Introdu Huma	RY AN	D MODE C	DF ACTION	4-54 4-54 5-* 5-* 5-* 5-*
efer)O \$.1	Summ rences SIMET Introdu Huma	RY AN	D MODE C	DF ACTION	4-54 4-54 5 5 5 5 5 5
efer 0 0 \$	Summ rences SIMET Introdu Huma	RY AN uction	D MODE C Figure 5-1	DF ACTION Schematic of the ozone exposure and response pathway. Ozone transport follows a path from exposure concentration, to inhaled dose, to net dose, to the local tissue dose. Chapter 5 discusses the concepts of dose and modes of action that result in the health effects discussed in Chapters 6 and 7	4-54 4-54 5 5 5 5 5 5
efe 0 0\$.1	Summ rences SIMET Introdu Huma 5.2.1	RY AN uction	D MODE C Figure 5-1	DF ACTION	4-54 4-54 5 5 5 5 5 5
efer 0 0 \$	Summ rences SIMET Introdu Huma 5.2.1	n and Ani Introduc	Conclusions D MODE C Figure 5-1 mal Ozone D ttion Figure 5-2 Figure 5-3 Jptake Table 5-1	DF ACTION	4-5. 4-5. 5- 5- 5- 5- 5- 5- 5- 5- 5- 5-
efe 0 0\$.1	Summ rences SIMET Introdu Huma 5.2.1	n and Ani Introduction Ozone I	Conclusions D MODE C Figure 5-1 mal Ozone D tion Figure 5-2 Figure 5-3 Jptake Table 5-1 Gas Transpo	DF ACTION Schematic of the ozone exposure and response pathway. Ozone transport follows a path from exposure concentration, to inhaled dose, to net dose, to the local tissue dose. Chapter 5 discusses the concepts of dose and modes of action that result in the health effects discussed in Chapters 6 and 7	4-5. 4-5. 5 5 5 5 5 5 5 5 5 5 5 5 5
efer)O \$.1	Summ rences SIMET Introdu Huma 5.2.1	n and Ani Introduction Ozone I 5.2.2.1 5.2.2.2	Conclusions D MODE C Figure 5-1 mal Ozone D ttion Figure 5-2 Figure 5-3 Jptake Table 5-1 Gas Transpo Target Sites	DF ACTION Schematic of the ozone exposure and response pathway. Ozone transport follows a path from exposure concentration, to inhaled dose, to net dose, to the local tissue dose. Chapter 5 discusses the concepts of dose and modes of action that result in the health effects discussed in Chapters 6 and 7	4-54 4-54 5 5 5 5 5 5 5 5 5 5 5
efe 00 \$	Summ rences SIMET Introdu Huma 5.2.1	n and Ani Introduction 0zone to 5.2.2.1 5.2.2.2 5.2.2.3	Conclusions D MODE C Figure 5-1 Figure 5-2 Figure 5-3 Jptake Table 5-1 Gas Transpo Target Sites Upper Resp	DF ACTION	4-54 4-58 5-1 5-2 5-2 5-2 5-3 5-4 5-8 5-4 5-10
Refe	Summ rences SIMET Introdu Huma 5.2.1	n and Ani Introduction 0zone to 5.2.2.1 5.2.2.2 5.2.2.3	Conclusions D MODE C Figure 5-1 Figure 5-2 Figure 5-3 Jptake Table 5-1 Gas Transpo Target Sites Upper Resp	DF ACTION Schematic of the ozone exposure and response pathway. Ozone transport follows a path from exposure concentration, to inhaled dose, to net dose, to the local tissue dose. Chapter 5 discusses the concepts of dose and modes of action that result in the health effects discussed in Chapters 6 and 7	4-54 4-58 5-1 5-2 5-2 5-2 5-3 5-5 5-1 5-10 5-10 5-12

5

		5.2.2.6	Interindividua	al Variability in Dose	5-15
		5.2.2.7	Physical Act	ivity	5-16
			Table 5-2	General adult human inhalation rates by activity levels.	5-17
			Figure 5-5	Modeled effect of exercise on tissue dose of the LRT.	5-18
		5.2.2.8	Summary _		5-19
	5.2.3			Reaction Products	
			Figure 5-6	Schematic overview of ozone interaction with PUFA in ELF and lung cells.	5-21
		5.2.3.1	Summary		5-28
			Figure 5-7	Details of the ozone interaction with the airway ELF to form secondary oxidation products.	5-29
5.3	Possib	le Pathw	ays/Modes of	Action	5-29
	5.3.1	Introduc	tion		5-29
	5.3.2			Reflexes	5-30
	5.3.3			ion	5-34
	5.3.4	Alteratio	on of Epithelia	Barrier Function	5-40
	5.3.5	Sensitiz	ation of Brond	chial Smooth Muscle	5-42
	5.3.6	Modifica	ation of Innate	Adaptive Immune System Responses	5-45
	5.3.7	Airways	Remodeling_		5-49
	5.3.8	-		n and Oxidative/Nitrosative Stress	
	5.3.9			erial Oxygen Transfer	5-52
	5.3.10	Summa	ry		5-52
			Figure 5-8	The modes of action/possible pathways underlying the health effects resulting from inhalation exposure to ozone.	5-53
5.4	Interine	dividual \	/ariability in R	esponse	5-56
	5.4.1	Dosime	tric Considera	tions	5-57
	5.4.2			ations	
				nment Interactions	5-58
		5.4.2.2	Pre-existing	Diseases and Conditions	5-62
				tatus	
			-		
				of Responses	
				es with Particulate Matter	
		5.4.2.7			5-74
			Figure 5-9	Some factors, illustrated in yellow, that likely contribute to the interindividual variability in responses resulting from inhalation of ozone.	5-75
5.5	Specie	s Homol	ogy and Inters	species Sensitivity	5-75
	5.5.1	Interspe	cies Dosimet	ry	5-76
			-	Humans and animals are similar in the regional pattern of ozone tissue dose distribution.	5-78
			-	Oxygen-18 incorporation into different fractions of BALF from humans and rats exposed to 0.4 and 2.0 ppm ¹⁸ O ₃ .	5-80
	5.5.2			y of Response	
	5.5.3	Summa	ry		5-84
5.6	Chapte	er Summ	ary		5-84
Refe	rences _				5-86
INT	EGRA [.]	TED HI		FECTS OF SHORT-TERM OZONE EXPOSURE	6-1
6.1					
6.2					
	6.2.1				
		6.2.1.1		uman Exposure	6-3
			Table 6-1	Activity levels used in controlled exposures of healthy young adults to ozone.	6-7
			Figure 6-1	Cross-study comparison of mean ozone-induced FEV ₁ decrements following 6.6 hours of exposure to ozone.	6-8

6

		Figure 6-2	Frequency distributions of FEV ₁ decrements observed by Schelegle et al. (2009) in young healthy adults (16 F, 15 M) following 6.6-hour exposures to ozone or filtered air.	6-17
	6.2.1.2	Epidemiolog	У	6-27
		Table 6-2	Mean and upper percentile ozone concentrations in epidemiologic studies of lung function in populations with increased outdoor exposures.	6-29
		Figure 6-3	Changes in FEV ₁ (mL) or PEF (mL/sec) in association with ambient ozone concentrations among children attending summer camp	6-32
		Table 6-3	Additional characteristics and quantitative data for studies represented in Figure 6-3.	6-33
		Figure 6-4	Percent change in FEV ₁ in association with ambient ozone concentrations among adults exercising outdoors.	6-35
		Table 6-4	Additional characteristics and quantitative data for studies represented in Figure 6-4 plus results from studies in children exercising outdoors.	6-36
		Figure 6-5	Percent change in FEV ₁ or FEV ₁ /FVC in association with ambient ozone concentrations among outdoor workers.	6-39
		Table 6-5	Additional characteristics and quantitative data for studies represented in Figure 6-5.	6-40
		Table 6-6	Associations between ambient ozone concentration and FEV ₁ decrements in different ranges of ambient ozone concentrations.	6-41
		Table 6-7	Mean and upper percentile concentrations of ozone in epidemiologic studies of lung function in children with asthma.	6-42
		Figure 6-6	Percent change in FEV ₁ in association with ambient ozone concentrations among children with asthma.	6-44
		Table 6-8	Characteristics and quantitative data for studies represented in Figure 6- 6, of FEV ₁ or FVC in children with asthma.	6-45
		Figure 6-7	Percent change in PEF or FEF _{25-75%} in association with ambient ozone concentrations among children with asthma.	6-46
		Table 6-9	Characteristics and quantitative data for studies represented in Figure 6- 7, of PEF or FEF _{25-75%} in children with asthma.	6-47
		Table 6-10	Mean and upper percentile concentrations of ozone in epidemiologic studies of lung function in adults with respiratory disease.	6-54
		Table 6-11	Mean and upper percentile concentrations of ozone in epidemiologic studies of lung function in populations not restricted to individuals with asthma.	6-56
		Figure 6-8	Percent change in FEV ₁ or FVC in association with ambient ozone concentrations in studies of children in the general population.	6-58
		Table 6-12	Characteristics and quantitative data for studies represented in Figure 6- 8, of lung function in children.	6-59
		Table 6-13	Associations between ambient ozone concentration and lung function in studies of adults.	6-61
		Figure 6-9	Comparison of ozone-associated changes in lung function in single- and co-pollutant models	6-65
		Table 6-14	Additional characteristics and quantitative data for studies represented in Figure 6-9.	6-66
	6.2.1.3	Toxicology: I	_ung Function	6-70
6.2.2			iveness	6-71
	6.2.2.1	Controlled H	uman Exposures	6-71
			Airway Hyperresponsiveness	
6.2.3	Pulmon	ary Inflammat	ion, Injury and Oxidative Stress	6-74
	6.2.3.1	Controlled H	uman Exposures	6-75
	6.2.3.2	Epidemiolog	У	6-81
		Table 6-15	Mean and upper percentile ozone concentrations in studies of biological markers of pulmonary inflammation and oxidative stress.	6-83
		Figure 6-10	Percent change in exhaled nitric oxide (eNO) in association with ambient ozone concentrations in populations with and without asthma.	6-84
		Table 6-16	Additional characteristics and quantitative data for studies represented in Figure 6-10.	6-85
		Table 6-17	Associations between short-term ambient ozone exposure and biological markers of pulmonary inflammation and oxidative stress.	6-86
	6.2.3.3	Toxicology: I	nflammation and Injury	6-95
		Table 6-18	Morphometric observations in non-human primates after acute ozone exposure.	6-98

6.2.4	Respira	tory Symptom	s and Medication Use	6-100			
	6.2.4.1	Children with) Asthma	_6-102			
		Table 6-19	Mean and upper percentile ozone concentrations in epidemiologic studies of respiratory symptoms, medication use, and activity levels in children with asthma.	6-103			
		Figure 6-11	Associations between ambient ozone concentrations and respiratory symptoms in children with asthma.	6-105			
		Table 6-20	Additional characteristics and quantitative data for studies presented in Figure 6-11.				
		Figure 6-12		6-110			
		Table 6-21	Additional characteristics and quantitative data for studies presented in Figure 6-12.	6-111			
	6.2.4.2	Adults with R	Respiratory Disease	6-113			
		Table 6-22	Mean and upper percentile ozone concentrations in epidemiologic studies of respiratory symptoms and medication use in adults with respiratory disease	 6-114			
	6.2.4.3	Populations	not Restricted to Individuals with Asthma	6-115			
		Table 6-23	Mean and upper percentile ozone concentrations in epidemiologic studies of respiratory symptoms in populations not restricted to individuals with asthma.	6-116			
		Figure 6-13	Associations between ambient ozone concentrations and respiratory symptoms in children in the general population.				
		Table 6-24	Additional characteristics and quantitative data for studies represented inFigure 6-13.				
	6.2.4.4	Confounding	in Epidemiologic Studies of Respiratory Symptoms and Medication Use	6-119			
		Table 6-25	Associations between ambient ozone concentrations and respiratory symptoms in single- and co-pollutant models.	_6-121			
	6.2.4.5 Summary of Epidemiologic Studies of Respiratory Symptoms and Asthma Medication Use						
6.2.5							
			Clearance				
			hiolar Transport Mechanism				
	6.2.5.3	5.3 Alveolar Macrophages					
	6.2.5.4	Infection and	Adaptive Immunity	6-126			
	6.2.5.5	Summary of	Lung Host Defenses				
6.2.6	Allergic and Asthma-Related Responses						
6.2.7			Emergency Department Visits, and Physicians Visits				
	6.2.7.1	Summary of	Findings from 2006 Ozone AQCD	6-132			
		Table 6-26	Mean and upper percentile concentrations of respiratory-related hospital admission and emergency department (ED) visit studies evaluated	6-134			
	6.2.7.2	•	nission Studies	6-136			
		Figure 6-14	Percent increase in respiratory hospital admissions from natural spline models with 8 df/yr for a 40 ppb increase in 1-h max ozone	6-140			
		Table 6-27	concentrations for each location of the APHENA study Corresponding effect estimates for Figure 6-14	0-140 6-141			
		Figure 6-15	Estimated relative risks (RRs) of asthma hospital admissions for 8-h max ozone concentrations at lag 0-1 allowing for possible nonlinear	6-148			
	6273	Emergency	relationships using natural splines	0-148 6-149			
	0.2.7.0		Risk ratio for respiratory ED visits and different ozone exposure metrics in Atlanta from 1993-2004.	6-151			
		Figure 6-17		_6-154			
	6.2.7.4	Outpatient a	nd Physician Visit Studies	6-156			
		Summary	· · · · · · · · · · · · · · · · · · ·	6-157			
			Percent increase in respiratory-related hospital admission and ED visits in studies that presented all-year and/or seasonal results.	_6-158			
		Table 6-28	Corresponding Effect Estimates for Figure 6-18.				
		Figure 6-19	Percent increase in respiratory-related hospital admissions and ED visits for studies that presented single and copollutant model results.	6-161			
		Table 6-29	Corresponding effect estimates for Figure 6-19.				

	6.2.8	Respira	tory Mortality_		6-163		
	6.2.9	9 Summary and Causal Determination					
6.3	Cardio						
	6.3.1	Controlled Human Exposure					
	6.3.2	Epidemi	iology		6-173		
			Arrhythmia		6-173		
			Table 6-30	Characterization of ozone concentrations (in ppb) from studies of arrhythmias.	6-174		
		6.3.2.2	Heart Rate/H	leart Rate Variability	6-176		
			Table 6-31	Characterization of ozone concentrations (in ppb) from studies of heart rate variability.	6-177		
		6.3.2.3	Stroke		6-180		
			Figure 6-20	Odds ratio (95% confidence interval) for ischemic stroke by quintiles of ozone exposure	6-182		
		6.3.2.4	Biomarkers_		6-182		
			Table 6-32	Characterization of ozone concentrations (in ppb) from studies of biomarkers.	6-183		
		6.3.2.5	Myocardial I	nfarction (MI)	6-188		
		6.3.2.6	Blood Press	ure	6-189		
			Table 6-33	Characterization of ozone concentrations (in ppb) from studies of blood pressure.	6-190		
		6.3.2.7	Hospital Adr	nissions and Emergency Department Visits	6-191		
			Table 6-34	Characterization of ozone concentrations (in ppb) from studies of hospital admissions and ED visits.	6-192		
			Figure 6-21		6-198		
			Table 6-35	Effect estimate (95% CI) per increment ppb increase in ozone for overall cardiovascular ED visits or hospital admissions in studies presented inFigure 6-21.	6-199		
			Figure 6-22	.	6-201		
			Table 6-36	Effect estimate (95% CI) per increment ppb increase in ozone for congestive heart failure ED visits or hospital admissions for studies in Figure 6-22.	6-202		
			Figure 6-23	Effect estimate (95% CI) per increment ppb increase in ozone for ischemic heart disease, coronary heart disease, myocardial infarction, and angina pectoris ED visits or hospital admissions.	6-203		
			Table 6-37	Effect estimate (95% CI) per increment ppb increase in ozone for ischemic heart disease, coronary heart disease, myocardial infarction, and angina pectoris Evisits or hospital admissions for studies presented			
			Figure 6-24		6-204		
			Table 6-38	ED visits or hospital admissions Effect estimate (95% CI) per increment ppb increase in ozone for stroke	6-205		
			Figure 6-25	ED visits or hospital admissions for studies presented in Figure 6-24 Effect estimate (95% CI) per increment ppb increase in ozone for arrhythmia and dysrhythmia ED visits or hospital admissions	6-206		
			Table 6-39	Effect estimate (95% CI) per increment ppb increase in ozone for arrhythmia and dysrhythmia ED visits or hospital admissions for studies	6-207		
			a "	presented in Figure 6-25	6-208		
				lar Mortality	6-208		
			-	Epidemiologic Studies	6-209		
	6.3.3	IOXICOIC	0,	scular Effects			
	004	0	Table 6-40				
	6.3.4			I Determination			
6.4	Centra	I Nervou					
			Table 6-41	Central nervous system and behavioral effects of short-term ozone	0.004		
	644	Nourse	doorine F#	exposure in rats	6-224		
	6.4.1	Neuroer		ts	6-225		
6 5	6.4.2			I Determination	6-226		
6.5	Effects	s on Othe	r Organ Syste	ems	6-227		

5.5.2 Effects on Cutaneous and Ocular Tissues	
Aortality	6-
Aortality	
i.6.1 Summary of Findings from 2006 Ozone AQCD	-66
Figure 6-26 Summary of mortality risk estimates for short-term ozone exposure and all-cause (nonaccidental) mortality from all-year and summer season analyses.	0-
Table 6-42 Corresponding effect estimates for Figure 6-26.	0 6
Table 6-43 Range of mean and upper percentile ozone concentrations in previous and recent multicity studies.	0 6-
6.6.2.1 Confounding	6-
Table 6-44 Correlations between PM and ozone by season and region.	6-
Figure 6-27 Scatter plots of ozone mortality risk estimates with versus without adjustment for PM_{10} in NMMAPS cities	6
Figure 6-28 Community-specific ozone-mortality risk estimates for nonaccidental mortality per 10 ppb increase in same-day 24-h average summertime ozone concentrations in single-pollutant models and copollutant models with sulfate.	6
Figure 6-29 Percent increase in all-cause (nonaccidental) and cause-specific mortality from natural spline models with 8 df/yr from the APHENA study for single- and copollutant models.	6
Table 6-45 Corresponding effect estimates for Figure 6-29.	0 6
Table 6-46 Sensitivity of ozone risk estimates per 10 μg/m³ increase in 24-h average ozone concentrations at lag 0-1 to alternative methods for adjustment of seasonal trend, for all-cause mortality using Berkey MLE and TLNISE Hierarchical Models.	0
6.6.2.2 Effect Modification	0 6
Table 6-47 Additional percent change in ozone-related mortality for individual-level characteristics.	6
Figure 6-30 Ozone mortality risk estimates and community-specific characteristics, U.S., 1987-2000.	6
Table 6-48 Percent change in all-cause mortality, for all ages, associated with a 40ppb increase in 1-h max ozone concentrations at Lag 0–1 at the 25th and 75th percentile of the center-specific distribution of selected effect modifiers.	6
Table 6-49 Percentage increase in daily mortality for a 10 ppb increase in 24-h average ozone concentrations during the previous week by geographic region in the U.S., 1987-2000.	0
Figure 6-31 Community-specific Bayesian ozone-mortality risk estimates in 98 U.S. communities.	6
Figure 6-32 Map of spatially dependent ozone-mortality coefficients for 8-h max ozone concentrations using summer data.	6
6.6.2.3 Interaction	6
6.6.2.4 Evaluation of the Ozone-Mortality C-R Relationship and Related Issues Table 6-50 Estimated effect of a 10 ppb increase in 8-h max ozone concentrations on mortality during the summer months for single-day and distributed lag models.	6 6
Figure 6-33 Estimated combined smooth distributed lag for 48 U.S. cities during the summer months.	0
Table 6-51 Estimated percent increase in cause-specific mortality (and 95% CIs) for a 10-µg/m ³ increase in maximum 8-hour ozone during June-August.	6
Figure 6-34 Estimated combined smooth distributed lag in 21 European cities during the summer (June-August) months.	6
Table 6-52 Percent excess all-cause mortality per 10 ppb increase in daily 8-h max ozone on the same day, by season, month, and age groups. Firmer 0.25 Firmer 0.25	6
Figure 6-35 Estimated combined C-R curve for nonaccidental mortality and 24-hour average ozone concentrations at lag 0-1 using the nonlinear (spline) model.	6
6.6.2.5 Associations of Cause-Specific Mortality and Short-term Ozone Exposure	6
Figure 6-36 Percent increase in cause-specific mortality.	6
Table 6-53 Corresponding effect estimates for Figure 6-36.	6
6.3 Summary and Causal Determination	6

Refe	erences _			Summary of causal determinations for short-term exposures to ozone.	
INT	EGRA	TED HI	EALTH EF	FECTS OF LONG-TERM OZONE EXPOSURE	
7.1	Introdu	uction			
7.2					
				Asthma	
			Figure 7-1	Interaction of heme-oxygenase genetic variants and O ₃ level on the Hazard Ratio (HR) of new-onset asthma in the 12 Children's Health Study communities.	
		7.2.1.2	Prevalence	of Asthma and Asthma Symptoms	
			Figure 7-2	Ozone modifies the effect of TNF GG genotype on bronchitic symptoms among children with asthma in the CHS.	
	7.2.2	Asthma	Hospital Adn	nissions and ED Visits	
			Figure 7-3	Ozone-asthma concentration-response relationship using the mean concentration during the entire follow-up period for first asthma hospital admission.	
	7.2.3	Pulmon	arv Structure	and Function	
	1.2.0		,	Structure and Function: Evidence from Epidemiology Studies	
				Structure and Function: Evidence from Toxicological Studies and Nonhuman I	
			Table 7-1	Respiratory effects in nonhuman primates and rodents resulting from long-term ozone exposure	
	7.2.4	Pulmon	ary Inflamma	tion, Injury, and Oxidative Stress	
	7.2.5	Allergic	Responses_		
	7.2.6	Host De	efense		
	7.2.7	Respira	tory Mortality		
	7.2.8	Summa	ry and Causa	I Determination	
			Table 7-2	Summary of selected key new studies examining annual ozone exposure and respiratory health effects	
			Table 7-3	Studies providing evidence concerning potential confounding by PM for available endpoints.	
7.3	Cardic	ovascular	Effects		
	7.3.1				
				ılar Epidemiology	
		7.3.1.2	Cardiovascu	ılar Toxicology	
			Table 7-3	Characterization of Study Details for Section 7.3.1.2.	
	7.3.2			ality	
	7.3.3		,	I Determination	
7.4	Repro	ductive a	nd Developm	ental Effects	
	7.4.1	Effects	on Sperm		
	7.4.2			ion	
	7.4.3	Birth We	eight		
			Figure 7-4	Birthweight deficit by decile of 24-h avg ozone concentration averaged over the entire pregnancy compared with the decile group with the lowest ozone exposure.	
			Table 7-4	Brief Summary of Epidemiologic Studies of Birth Weight.	
	7.4.4	Preterm	Birth	,	
			Table 7-5	Brief summary of epidemiologic studies of PTB	
	7.4.5	Fetal G	rowth		
	-		Table 7-6	Brief summary of epidemiologic studies of fetal growth.	_
	7.4.6	Postnat			
	7.4.7		efects		
			Table 7-7	Brief summary of epidemiologic studies of birth defects	
	7.4.8	Develop	omental Resp	iratory Effects	
	7.4.9			al Nervous System Effects	

		7.4.9.1 Laterality		7-7 [.]			
		7.4.9.2 Brain Morpho	blogy and Neurochemical Changes	7-7			
		7.4.9.3 Neurobehavi	oral Outcomes	7-72			
		7.4.9.4 Sleep Aberra	tions after Developmental Ozone Exposure	7-7			
	7.4.10	Early Life Mortality		7-7			
			ty, Less than 1 Year				
		7.4.10.3 Neonatal Mo	rtality, Less than 1 Month	7-7			
		7.4.10.4 Postneonata	Mortality, 1 Month to 1 Year	7-7			
		7.4.10.5 Sudden Infar	nt Death Syndrome	7-7			
		Table 7-8	Brief summary of infant mortality studies.	7-7			
		Table 7-9	Summary of key reproductive and developmental toxicological studies.	7-7			
	7.4.11	Summary and Causal	Determination	7-7			
7.5	Centra	I Nervous System Effe	cts	7-8			
	7.5.1	Effects on the Brain a	nd Behavior	7-8			
		Table 7-10	Central nervous system effects of long-term ozone exposure in rats.	7-8			
	7.5.2	Summary and Causal	Determination	7-8			
7.6	Carcin	ogenic and Genotoxic I	Potential of Ozone	7-8			
	7.6.1	7.6.1 Introduction					
	7.6.2		e and Mortality				
	7.6.3	-	·				
	7.6.4		Determination				
7.7	Mortal	ty		7-9			
		Figure 7-5	Adjusted ozone-mortality relative risk estimates (95% CI) by time period of analysis per subject-weighted mean ozone concentration in the Cancer Prevention Study II by the American Cancer Society.	7-9			
		Table 7-11	Relative risk (and 95% CI) of death attributable to a 10-ppb change in the				
			ambient ozone concentration.	7-9			
	7.7.1	Summary and Causal	Determination	7-9			
7.8	Overa	I Summary		7-9			
		Table 7-12	Summary of causal determinations for long-term exposures to ozone.	7-9			
Refe	rences		· · · · · · · · · · · · · · · · · · ·	7-98			

8 POPULATIONS POTENTIALLY AT INCREASED RISK FOR OZONE-RELATED HEALTH EFFECTS _____

EFF	ECTS	·		
		Table 8-1	Classification of Evidence for Potential At-Risk Factors.	
8.1	Genet	ic Factors		
		Table 8-2	Summaries of results from epidemiologic and controlled human exposures studies of modification by genetic variants.	
8.2	Preex	isting Disease/Conditi	ons	
		Table 8-3	Prevalence of respiratory diseases, cardiovascular diseases, and diabetes among adults by age and region in the U.S.	
	8.2.1	Influenza/Infections		8
	8.2.2	Asthma		8
		Table 8-4	Prevalence of asthma by age in the U.S	8
	8.2.3	Chronic Obstructive	Pulmonary Disease (COPD)	8
	8.2.4	Cardiovascular Dise	ease (CVD)	8
	8.2.5	Diabetes		8
	8.2.6			
8.3	Socio	demographic Factors		8
	8.3.1	Lifestage		8
			ts	
	8.3.2			
	8.3.3	Socioeconomic Stat	us	
	8.3.4	Race/Ethnicity		
8.4	Behav	vioral and Other Facto	rs	8

	8.4.1	Diet		8-31	
				8-32	
	8.4.3 Smoking				
				8-35	
				8-35	
8.5				8-36	
		Table 8-5	Summary of evidence for potential increased risk of ozone-related health effects.	8-37	
References					

9 ENVIRONMENTAL EFFECTS: OZONE EFFECTS ON VEGETATION AND ECOSYSTEMS 9-1

9.1	Introdu		9-1
		Figure 9-1 An illustrative diagram of the major pathway through which ozone enters plants and the major endpoints that ozone may affect in plants and ecosystems.	9-3
9.2	Experi	imental Exposure Methodologies	9-3
	9.2.1	Introduction	9-3
	9.2.2	"Indoor," Controlled Environment, and Greenhouse Chambers	9-4
	9.2.3	Field Chambers	
	9.2.4	Plume and FACE-Type Systems	9-6
	9.2.5	Ambient Gradients	9-8
	9.2.6	Comparative Studies	9-9
9.3	Mecha	anisms Governing Vegetation Response to Ozone	9-10
	9.3.1	Introduction	9-10
	9.3.2	Ozone Uptake into the Leaf	9-12
		9.3.2.1 Changes in Stomatal Function	
		Figure 9-2 The microarchitecture of a dicot leaf	
		Figure 9-3 Possible reactions of ozone within water.	
		Figure 9-4 The Crigee mechanism of ozone attack of a double bond.	
	9.3.3		
		9.3.3.1 Ozone Detection and Signal Transduction	9-17
			9-18
		Figure 9-5 Composite diagram of major themes in the temporal evolution of the genetic response to ozone stress.	9-23
		9.3.3.3 Role of Phytohormones in Plant Response to Ozone	9-24
		Figure 9-6 The oxidative cell death cycle.	9-26
	9.3.4	Detoxification	9-27
		9.3.4.1 Overview of Ozone-induced Defense Mechanisms	
		9.3.4.2 Role of Antioxidants in Plant Defense Responses	
	9.3.5		
		9.3.5.1 Light and Dark Reactions of Photosynthesis	
			9-33
		9.3.5.3 Secondary Metabolism	9-34
		Summary	9-36
9.4		e of Effects on Vegetation and Ecosystems	
	9.4.1	Introduction	
		9.4.1.1 Ecosystem Scale, Function, and Structure	
	0.4.0	9.4.1.2 Ecosystem Services	9-40
	9.4.2	Visible Foliar Injury and Biomonitoring	9-41 9-43
		9.4.2.1 Biomonitoring	9-43 9-45
	9.4.3	9.4.2.2 Summary Growth, Productivity and Carbon Storage in Natural Ecosystems	
	9.4.5	9.4.3.1 Plant Growth and Biomass Allocation	9-40 9-46
		9.4.3.2 Summary	
		9.4.3.3 Reproduction	9-50 9-50
		Table 9-1 Ozone effects on plant reproductive processes	
		9.4.3.4 Ecosystem Productivity and Carbon Sequestration	
			002

			Table 9-2	Comparison of models used to simulate the ecological consequences of ozone exposure	9-54
		9.4.3.5	Summary		9-58
			Table 9-3	Modeled effects of ozone on primary production, C exchange, and C sequestration.	9-60
	9.4.4	Crop Yi	eld and Qualit	y in Agricultural Systems	
	•••••			,	
					9-67
		0.4.4.0	Table 9-4	Summary of recent studies of ozone effects on crops (exclusive of growth	0 0/
				and yield)	
			Table 9-5	Modeled effects of ozone on crop yield loss at regional and global scales	
	9.4.5	Water C	Cycling		9-72
			Figure 9-7	, , , , , , , , , , , , , , , , , , , ,	
	9.4.6	Below-0		SSES	9-76
			Figure 9-8	Conceptual diagram showing where ozone alters C, water and nutrient flow in a tree-soil system, including transfer between biotic and abiotic components below ground that influence soil physical and chemical properties.	9-77
		9.4.6.1	Litter Carbor	h Chemistry, Litter Nutrient and Their Ecosystem Budgets	9-78
			Table 9-6	The effect of elevated ozone on leaf/litter nutrient concentrations.	9-79
		9.4.6.2	Decomposer	Metabolism and Litter Decomposition	
				tion and Carbon Formation	
			Table 9-7	The temporal variation of ecosystem responses to ozone exposure at Aspen FACE site	
		9464	Nutrient Cvc		
				rganic Carbon and Biogenic Trace Gases Emission	
	9.4.7			ion	9-86
	9.4.7				
				nd Agricultural Land	
	0.4.0		Summary		
	9.4.8		-	unctional and Growth Response	
			Genetics		
				al Biological Factors	
			•	tors	
		9.4.8.4		with other Pollutants	9-93
			Table 9-8	Response of plants to the interactive effects of elevated ozone exposure and nitrogen enrichment.	9-94
	9.4.9	Insects	and Other Wil	dlife	9-97
		9.4.9.1	Insects		9-97
		9.4.9.2	Wildlife		9-100
		9.4.9.3	Indirect Effect	cts on Wildlife	9-101
		9.4.9.4	Summary		9-104
9.5	Effects			oosure Indices and Dose Modeling	
	9.5.1			<u> </u>	
	9.5.2			Ire Indices Available in the Literature	
	0.0.2	Descrip	Figure 9-9	Diagrammatic representation of several exposure indices illustrating how they weight concentration and accumulate exposure.	
	9.5.3	Importo	nt Component	ts of Exposure Indices	
	9.0.0			centration	9-112
		9.0.3.1		Trends in May to September: 12-hour SUM06, Peak 1-hour ozone	9-112
			Figure 9-10	concentration and number of daily exceedances of 95 ppb for the Crestline site in 1963 to 1999; in relation to trends in mean daily maximum temperature for Crestline and daily reactive organic gases	

			Figure 9-11	The number of hourly average concentrations between 50 and 89 ppb for the period 1980-2000 for the Crestline, San Bernardino County, CA, monitoria ait	9-115
		0 5 2 2	Diumaland	monitoring site	_9-115 9-115
		9.5.3.2		Seasonal Exposure Diurnal (a) conductance through boundary layer and stomata (gbs), (b)	_9-115
			rigule 9-12	ozone concentration, and leaf-level stomatal ozone flux (Fstol) in control plots from mid-June through August in (c) 2004 and (d) 2005 in the Aspen	0.440
			Figure 9-13	FACE experiment Maximum 3-month, 12-h W126 plotted against maximum 6-month, 12-h W126	_9-118 _9-120
	9.5.4	Ozone I	Intake/Dose M	12-h W126 Modeling for Vegetation	
	9.5.5				9-123
9.6			-	onse Relationships	_
0.0	9.6.1		tion		9-125
	9.6.2			Id Loss and Tree Seedling Biomass Loss in the 1996 and 2006 Ozone AQCDs	
	0.0.2	20111141		Quantiles of predicted relative yield loss for 34 NCLAN crop experiments.	
				Quantiles of predicted relative yield loss for 4 crop species in NCLAN experiments.	9-131
			Figure 9-16	Quantiles of predicted relative biomass loss for 49 tree species in NHEERL/WED experiments.	9-132
			Figure 9-17	Quantiles of predicted relative biomass loss for 4 tree species in NHEERL/WED experiments.	
			Table 9-9	Ozone exposures at which 10 and 20% yield loss is predicted for 50 and 75% of crop species.	_9-134
			Table 9-10	Ozone exposures at which 10 and 20% yield loss is predicted for 50 and 75% of crop species (Droughted versus Watered conditions).	_9-134
			Table 9-11	Ozone exposures at which 10 and 20% biomass loss is predicted for 50 and 75% of tree species	_9-135
	9.6.3			d 2006 Ozone AQCD Models and Methodology Using the 90 day 12-h W126 a	
			_		_9-135
		9.6.3.1	Table 9-12	of NCLAN-Based Prediction and SoyFACE Data Comparison between change in yield observed in the SoyFACE experiment between elevated and ambient ozone, and change predicted at the same values of ozone by the median composite function for	_9-137
			Table 9-13	NCLAN Comparison between yield observed in the SoyFACE experiment and yield predicted at the same values of ozone by the median composite function for NCLAN.	_9-138 _9-138
			Figure 9-18	Comparison of yield observed in SoyFACE experiment in a given year with yield predicted by the median composite function based on NCLAN.	9-139
			Figure 9-19	Comparison of composite functions for the quartiles of 7 curves for 7 genotypes of soybean grown in the SoyFACE experiment, and for the quartiles of 11 curves for 5 genotypes of soybean grown in the NCLAN project.	9-140
		9.6.3.2	Comparison Data	of NHEERL/WED-Based Prediction of Tree Biomass Response and Aspen FA	_
			Table 9-14	Comparison between change in above-ground biomass elevated and ambient ozone in Aspen FACE experiment in 6 year, and change predicted at the same values of ozone by the median composite function for NHEERL/WED.	9-142
			Table 9-15	Comparison between above-ground biomass observed in Aspen FACE experiment in 6 year and biomass predicted by the median composite function based on NHEERL/WED	9-142
			Figure 9-20	Comparison between above-ground biomass observed in Aspen FACE experiment in 6 year and biomass predicted by the median composite function based on NHEERL/WED	_9-143
		9.6.3.3	Exposure-Re	esponse in a Gradient Study	_9-144
				Above-ground biomass for one genotype of cottonwood grown in seven locations for one season in 3 years.	9-145
		9.6.3.4	Meta-analyse	es of growth and yield studies	_9-146
			Table 9-16		_9-146
		9.6.3.5	Additional ex	posure-response data	_9-148
	9.6.4	Summa	ry		_9-148

	Table 9-17	Summary of studies of effects of ozone exposure on growth and yield of agricultural crops	9-149
	Table 9-18	Summary of studies of effects of ozone exposure on growth of natural vegetation.	9-153
9.7	Summary and Conclusions_		9-155
	Table 9-19	Summary of ozone causal determinations for vegetation and ecosystem effects	9-156
Refe	rences		9-157

10 THE ROLE OF TROPOSPHERIC OZONE IN CLIMATE CHANGE AND UV-B EFFECTS__10-1

10.1	Introduction10				
10.2	Physics of the Earth's Radiation Budget			_ 10-1	
		Figure 10-1	Diagram of the factors that determine human exposure to ultraviolet		
			radiation.		
10.3	Effects of Tropospheric Ozone on Climate				
				10-4	
	10.3.2 Climate Change Evidence and the Influence of Tropospheric Ozone				
	10.3.2.1 Climate Change in the Recent Past				
	10.3.2.2 Projections of Future Climate Change				
	10.3.2.3 Metrics of Potential Climate Change				
	10		c Ozone as a Greenhouse Gas		
		-	Schematic illustrating the effects of tropospheric ozone on climate.	10-9	
		Figure 10-3	Global average radiative forcing (RF) estimates and uncertainty ranges in 2005 for anthropogenic CO ₂ , CH ₄ , ozone and other important agents and mechanisms.	_10-10	
	10.3.3 Fa	actors that Influence	e the Effect of Tropospheric Ozone on Climate		
			e Concentration of Tropospheric Ozone		
			f Surface Albedo on Ozone Radiative Forcing		
			f Vertical Distribution on Ozone Radiative Forcing		
	10.3.3.4 Feedback Factors that Alter the Climate Response to Changes in Ozone Radiative				
	10.3.3.5 Indirect Effects of Tropospheric Ozone on the Carbon Cycle				
	10.3.4 Competing Effects of Ozone Precursors on Climate10				
			Components of radiative forcing for emissions of principal gases, aerosols, aerosol precursors, and other changes.		
	10.3.5 C	alculating Radiative	Forcing and Climate Response to Past Trends in Tropospheric Ozone	_10-19	
		Figure 10-5	Ensemble average 1900-2000 radiative forcing and surface temperature trends (°C per century) in response to tropospheric ozone changes.	_10-20	
	10.3.6 C	alculating Radiative	Forcing and Climate Response to Future Trends in Tropospheric Ozone	_10-20	
	10.3.6.1 Emissions of Anthropogenic Ozone Precursors Across the 21st Century			_10-21	
	10.3.6.2 Impact of 21st Century Trends in Emissions on Tropospheric Ozone		_10-22		
		Table 10-1	2000-2030 changes in anthropogenic emissions, and CH_4 and tropospheric ozone concentrations, and the associated tropospheric		
			ozone forcing for three scenarios.		
		•	, , , , , , , , , , , , , , , , , , , ,	_10-23	
	10		rcing and Climate Response from 21st Century Trends in Tropospheric Ozone	_10-24	
		-	Global mean radiative forcing estimates calculated by a set of models for the 2000-2100 change in tropospheric ozone		
10.4	UV-B Related Effects and Tropospheric Ozone				
	10.4.1 Background				
	10.4.2 Human Exposure and Susceptibility to Ultraviolet Radiation				
	10.4.3 Human Health Effects due to UV-B Radiation			_10-27	
	10.4.4 Ecosystem and Materials Damage Effects Due to UV-B Radiation			_10-28	
	10.4.5 UV-B Related Effects Associated with Changes in Tropospheric Ozone Concentrations			_10-30	
10.5	Summary and Causal Determinations			10-31	
				_10-31	
	10.5.2 Summary of UV-B Related Effects on Human Health, Ecosystems, and Materials Relating to Changes			es in _10-33	
	10.5.3 Si	ummary of Ozone C	Causal Determinations	_10-33	

Table 10-2	Summary of ozone causal determinations for climate and UV-B effects.	10-33
		10-34

References _____

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ACRONYMS AND ABBREVIATIONS

129	mouse strain (129S1/SvlmJ)	AOT60	seasonal sum of the difference
α	alpha, ambient exposure factor	ACTO	between an hourly concentration
α-ATD	alpha 1-antitrypsin deficiency		at the threshold value of 60 ppb,
a-SMA	alpha-smooth muscle actin		minus the threshold value of 60 ppb
a-tocopherol	alpha-tocopherol	AOTx	family of cumulative, cutoff
α-ΤΟΗ	alpha tocopherol	NOTX	concentration-based exposure
a	air exchange rate of the		indices
u	microenvironment	AP	activated protein
A2	climate scenario in IPCC	A2p	climate scenario in IPCC (preliminary version of A2)
AADT	annual average daily traffic	APEX	Air Pollutants Exposure (model)
A1B	climate scenario in IPCC	APHEA(2)	Air Pollution on Health: a
ABA	abscisic acid		European Approach (study)
ABI	abscisic acid insensitive	APHENA	Air Pollution and Health: A
A1c	glycosylated hemoglobin blood test		European and North American Approach
Ach	acetylcholine	АроВ	apolipoprotein B
ACM	(Harvard University) Atmospheric	ApoE	apolipoprotein E
	Chemistry Modeling (Group)	APX	ascorbate peroxidase
ACS	American Cancer Society	aq	aqueous form: (aq)O ₃
ACS-CPSII	ACS Cancer Prevention Study II	AQCD	Air Quality Criteria Document
ADC	arginine decarboxylase	AQI	Air Quality Index
ADSP	Adirondack State Park, NY	AQS	(U.S. EPA) Air Quality System
AER	air exchange rate		(database)
	ascorbic acid; ascorbate	AR	acoustic rhinometry
AHR	airway(s) hyperresponsiveness, airway(s) hyperreactivity	AR4	Fourth Assessment Report (AR4) from the IPCC
AhR	aryl hydrocarbon receptor	AR5	Fifth Assessment Report (AR5)
AHSMOG	(California Seventh Day) Adventist Heath and Smog (Study)	ARG	from the IPCC arginase variants (ex., ARG1,
AI	alveolar interstitial	1510	ARG2, ARG1h4)
AIC(s)	Akaike's information criterion	ARIC	Atherosclerosis Risk in Communities
AIRS	Aerometric Information Retrieval System; Atmospheric Infrared Sounder (instrument)	ARIES	(Atlanta) Aerosol Research and Inhalation Epidemiology Study
A/J	mouse strain	atm	atmosphere
Ala-9Val	genotype associated with	ATP	adenosine triphosphate
	Manganese superoxide dismutase (MnSOD) gene	ATPase	adenosine triphosphatase; adenosine triphosphate synthase
AM	alveolar macrophage(s)	ATS	American Thoracic Society
ANF	atrial natriuretic factor	avg	average
AOT20	seasonal sum of the difference between an hourly concentration	AVHRR	advanced very high resolution radiometer
	at the threshold value of 20 ppb, minus the threshold value of 20 ppb	β	beta, beta coefficient; regression coefficient; standardized coefficient; shape parameter;
AOT30	seasonal sum of the difference		scale parameter
	between an hourly concentration at the threshold value of 30 ppb,	В	boron
	minus the threshold value of	B1	climate scenario in IPCC
	30 ppb	B6	mouse strain (C57BL/6J)
AOT40	seasonal sum of the difference	BAL	bronchoalveolar lavage
	between an hourly concentration at the threshold value of 40 ppb,	BALB/c	mouse strain
	minus the threshold value of	BALF	bronchoalveolar lavage fluid
	40 ppb	bb	bronchials

DD	han shiel similar	CAMP	
BB BC	bronchial airways black carbon	CAMP	Childhood Asthma Management Program
B cells	bone-marrow-derived lymphocytes; B lymphocytes	CAMx	Comprehensive Air Quality Model, with extensions
B6C3F1	mouse strain	CAN	Canada
BDNF	brain-derived neurotrophic factor	CAP(s)	concentrated ambient particles
BEAS-2B	human bronchial epithelial cell line	CAR	centriacinar region
BEIS	Biogenic Emissions Inventory System	CASAC	Clean Air Scientific Advisory Committee
BELD	Biogenic Emissions Landcover Database	CASTNET	Clean Air Status and Trends Network
BIPM	International Bureau of Weights	CAT	catalase
BM	and Measures basement membrane	СВ	carbon black; CMAQ mechanisms (ex., CB04, CB05, CB06)
BMI		C57BL/6	mouse strain
	body mass index	C57BL/6J	mouse strain
BNP	β -type natriuretic peptide	CBSA	core-based statistical area
BP	blood pressure	C/C	carbon of total carbon
BPD	biparietal diameter	CCSP	Clara cell secretory protein
bpm Dr	breaths per minute	CD	cluster of differentiation (various
Br	bromine	02	receptors on T-cells: CD8+, CD44,
BRFSS	Behavioral Risk Factor Surveillance System		etc.);
BS	black smoke	CD-1	criteria document (see AQCD) mouse strain
BSA	bovine serum albumin;	CDC	Centers for Disease Control and
	body surface area	CDC	Prevention
Bsp, BSP	black smoke particles	CF	charcoal-filtered; carbon filtered air
Bt, BT, bt	Bacillus thuringiensis; bacterium proteins used in pesticides (or	CF2	twice-filtered air (particulate filter and activated charcoal filter)
577	genetically engineered plants produce Bt toxin)	C-fibers	afferent, slow, unmylenated nerves innervating the respiratory system
BTEX	family of compounds (benzene, toluene, ethylbenzene, and xylene)	CFR	Code of Federal Regulations
BW	body weight	CGRP	calcitonin gene-related peptide
C	carbon; concentration;	CH₃	methyl group
-	(Vitamin C, ascorbate)	CH_4	methane
°C	degrees Celsius	C_2H_2	acetylene
¹³ C	carbon-13 isotope	C_2H_4	ethylene
C3	mouse strain (C3H/HEJ)	C3H	mouse strain (C3H/HEJ or
C3	plants that use only the Calvin		C3H/OuJ)
	cycle for fixing the carbon dioxide from the air	C ₃ H ₆	propylene
C4	plants that use the Hatch-Slack	CHAD	Consolidated Human Activity Database
	cycle for fixing the carbon dioxide	CH₃Br	methyl bromide
040.0	from the air	CH₃-CHO	acetaldehyde
C16:0	palmitic acid (saturated fatty acid)	CH₃CI	methyl chloride
C18:1	unsaturated fatty acid	CH₃-CO	acetyl radical(s)
Ca	calcium	CHD	coronary heart disease
C _a	ambient concentration	CHF	congestive heart failure
[Ca]	calcium concentration	C₂H₅–H	ethane
Ca ²⁺	calcium ion	C3H/HeJ	mouse strain
CA	Canada (ICD-10-CA)	CH₃I	methyl iodide
CAA	Clean Air Act	CHIP	Effects of Elevated Carbon Dioxide
CALINE4	California line source dispersion model for predicting air pollutant concentrations near roadways		and Ozone on Potato Tuber Quality in the European Multiple Site Experiment
CAM	plants that use crassulacean acid	CH ₃ O ₂ •	methyl peroxy (radical)
	metabolism for fixing the carbon dioxide from the air	CH ₃ OOH	acetic acid; methyl hydroperoxide
		CHS	Child Health Study

CI	confidence interval(s)	схс	chemokine family of cytokines,
C _j	airborne O ₃ concentration at		with highly conserved motif:cys- xxx-cys (CXC) amino acid group
CI	microenvironment j chlorine	CXCR2	CXC chemokine receptor 2
CI	chlorine ion		(CXCR2)
Cl ₂	chlorine gas	CXR	Chest (x-ray) radiograph(s)
CLE	Current Legislation (climate	CyS	protein cysteines
CLM	scenario in IPCC) chemiluminescence method	Cys-LT	cysteinyl leukotrienes (LTC ₄ , LTD ₄ , LTE ₄)
	nitryl chloride	cyt	cytosolic-free
-	centimeter(s)	Δ, δ	delta, difference; change
cm cm²	square centimeters	ΔFEV_1	change in FEV ₁
CM	Clinical Modification (ICD-9-CM)	ΔV_{D}	change in dead space volume of
CMAQ	Community Multi-scale Air Quality	_	the respiratory tract
CINAQ	modeling system	2-D	two-dimensional
CN	constant atmospheric nitrogen	3-D	three-dimensional
	deposition (in PnET-CN ecosystem model)	DAHPS	3-deoxy-D-arabino-heptulosonat- 7-phosphate synthase
CNA	continental North America	DBP	diastolic blood pressure
CNS	central nervous system	DC(s)	dendritic cell(s)
СО	carbon monoxide; Cardiac output	DDM	direct decoupled method
CO_2	carbon dioxide	DEP(s)	diesel exhaust particle(s)
COD	coefficient of divergence;	df	degrees of freedom
Col-0	coefficient of determination (Arabidopsis ecotype) Columbia-0	DGGE	denaturing gradient gel electrophoresis
COP	Conference of Parties (to the	DHA	dehydroascorbate
001	UNFCCC)	DHAR	dehydroascorbate reductase
COPD	chronic obstructive pulmonary	DHBA	2,3-dihydroxybenzoic acid
	disease	DLEM	Dynamic Land Ecosystem Model
COX-2	cyclooxygenase 2 enzyme	dm ³	cubic decimeter(s)
C-R	concentration-response	DNA	deoxyribonucleic acid
CRA	Centro di ricerca per la cerealicoltura (CRA) [The Centre	DOAS	differential optical absorption spectroscopy
	for Cereal Research] – Unit 5: The	DOC	dissolved organic carbon
	Research Unit for Cropping Systems in Dry Environments in	DR	type of human leukocyte antigens
	Bari, Italy (water-stressed		(HLA-DR)
	conditions)	dt	Portion of time-period spent in
CRP	C-reactive protein		microenvironment j
CS	corticosteroid	DTH	delayed-type hypersensitivity
CSA	cross-sectional area; combined	DU	Dobson unit(s)
aah Cah	statistical area	DW	dry weight
csb, Csb	cockayne syndrome (cb) gene/protein group A	E	embryonic day (ex., E15, E16, etc); [Vitamin] E
CSF	colony-stimulating factor	Ea	exposure to pollutant of ambient
CST	central standard time		origin
CSTR	continuous stirred tank reactor	EBC	exhaled breath condensate (fluid)
CSV	comma-separated values (a spreadsheet format)	EC	elemental carbon
СТ	computer tomography	ECE	endothelin converting enzyme(s) [i.e., ECE-1]
CTM(s)	chemical transport model(s)	ECG	electrocardiogram
()	,	ECOPHYS	v
cum avg CUOt	cumulative average The cumulative stomatal uptake of	ECOFIIIS	physiological process modeling to predict the response of aspen
CUOL	O_3 , using a constant O_3 uptake rate threshold (t) of nmol/m ² /sec		forest ecosystems (modeling growth and environmental stress in
CV, C.V.	coefficient of variation	50	Populus)
CV, C.V.	cultivar	ED	emergency department; embryonic day (ex., ED5, ED20)
CVD	cardiovascular disease		

5054			
EGEA	(The) Epidemiology (study on) Genetics and Environment of	FEM	Federal equivalent method
	Asthma, (adults and children with	FeNO	exhaled nitric oxide fraction
EGEA2	asthma) follow-up study on EGEA (adults	FEV ₁	forced expiratory volume in 1 second
-	with asthma only)	FHM	(USDA Forest Service) Forest Health Monitoring Program
EHC-93	ambient PM reference sample (urban dust [air particles] collected in Ottawa Canada)	FIA	(USDA Forest Service) Forest Inventory and Analysis Program
ELF	extracellular lining fluid	F _{inf}	infiltration factor
EMI	(U.S. EPA) Exposure Model for Individuals	F _{inf,i}	infiltration factor for indoor environment (i)
E _{na}	exposure to pollutant of nonambient origin	FLAG	Federal land managers' air quality related values workgroup
ENA-78	epithelial cell-derived neutrophil- activating peptide 78	F _{LRT}	fractional uptake efficiency of the lower respiratory tract (LRT)
eNO	exhaled nitric oxide	F _{nose}	fractional uptake efficiency via
eNOS	endothelial nitric oxide synthase		nasal absorption
ENVISAT	(EAS) Earth Observation satellite	Fo	fraction of time spent in outdoor
EOTCP	European Open Top Chamber	FPM	microenvironments Forest Pest Management
EP	Programme	FR	Federal Register
EPA	epithelial cells U.S. Environmental Protection	FRAP	ferric reducing ability of plasma
EPA	Agency	FRC	functional residual capacity
EPIC	European Prospective	FRM	Federal reference method
	Investigation into Cancer and Nutrition	F _{RT}	fractional uptake efficiency of the respiratory tract (RT)
ER	emergency room	Fst0₁	flux cut off threshold
ESA ET	European Space Agency extrathoracic; endothelin	F _{URT}	fractional uptake efficiency of the upper respiratory tract (URT)
	(i.e., ET-1)	FVC	forced vital capacity
ET ₁	anterior nasal passages within the extrathoracic (ET) region	Fv/Fm	a ratio: a measure of the maximum efficiency of Photosystem II
ET ₂	oral airway and posterior nasal	FVI	fruits and vegetables index
	passages within the extrathoracic (ET) region	γ	gamma
ETS	environmental tobacco smoke	γ-ΤΟΗ	gamma-tocopherol
EU	European Union	g, kg, mg, µg, ng, pg	gram(s), kilogram(s), milligram(s),
EUS	eastern U.S.		microgram(s), nanogram(s), picogram(s)
Φ	Phi; calculated efficiency	G	granulocyte; guanosine
Φ ΦPSII-max	-		gram(s); gaseous form: $(g)O_3$
ΨF SII-IIIdx	maximum photochemical effective quantum yield of PSII	g GAM	
f	Fraction of the relevant time period		generalized additive model(s)
F	female	g _{bs}	conductance through boundary layer and stomata
F344	Fischer 344 (rat strain)	GCLC	(glutathione genetic variant)
F2a	8-isoprostane (major F2 prostaglandin [8 iso-PGF2a])		glutamate-cysteine ligase catalytic subunit
FA	filtered air	GCLM	(glutathione genetic variant)
FACE	free-air– CO_2 enrichment (system)		glutamate-cysteine ligase modifier
FACES	Fresno Asthmatic Children's	C CSE	subunit
	Environment Study	G-CSF	granulocyte colony-stimulating factor (receptor)
f _Β FC	frequency of breathing	GD	gestational day
	fibrocartilaginous coat	GEE	generalized estimating equations
FEF	forced expiratory flow	GEOS	(NASA) Goddard Earth Observing
FEF ₂₅₋₇₅	forced expiratory flow between the times at which 25% and 75% of	CEOSE	System model
	the vital capacity is reached	GEOS5	GEOS version 5
FEFx	forced expiratory flow after (x)% vital capacity (e.g., after 25, 50, or	GEOS-Chem	GEOS-Chemistry (tropospheric model)
	75% vital capacity)	GFAP	glial fibrillary acidic protein

GH	growth hormone	HeJ	O ₃ -resistant C3H mouse strain
GHG	greenhouse gas		(C3H/HeJ)
GLM(s)	generalized linear model(s)	HEPA	high efficiency particle air (filter)
GMAO	(NASA) Global Modeling and Assimilation Office	HERO	Health and Environmental Research Online, NCEA Database System
GM-CSF	granulocyte macrophage colony- stimulating factor	12-HETE	12-Hydroxyeicosatetraenoic acid
GOME	(ESA) Global Ozone Monitoring Experiment (spectrometer)	HF	(HRV signal) high-frequency power
GOMOS	Global Ozone Monitoring by	HFCs	hydrofluorocarbons
	Occultation of Stars (ESA	Hg	mercury
	ENVISAT spectrometer measuring	HHP-C9	1-hydroxy-1-hydroperoxynonane
000	long-term trends in O_3)	HIST	histamine
G6P	glucose-6-phosphate	HLA	human leukocyte antigen
G6PD	glucose-6-phosphate dehydrogenase	HLA-DR	human leukocyte antigen receptor genes
GPP	gross primary production	HMOX	Heme oxygenase
G-proteins	GTPases	HMOX-1	heme-oxygenase-1
GPT	gas phase titration		(polymorphism)
GR	glutathione reductase	HNE	4-hydroxynonenal
GSH	glutathione; reduced glutathione	HNO ₂	nitrous acid
GSO3 ⁻ /GSO3 ²⁻	guanine sulfonates	HNO ₃	nitric acid
GSR	glutathione reductase	HNO ₄	pernitric acid
GSS	glutathione synthetase	НО	hydroxyl; heme oxygenase
GSSG	glutathione disulfide	HO•	hydroxyl radical
GST	glutathione S-transferase	HO-1	heme oxygenase 1
GSTM1	glutathione S-transferase polymorphism M1 genotypes	HO ₂ •	hydroperoxyl; hydroperoxy radical; protonated superoxide
GSTP1	(GSTM1-null, -GSTM1-sufficient) glutathione S-transferase	HO₃•	protonated ozone radical
GOTFT	polymorphism P1 genotypes	H ₂ O	water
GTP	guanosine triphosphate	H_2O_2	hydrogen peroxide
GTPases	G-proteins/enzymes	H_3O^+	hydronium ion
GWP	global warming potential	HOCH ₂ OOH	hydroxymethylhydroperoxide
GxE	gene-environmental interaction	HONO	nitrous acid
h	hour(s)	HO ₂ NO ₂	peroxynitric acid
h/day	hour(s) per day	HOONO	pernitrous acid
H; H+; H∙	atomic hydrogen, hydrogen ion;	HOX	hydrogen radical(s)
	hydrogen radical	hPa	hectopascal
³ Н	radiolabeled hydrogen; tritium	HPLC	high-pressure liquid
H ₂	molecular hydrogen		chromatography
ha	hectare	HPOT	13-hydroperoxide linolenic acid
HA	hyaluronic acid, hospital admission	HR	heart rate, hazard ratio
HA(s)	hospital admission(s)		maximum heart rate
Hb	hemoglobin	HRP	horseradish peroxidase
HbA1c	glycosylated hemoglobin (blood	HRV	heart rate variability
	test)	HSC	Houston Ship Channel (Texas)
HC(s)	hydrocarbon(s)	hs-CRP	high-sensitivity C-reactive protein
HCFC(s)	hydrochlorofluorocarbon(s)	H ₂ SO ₄	sulfuric acid
НСНО	formaldehyde	HSP	high speed pellet (after centrifuge
H ₂ CO	formaldehyde	HSP70	spin) boat shock protoin 70
HCO•	formyl (radical)	HSS	heat shock protein 70 high speed supernatant (after
HDM	house dust mite	100	centrifuge spin)
2HDM	second-highest daily maximum	5-HT	5-hydroxytryptamine
	house dust mite allergen	hv	Energy per photon of
³ He	non-radioactive isotope of helium		electromagnetic energy at frequency v

HVAC	heating, ventilation, and air conditioning	INRA	National agronomical research institute (INRA) in Thiverval-
Hz	hertz		Grignon. France (adequately-
I	iodine		watered conditions)
IARC	International Agency for Research on Cancer	INTRASTAND	a stand-level model designed for hourly, daily and annual integration of forest carbon and water cycle
IAS	interalveolar septum		fluxes
IBM	individual-based model or modeling	I/O IOM	indoor-outdoor ratio Institute of Medicine
IC	inspiratory capacity;	i.p.	intraperitoneal (route)
	intracloud (lightning flash)	IPCC	Intergovernmental Panel on
ICAM-1	intercellular adhesion molecule 1		Climate Change
ICARTT	International Consortium for Atmospheric Research on Transport and Transformation	IPCC-A2	Intergovernmental Panel on Climate Change 2nd Assessment Report
ICAS	Inner City Asthma Study	IPCC-AR4	Intergovernmental Panel on
ICC	intraclass correlation coefficient		Climate Change 4th Assessment
ICD	implantable cardioverter defibrillator(s); International Classification of Diseases	IPCC-AR5	Report Intergovernmental Panel on Climate Change 5th Assessment
ICD-9	International Classification of		Report
ICD-10	Disease 9th revision International Classification of Disease 10th revision	IPCC-TAR	Intergovernmental Panel on Climate Change Third Assessment Report
ICEM	Indoor Chemistry and Exposure Model	IPMMI	International Photolysis Frequency Measurement and Modeling Inter-
ICNIRP	International Commission on Non-ionizing Radiation Protection	IQR	comparison interquartile range
ICP Forests	International Cooperative	IR	infrared
	Programme on Assessment of Air	I/R	ischemia-reperfusion
ICU	Pollution Effects on Forests Intensive Care Unit	IRIS	Integrated Risk Information System
ICVE	ischemic cerebrovascular events	IRP	Integrated Review Plan for the
IDW	inverse-distance-weighted		Ozone National Ambient Air
IFN	interferon (e.g., IFN-())		Quality Standards
IFN-y	interferon-gamma	ISA	Integrated Science Assessment
lg	immunoglobulin (e.g., IgE)	ISCCP	International Satellite Cloud Climatology Project
lgA	immunoglobulin A	ISO	International Standards
lgE	immunoglobulin E	100	Organization
IGF-1	insulin-like growth factor 1	8-iso-PGF	8-isoprostane
lgG	immunoglobulin G	IT	intratracheal
lgM	immunoglobulin M	IU	International Units
IHD	ischemic heart disease	IUGR	intrauterine growth restriction
IL	interleukin (e.g., IL-2, IL-4, IL-6,	i.v.	intravenous (route)
	IL -8, etc.)	IVF	in vitro fertilization
IL-1β	interleukin-1β	j	Microenvironment
lle	isoleucine	JA	jasmonic acid
i.m.	intramuscular (route)	Jmax	maximum rate of electron transport
IMPACT	Interactive Modeling Project for		(for regeneration of RuBP)
	Atmospheric Chemistry and Transport	JNK	jun N-terminal kinase
IMPROVE		JPL	Jet Propulsion Laboratory
IMFROVE	Interagency Monitoring of Protected Visual Environment	К	kappa
IN	intranasal	кВ	kappa B
INF	interferon	k	dissociation rate; root:shoot
inh	inhalation		allometric coefficient; rate of O_3
iNKT	invariant (type I) natural killer T-cell	к	loss in the microenvironment potassium
iNOS	inducible nitric oxide synthase	K ⁺	potassium ion
	madelible mittle oxide synthase		

Ka	intrinsic mass transfer	LOSU	level of scientific understanding
	coefficient/parameter	LOWESS	locally weighted scatter plot
KC	keratinocyte-derived chemokine		smoother
kg	kilogram	LOX-1	Lipoxygenase; lectin-like oxidized low density lipoprotein receptor-1
K _g	mass transfer coefficient for gas phase	LPS	lipopolysaccharide
kHz	, kilohertz	LRS	lower respiratory symptoms
kJ	kilojoules	LRT	lower respiratory tract; lower
KI	mass transfer coefficient for liquid		airways; Long range transport
	phase	LST	local standard time
km	kilometer	LT	leukotriene (e.g., LTB4 , LTC 4,
KM	particle optical reflectance		LTD4, LTE4); local time
KML	keyhole markup language	LT-α LTA	lymphotoxin-α
KMZ	zipped KML computer language	LUR	lymphotoxin-alpha land use regression
КО	knockout	LVEDD	left ventricular chamber
Kr	reaction rate constant		dimensions at end diastole
KROFEX	Krauzberg Ozone Fumigation Experiment	LVEDP	left ventricular end diastolic pressure
L, dL, mL, µL	Liter, deciLiter, milliLiter, microLiter	LWC	liquid water content
LO	Lag (e.x., Lag 0, Lag 1, etc.)	μ	mu, micro
LAI	leaf area index	hed	microequivalent
LBL	Lawrence Berkeley Laboratory	μg	microgram
LBLX	Lawrence Berkeley Laboratory model including airflow from	µg/m ³	micrograms per cubic meter
	natural ventilation	μm	micrometer, micron
Lb(s)	pound(s)	m, cm, µm, nm	meter(s), centimeter(s),
LBW	low birth weight		micrometer/[micron](s),
LC ₅₀	median lethal concentration	Μ	nanometer(s) male
LCL	lower 95th% confidence limit	M, mM, μM, nM, pM	Molar, milliMolar, microMolar,
LDH	lactate dehydrogenase	ινι, πινι, μινι, πινι, μινι	nanoMolar, picoMolar
LDL	low-density lipoprotein ; lower	m²	square meters
LF	detectable level	m³	cubic meters
	(HRV signal) low-frequency power	M#	Month (M1 Month1; M2 Month2;
LEULK	low frequency/high frequency (ratio)		M3 Month3; M4 Month4)
LFT	lower free troposphere	M2	type of muscarinic receptor
LI	labeling index	M7	7-hour seasonal mean
LIDAR	Light Detection and Ranging	M12	12-hour seasonal mean of O_3
	(remote sensing system)	ma	moving average
LIF	laser-induced fluorescence	mAOT	modified accumulated exposure over threshold
LINKAGES	individual-based model of forest succession	МАР	mitogen-activated protein; mean arterial pressure
LIS	lateral intercellular space	МАРК	mitogen-activated protein
LLJ	low-level jet		kinase(s), MAP kinase
L/min	liters per minute	MAQSIP	Multiscale Air Quality Simulation
Ln	Natural logarithm		Platform (model)
LnRMSSD	natural log of RMSSD; measure of HRV	MARAT	Mid-Atlantic Regional Assessment Team
InSDNN	natural log of the standard deviation of NN intervals in an FKG	MARCO	Macrophage receptor with collagenous structure
LOAEL	lowest observed adverse effect	max	maximum
	level	MBL	marine boundary layer
LOD	limit of detection	MCA	minimum cross-sectional area
LOEL	lowest-observed-effect level	MCCP	Mountain Cloud Chemistry Program
LOESS			
	locally weighted scatterplot	Mch: MCh	methacholine
LOP	locally weighted scatterplot smoothing lipid ozonation products	Mch; MCh MCM	methacholine master chemical mechanism

MCP-1	monocyte chemotactic protein 1	MOBILE6	vehicle emissions modeling software version 6; replaced by
MDA	malondialdehyde		MOVES
MDAR	monodehydroascorbate reductase	MODNR	Missouri Department of Natural
MDI	Mediterranean diet index	-	Resources
MDL	minimum detection level	MONICA	Monitoring of Trends and
MED	minimal erythema dose		Determinants in Cardiovascular
MEF _{50%}	maximal midexpiratory flow at 50%	No	Disease
	of forced vital capacity	MoOx	molybdenum oxides
MEGAN	model of emissions of gases and aerosols from nature	MOSES	Met Office Surface Exchange Scheme
MeJA	methyl jasmonate	MOVES	Motor Vehicle Emission Simulator
MENTOR	Modeling Environment for Total Risk Studies		(replaced MOBILE6; for estimating emissions from cars, trucks, and
METs	metabolic equivalent unit(s) [of work]	MOZAIC	motorcycles Measurement of Ozone and Water
MFR	Maximum Feasible Reduction		Vapor by Airbus In-Service Aircraft
Mg	magnesium	MOZART	Model for Ozone and Related chemical Tracers
MGDG	monogalactosyl diacylglycerol	MPAN	peroxymethacryloyl nitrate;
mg/m ³	milligrams per cubic meter		peroxy-methacrylic nitric anhydride
MHC	major histocompatibility complex	MPO	myeloperoxidase
mi	mile(s)	MQL	Minimum quantification limit
MI	myocardial infarction, "heart	MRI	magnetic resonance imaging;
	attack"		Midwest Research Institute;
MIESR	matrix isolation electron spin		Meteorological Research Institute
	resonance (spectroscopy)	mRNA	messenger RNA
min	minute; minimum	ms	millisecond(s)
MIP MIP-2	macrophage inflammatory protein	MS	mass spectrometry; Mt. Moosilauke site
WIF-2	macrophage inflammatory protein 2	MSA	Metropolitan Statistical Area;
mL	milliliter		methane sulfonic acid
mL/min	milliliter(s) per minute	MSL	mean sea level
MLN	mediastinal lymph node	MS/MS	tandem mass spectrometry
Mm	megameter	MT	million ton(s); metric ton(s)
mm	millimeter(s)	MT, Mt	metallothionein
MM Mt.	Mt. Mitchell site	MT1	mitochondria
MM5	National Center for Atmospheric	MTBE	methyl-tertiary-butyl ether
	Research/Penn State Mesoscale	mtDNA	mitochondrial DNA
	Model (version 5)	Mtn	mountain
MMAD	mass median aerodynamic diameter; mass median	MW	molecular weight
MMEF	aerodynamic density maximal midexpiratory flow	MyD88	myeloid differentiation primary response gene 88
		n, N	number; number of observations
mmHg MMMD	millimeters of mercury mean maximum mixing height	Ν	nitrogen; North; nasal exposure by natural breathing
MMP-2	depth matrix metalloproteinase-2	¹⁵ N	nitrogen-15, stable isotope of
MMP-3	matrix metalloproteinase-3		nitrogen
MMP-9	metalloproteinase-9	N ₂	molecular nitrogen; nonreactive nitrogen
MMSP	Mount Mitchell State Park, NC	Na	sodium
Mn	manganese	NA	noradrenaline: North American
M/N	pooled data from mouth and nasal	NA; N/A	not available; not applicable
	exposure	Na ⁺	sodium ion
MnSOD	Manganese superoxide dismutase	NAAQS	National Ambient Air Quality
mo	month(s)		Standards
MOA(s)	mode(s) of Action	NAD	nicotinamide adenine nucleotide
MOBILE	(U.S. EPA) mobile vehicle emission factor model (on-road vehicles)	NADH	reduced nicotinamide adenine dinucleotide; nicotinamide adenine dinucleotide dehydrogenase

NADP	National Atmospheric Deposition	NH ₃	ammonia
	Program	NH_4^+	ammonium ion
NADPH	reduced nicotinamide adenine dinucleotide phosphate	NH ₄ HSO ₄	ammonium bisulfate
NADPH-CR	reduced nicotinamide adenine	(NH ₄) ₂ HSO ₄	ammonium sulfate
NADEN-CK	dinucleotide phosphate - cytochrome c reductase	NHANES	National Health and Nutrition Examination Survey
NaE	sodium erythorbate	NHANES III	National Health and Nutrition Examination Survey III
NAG	N-acetyl-glucosaminidase	NHAPS	,
Na-K-ATPase	sodium-potassium-dependent adenosine triphosphatase	-	National Human Activity Pattern Survey
NAMS	National Ambient Monitoring Stations	NHEERL	(U.S. EPA) National Health and Environmental Effects Research Laboratory
NAPAP	National Acid Precipitation Assessment Program	NHIS	National Health Interview Survey
NAPBN	National Air Pollution Background	(NH ₄) ₂ SO ₄ NIH	ammonium sulfate National Institutes of Health
	Network	NIST	National Institute of Standards and
NARE	North Atlantic Regional Experiment	-	Technology
NARSTO	North American Regional Strategy	NK	natural killer cells; neurokinin
	for Tropospheric Ozone	NKT	natural killer T-cells
NAS	National Academy of Sciences; Normative Aging Study	NL	nasal lavage
NACA	National Aeronautics and Space	NLF	nasal lavage fluid
NASA	Administration	NM	National Monument
NBS	National Bureau of Standards	NMHC(s)	nonmethane hydrocarbon(s)
NBTH	3-methyl-2-benzothiazolinone acetone azine	NMMAPS	National Morbidity, Mortality, and Air Pollution Study
NCEA	National Center for Environmental	NMOC(s)	nonmethane organic compound(s)
-	Assessment	NMVOCs	nonmethane volatile organic compounds
NCEA-RTP NCHS	NCEA Division in Research Triangle Park, NC National Center for Health	NN	normal-to-normal (NN or RR) time interval between each QRS complex in the EKG
	Statistics	NNK	4-(N-nitrosomethylamino)-1-
NCICAS	National Cooperative Inner-City Asthma Study		(3-pyridyl)-1-butanone
NCLAN	National Crop Loss Assessment Network	nNOS	neuronal nitric oxide synthase (NOS)
NCore	National Core multi-pollutant	NO	nitric oxide
	monitoring network	·NO	nitric oxide concentration (interpunct NO)
NC-R	resistant clones of white clover	NO ₂	nitrogen dioxide
NC-S	sensitive clones of white clover	NO ₃ ; NO ₃ •	nitrate, nitrate radical
ND; n.d.	not detectable; not detected; no data	NO ₃ ⁻	nitrate, nitrate ion
2ndHDM	second-highest daily maximum	N ₂ O	nitrous oxide
NDF	neutral detergent fiber	N_2O_5	dinitrogen pentoxide
NEE	net ecosystem CO ₂ exchange	NOAA	National Oceanic and Atmospheric Administration
NEI	National Emissions Inventory	NOAEL	no observed adverse effect level
NEM	National Ambient Air Quality Standards Exposure Model	NOS	nitric oxide synthase (types,
NEP	Net Ecosystem Production	NO	NOS-1, NOS-2, NOS-3)
NERL	National Exposure Research Laboratory	NO _x	nitrogen oxides, oxides of nitrogen (NO + NO ₂)
NESCAUM	Northeast States for Coordinated Air Use Management	NO _Y	sum of NO _x and NO _z ; odd nitrogen species; total oxidized nitrogen
NF	National Forest; non-filtered air	NOz	sum of all inorganic and organic
NF-ĸB	nuclear factor kappa B		reaction products of NO _X (HONO, HNO ₃ , HNO ₄ , organic nitrates,
ng	nanogram(s)		particulate nitrate, nitro-PAHs,
NGF	nerve growth factor		etc.)
NH	northern hemisphere	NP	National Park

NPP	not primary production	OPECs	Outdoor Plant Environment
NPS	net primary production National Park Service, U.S.	OFECS	Chambers
	Department of the Interior	OR	odds ratio
NQO1	NAD(P)H-quinone oxidoreductase (genotype)	ORD	Office of Research and Development
NQO1wt	NAD(P)H-quinone oxidoreductase wild type (genotype)	OSHA	Occupational Safety and Health Administration
NR	not reported	OTC	open-top chamber
Nr	reactive nitrogen	OuJ	O ₃ -sensitive C3H mouse strain
NRC	National Research Council		(C3H/OuJ)
Nrf-2	nuclear factor erythroid 2-related	OVA	ovalbumin
	factor 2	OX	odd oxygen species; total oxidants
Nrf2-ARE	NF-e2-related factor 2-antioxidant response element	OxComp	oxidative capacity of the atmosphere
NS; n.s.	nonsignificant; non-smoker; national seashore; natural spline	oz P	ounce(s) pressure in atmospheres; plants
NSAID	non-steroidal anti-inflammatory agent		grown in pots; phosphorus; penetration fraction of O_3 into the
NSBR	nonspecific bronchial responsiveness		microenvironment; pulmonary region
NSF	National Science Foundation	р	probability value
NTE	nasal turbinate epithelial (cells)	P450	cytochrome P450
NTN	National Trends Network	p53	cell cycle protein gene
NTP	National Toxicology Program	P90	90th percentile of the absolute
NTRMs	NIST Traceable Reference Materials	PACF	difference in concentrations partial autocorrelation function of
NTS	nucleus of the solitary tract (in brainstem)	PAD	the model residuals peripheral arterial disease;
NWR	national wildlife refuge	5.5	pollutant-applied dose
NWS	National Weather Service	PAF	platelet-activating factor; paroxysmal atrial fibrillation
NZW	New Zealand white (rabbit)	PAH(s)	polycyclic aromatic hydrocarbon(s)
0 ¹⁸ 0	oxygen; horizon forest floor oxygen-18, stable isotope of	PAI-1	plasminogen activator fibrinogen inhibitor-1
-	oxygen	PAL	phenylalanine ammonia lyase
O ₂	molecular oxygen	PAMS	Photochemical Assessment
O ₂ ⁻	superoxide		Monitoring Stations network
O ₂ •	superoxide radical	PAN	peroxyacetyl nitrate
¹ O ₂	singlet oxygen	PaO ₂	arterial oxygen pressure
O ₃	ozone	PAPA	Public Health and Air Pollution in
¹⁸ O ₃	(oxygen-18 labeled) ozone		Asia
O ₃ *	electronically excited ozone	PAR	photosynthetically active radiation; proximal alveolar region
OAQPS	Office of Air Quality Planning and Standards	P _{atm}	Pressure in atmospheres
OAR	Office of Air and Radiation	p-ATP	para-acetamidophenol
OBMs	observationally based methods	Pb	Lead
OC	organic carbon	PBL	planetary boundary layer;
OD	outer diameter; optical density		peripheral blood lymphocytes
O(¹ D)	electronically excited oxygen atom	PBM	population-based model or
OH, OH•	hydroxyl group, hydroxyl radical		modeling
8-OHdG	8-hydroxy-2'-deoxyguanosine	PBN	C-phenyl N-tert-butyl nitrone
OLS	ordinary least squares	PBPK	physiologically based pharmacokinetic (model)
OMI	Ozone Monitoring Instrument	PBS	phosphate buffered saline
ON	Ontario	PC	phosphatidylchloline
ONOO ⁻	peroxynitrate ion	PC ₂₀	provocative concentration that
O(³ P)	ground-state oxygen atom		produces a 20% decrease in
OPE	ozone production efficiency		forced expiratory volume in 1 second

$PC_{20}FEV_1$	provovative concentration that produces a 20% decrease in FEV ₁	рН	relative acidity; Log of the reciprocal of the hydrogen ion
PC ₅₀	provocative concentration that	DUA	
	produces a 50% decrease in forced expiratory volume in	PHA	phytohemagglutinin A
	1 second	PI	phosphatidylinositol; probability interval; posterior interval
PCA	principal component analysis	PIF	peak inspiratory flow
PC-ALF	1-palmitoyl-2-(9-oxonononoyl)-sn-	PiZZ	respiratory phenotype
505	glycero-3-phosphocholine	PK	pharmacokinetics
PCD	programmed cell death	рКа	dissociation constant
PCI	picryl chloride	PLFA	phospholipid fatty acid
pCNEM	Canadian version of National Ambient Air Quality Standards	PM	particulate matter
	Exposure Model	PM _X	Particulate matter of a specific size
PCO ₂	Average partial pressure of O ₂ in lung capillaries		range not defined for regulatory use. Usually X refers to the 50% cut point, the aerodynamic
pCO ₂	partial pressure of carbon dioxide		diameter at which the sampler
PCR	polymerase chain reaction		collects 50% of the particles and
PCR-DGGE	PCR–denaturing gradient gel electrophoresis		rejects 50% of the particles. The collection efficiency, given by a penetration curve, increases for
PD	pregnancy day		particles with smaller diameters
PD ₂₀	provocative dose that produces a 20% decrease in FEV ₁		and decreases for particles with larger diameters. The definition of
$PD_{20}FEV_1$	provocative dose that produces a 20% decrease in FEV ₁		PM _X is sometimes abbreviated as "particles with a nominal aerodynamic diameter less than or
PD ₁₀₀	provocative dose that produces a 100% increase in sRAW		equal to X μ m" although X is usually a 50% cut point.
$PD_{100}S_{Raw}$	provocative dose that produces a 100% increase in $S_{\mbox{\scriptsize Raw}}$	PM _{2.5}	In general terms, particulate matter with an aerodynamic diameter less
PDI	pain on deep inspiration		than or equal to a nominal 2.5 μm;
PE	postexposure, phosphatidylethanolamine		a measurement of fine particles in regulatory terms, particles with an upper 50% cut-point of 2.5 µm
PEF	peak expiratory flow		aerodynamic diameter (the 50%
PEF _{0.75}	peak expiratory flow in 0.75 second		cut point diameter is the diameter at which the sampler collects 50%
PEFR	peak expiratory flow rate		of the particles and rejects 50% of the particles) and a penetration
PEFT	time to peak flow		curve as measured by a reference
PEG-CAT	polyethylene glycol-catalase		method based on Appendix L of 40 CFR Part 50 and designated in
PEG-SOD	polyethylene glycol-superoxide dismutase		accordance with 40 CFR Part 53, by an equivalent method
PEM(s)	personal exposure monitor(s)		designated in accordance with 40
Penh	enhanced pause		CFR Part 53, or by an approved
PEPc	phosphoenolpyruvate carboxylase		regional method designated in accordance with Appendix C of 40
PFD	photosynthetic flux density		CFR Part 58.
PFT	pulmonary function test		
pg	picogram(s)		
PG	prostaglandin (e.g., PGE2 ,PGF2); phosphatidylglycerol		
6PGD	6-phosphogluconate dehydrogenase		
PGE2	prostaglandin E2		
PGF2α	prostaglandin F2-alpha		
PGHS-2	prostaglandin endoperoxide G/H synthase 2		
PGP	protein gene product (e.g., PGP9.5)		
PGSM	Plant Growth Stress Model		

PM ₁₀	In general terms, particulate matter with an aerodynamic diameter less than or equal to a nominal 10 µm;	PNN50	proportion of interval differences of successive normal-beat intervals greater than 50 ms in EKG
	a measurement of thoracic	PO ₂	partial pressure of oxygen
	particles (i.e., that subset of	POC	particulate organic carbon
	inhalable particles thought small enough to penetrate beyond the	POD	peroxidase
	larynx into the thoracic region of the respiratory tract) in regulatory	polyADPR	poly(adenosinediphosphate- ribose)
	terms, particles with an upper 50% cut-point of 10 \pm 0.5 μ m aerodynamic diameter (the 50%	POMS	Portable Ozone Monitoring Systems
	cut point diameter is the diameter	ppb	parts per billion
	at which the sampler collects 50% of the particles and rejects 50% of	ppb-h	parts per billion per hour
	the particles) and a penetration	ppbv	parts per billion by volume
	curve as measured by a reference	pphm	parts per hundred million
	method based on Appendix J of 40	ppm	parts per million
	CFR Part 50 and designated in accordance with 40 CFR Part 53 or by an equivalent method	ppm-h	parts per million hours; weighted concentration values based on hourly concentrations: usually
PM _{10-2.5}	designated in accordance with 40 CFR Part 53. In general terms, particulate matter		summed over a certain number of hours, day(s), months, and/or season.
	with an aerodynamic diameter less	nnmu	parts per million by volume
	than or equal to a nominal 10 μm	ppmv	
	and greater than a nominal 2.5 µm; a measurement of thoracic coarse particulate matter or the	PPN	peroxypropionyl nitrate; peroxypropionic nitric anhydride
	coarse fraction of PM_{10} in	PPPs	power plant plumes
	regulatory terms, particles with an	ppt	parts per trillion
	upper 50% cut-point of 10 µm aerodynamic diameter and a lower	pptv	parts per trillion by volume
	50% cut-point of 2.5 μ m	PQH2	plastoquinone
	aerodynamic diameter (the 50%	PR	pathogenesis-related (protein)
	cut point diameter is the diameter at which the sampler collects 50%	PR-1	promoter region 1
	of the particles and rejects 50% of	PRB	policy-relevant background
	the particles) as measured by a	preproET-1	pre-protein form of ET-1 mRNA
	reference method based on Appendix O of 40 CFR Part 50 and	PRYL	predicted relative yield (biomass) loss
	designated in accordance with 40 CFR Part 53 or by an equivalent	PS	penalized spline
	method designated in accordance	PS	paradoxical sleep
PM _{10C}	with 40 CFR Part 53. The PM _{10-2.5} concentration of PM _{10-2.5} measured by the 40 CFR	PS II	Photosystem II: enzyme that uses light to obtain electrons from water (for photosynthesis).
	Part 50 Appendix O reference	PSA	picryl sulfonic acid
	method which consists of currently	PSC	polar stratospheric clouds
	operated, co-located low-volume (16.7 Lpm) PM ₁₀ and PM _{2.5}	PTB	preterm birth
p38MAPK	reference method samplers. p38 mitogen-activated protein	PTR-MS	proton-transfer-reaction mass spectroscopy
	kinase(s)	PU, PUL	pulmonary
PM-CAMx	Comprehensive Air Quality Model	PUFA(s)	polyunsaturated fatty acid(s)
	with extensions and with	PV	potential vorticity
PMN(s)	particulate matter chemistry polymorphonuclear leukocyte(s)	PVCD	peripheral vascular and cerebrovascular disease
PMT	photomultiplier tube	PVD	peripheral vascular disease
PND	post natal day	PVOCs	photochemical volatile organic
pNEM	probabilistic National Exposure Model	PWM	compounds pokeweed mitogen
PnET	Photosynthetic EvapoTranspiration	PWTES	(left ventricular) posterior wall
PNN	model proportion of interval differences of		thickness at end systole
	successive normal-beat intervals	Pxase	
	in EKG	QA	Quality Assurance
		QC	quality control

QCE		Rn	nanal registeries
	quasi continuous exercise	RNA	nasal resistance ribonucleic acid
qNP	non-photochemical quenching		
q _{NP}	non-photochemical quenching	RO₂	organic peroxyl; organic peroxy
qP	photochemical quenching	ROG	reactive organic gases
QRS	A complex of three distinct electrocardiogram waves which	ROI	reactive oxygen intermediate/superoxide anion
	represent the beginning of ventricular contraction	RONO ₂	organic nitrate
QT	interval measure of the time	ROOH	organic peroxides
	interval between the start of the Q	ROONO ₂ , RO ₂ NO ₂	peroxy nitrate
	wave and the end of the T wave in	ROS	reactive oxygen species
	the heart's electrical cycle	RPD	relative percent difference
QTc	corrected QT interval	RR	normal-to-normal (NN or RR) time
r	Pearson correlation coefficient		interval between each QRS complex in the EKG; risk ratio;
R, r	correlation coefficient		relative risk; respiratory rate
r ²	correlation coefficient	RRMS	relatively remote monitoring sites
R ²	multiple regression correlation coefficient	RT	respiratory tract
R^2 , r^2	coefficient of determination	RT	transepithelial resistance
RACM	Regional Atmospheric Chemistry	RTLF	respiratory tract lining fluid
-	Mechanism	RuBisCO; Rubisco	ribulose-1,5-bisphosphate carboxylase/oxygenase
RADM	Regional Acid Deposition Model	RuBP	ribulose bisphosphate
rALP	recombinant antileukoprotease		sigma, standard deviation
RAMS	Regional Atmospheric Modeling	σ	-
RANTES	System regulated upon activation, normal	σg	sigma-g; (geometric standard deviation)
	T-cell expressed and secreted (cells)	S	second
Raw		S	Short; smoker; sulfur; South
RB	airway resistance respiratory bronchiole	S.C.	subcutaneous (route)
RBC(s)		SA	salicylic acid
rbcL	red blood cell(s); erythrocyte(s)	SAB	Science Advisory Board
rbcS	Rubisco large subunit Rubisco small subunit	SAC	Staphylococcus aureus Cowan 1 strain
R'CO acyl		SAG21	
$R'C(O) - O_2$	acyl carrier protein	SAG21	Senescence
rcd1	acyl peroxy Arabidopsis mutant radical	S-allele	Systems Applications International short-allele
ICUT	induced cell death	SAMD	S-adenosyl methionine
RCD3	rod-cone dysplasia 3	SAMD	decarboxylase
RCP	Representative Concentration	SaO ₂	oxygen saturation of arterial blood
	Pathways	SAPALDIA	Study of Air Pollution and Lung
RDBMS	Relational Database Management Systems	SAPRC	Diseases in Adults Stratospheric Processes and their
Re	Reynolds number	0AI NO	Role in Climate; Statewide Air
REHEX	Regional Human Exposure Model		Pollution Research Center,
RER	rough endoplasmic reticulum; Respiratory exchange ratio	SAR	University of California, Riverside systemic acquired resistance
RF	radiative forcing	SAROAD	Storage and Retrieval of
RGR	relative growth rate		Aerometric Data (U.S. EPA
RH	relative humidity		centralized database; superseded by Aerometric Information
RIOPA	Relationship of Indoor, Outdoor,		Retrieval System [AIRS])
	and Personal Air (study)	SAWgrp	small airway function group
RL	total pulmonary resistance	SBNF	San Bernardino National Forest,
RLKs	receptor-like/Pelle kinase group		California
RMNP	Rocky Mountain National Park,	SBP	systolic blood pressure
	Colorado	SBUV	Solar Backscatter Ultraviolet
RMR	resting metabolic rate	80	Spectrometer
rMSSD	root mean squared differences	SC	stratum corneum
	between adjacent normal-to- normal heartbeat intervals	Sc	scandium

SCAQS	Southern California Air Quality	SOS	Southern Oxidant Study
SCAQS	Study	SO3 SO _x	sulfur oxides
SCE(s)	sister chromatid exchange(s)	SoyFACE	Soybean Free Air gas
SD	standard deviation; Sprague-Dawley rat	00)0	Concentration Enrichment (Facility)
SDNN	standard deviation normal-to- normal (NN or RR) time interval	SP	surfactant protein (e.g., SPA, SPD); substance P
	between each QRS complex in the EKG	SP-A	surfactant protein-A
SE	standard error	SPF	specific pathogen free
SEBAS	Social Environment and	SPMs	special purpose monitors
	Biomarkers of Aging Study second	SP-NK	substance P – neurokinin receptor complex
sec Sess.	session	sRaw,	specific airway resistance
SEM	simultaneously extracted metal;	SRBC	sheep red blood cell
SEM	standard error of the mean; scanning electron microscopy	SRES	Special Report on Emissions Scenarios
SENP	Sequoia National Park, California	SRM	standard reference method
SES	socioeconomic status	SRP	standard reference photometers
SF	San Francisco Bay Area	SSCP	single-strand conformation
SF6	sulfur hexafluoride (tracer gas)		polymorphism
SGA	small for gestational age	129S1/SvlmJ	mouse strain
sRaw	specific airway conductance	STE	stratosphere-troposphere exchange
SH	Shenandoah National Park site	STEP	Stratospheric-
SHEDS	Stochastic Human Exposure and		Tropospheric-exchange Project
CHEN	Dose Simulation	STN	speciation trends network
SHEN sICAM-1	Shenandoah National Park	sTNFR1	soluble tumor necrosis factor
	soluble intercellular adhesion molecule	STP	receptor 1 standard temperature and
SIDS	sudden infant death syndrome	0700	pressure
SIGMOID	sigmoid weighted summed concentration	STPD	standard temperature and pressure, dry
SINIC	Simple Nitrogen Cycle model	STRF	Spatio-Temporal Random Field
SIP	State Implementation Plan	and a solution	(theory)
SIPK	salicylic acid (SA) induced protein kinase	subscript i subscript o	Index of indoor microenvironments Index of outdoor
SK	shikimate kinase		microenvironments
SLA	specific leaf area	subscript o,i	Index of outdoor microenvironments adjacent to a
SLAC1	(protein) slow anion channel associated 1	0	given indoor microenvironment i
SLAMS	State and Local Air Monitoring Stations	SUM00	sum of all hourly average concentrations
SM	smooth muscle	SUM06	seasonal sum of all hourly average
SMD	soil moisture deficit	CUM07	concentrations ≥ 0.06 ppm
SME	soybean oil methyl ester	SUM07	seasonal sum of all hourly average concentrations ≥ 0.07 ppm
SMNP	Great Smoky Mountain National Park (North Carolina and	SUM08	seasonal sum of all hourly average concentrations ≥ 0.08 ppm
SMOKE	Tennessee) Spare-Matrix Operator Kernel	SURE	Sulfate Regional Experiment
SMORE	Emissions	SVE	Program supraventricular ectopy
S _N	normalized slope of the alveolar	S-W	square-wave
	plateau	SWS	slow wave sleep
SNAAQS	Secondary National Ambient Air	SZA	solar zenith angle
SNP(s)	Quality Standards single-nucleotide polymorphism	T	tau, photochemical lifetime;
SO ₂	sulfur dioxide		atmospheric lifetime
SO4 ²⁻	sulfate	t 	t-test statistical value; t statistic
SOC	soil organic carbon	Т	time; duration of exposure
SOD	superoxide dismutase		

T-cell(s)	T lymphocyte(s), thymus- dependent lymphocytes	TOMS	Total Ozone Mapping/Monitoring Satellite; total ozone mapping
T1	first trimester	TOPOE	spectrometer
T2	second trimester	TOPSE	Tropospheric Ozone Production About the Spring Equinox
T ₃	triiodothyronine	tPA	tissue plasminogen activator
T3	third trimester	TPLIF	two-photon laser-induced
T ₄	thyroxine		fluorescence
TAR	IPCC Third Assessment Report	TRAMP	TexAQS-II Radical and Aerosol
TAR WGI	IPCC Third Assessment Report of Working Group I	705000	Measurement Project
ТВ	tracheobronchial; terminal	TREGRO	Tree Growth Model
	bronchioles; tuberculosis	TRIFFID	Top-down Representation of Interactive Foliage and Flora
ТВА	thiobarbituric acid		Including Dynamics
TBARS	thiobarbituric acid reactive	TRIM	Total Risk Integrated Methodology
	substances		(model)
TC	total carbon	TRIM.Expo	Total Risk Integrated Methodology Exposure Event (model)
^{99m} Tc	Technetium-99m	TRP	transient receptor potential (ion
T-cells	T-lymphocytes, Thymus-derived lymphocytes		channel[s], ex., TRP-A1, TRP-V1, TRP-M8)
99mTc-DTPA	99mTc- diethylenetriaminepentaacetic acid	TSH	thyroid stimulating hormone
Тсо	core temperature	TSP	total suspended particles
TDLAS	Tunable Diode Laser Absorption Spectrometer	TTFMS	two-tone frequency-modulated spectroscopy
Те	expiratory time	TWA	time-weighted average
TEM	transmission electron microscopy;	ТХ	thromboxane (e.g., TXB ₂)
	Terrestrial Ecosystem Model	TXB ₂	thromboxane B2
TES	Tropospheric Emission	UA	uric acid; urate
	Spectrometer	UAM	Urban Airshed Model
TexAQS	Texas Air Quality Field Study	UCL	upper 95th% confidence limit
Tg	teragram(s)	UDGT	UDP -galactose-1,2,-diacylglycerol
TGF	transforming growth factor		galactosyltransferase
TGF β	transforming growth factor beta	UDP	uridine diphosphate
Th	T helper cell type	U.K.	United Kingdom
Th2	T helper cell type 2	UNECE	United Nations Economic Commission for Europe
THC	Total hydrocarbon content	UNEP	United Nations Environmental
tHcy T:	total homocysteine	UNE!	Programme
Ti Ti	inspiratory time	UNFCCC	United Nations Framework
TIA	titanium transient ischemic attack		Convention on Climate Change
TIMP-2	tissue inhibitor of matrix	U-O	epioxides formed from uric acid
111111 -2	metalloprotease-2		peroxides formed from uric acid
TiO ₂	titanium dioxide		ozonides formed from uric acid
TLC	total lung capacity	URI	upper respiratory infection
TLNISE	two-level normal independent	URS	upper respiratory symptoms
	sampling estimation	URT	upper respiratory tract; upper airways
Tlr	Toll-like receptor gene	U.S.	United States (of America)
TLR	Toll-like receptor protein (ex.,	USC; U.S.C.	U.S. Code
ТМРО	TLR2, TLR4) tetramethylphrrolise 1-oxide	USDA	U.S. Department of Agriculture
TNC	total nonstructural carbohydrate	USFS	U.S. Forest Service
TNF	tumor necrosis factor (e.g., TNF- α)	USGCRP	U.S. Global Change Research
TNF-308	tumor necrosis factor genotype		Program
TNF-α	tumor necrosis factor alpha	USGS	U.S. Geological Survey
TNFR	tumor necrosis factor receptor	UV	ultraviolet radiation
		UV-A	ultraviolet radiation at wavelengths of 320 to 400 nm

UV-B	ultraviolet radiation at wavelengths	WED	(U.S. EPA NHEERL) Western
UV-C	of 280 to 320 nm ultraviolet radiation at wavelengths	WF, WFM	Ecology Division White Face Mountain site
	of 200 to 280 nm	WHI	Women's Health Initiative
UV-DIAL	Ultraviolet Differential Absorption	WHO	World Health Organization
	Lidar	W/m^2 , $W m^{-2}$	watts per square meter
V	vanadium	WMO	World Meteorological Organization
V, mV, µV	volt, millivolt, microvolt	WMO/UNEP	World Meteorological
VA	alveolar ventilation		Organization/United Nations
Val	valine		Environment Program
VC	vital capacity	WRF	Weather Research and Forecasting model
VCAM	vascular cell adhesion molecule	Ws	Wassilewskija Arabidopsis ecotype
V _d	deposition rate, deposition velocity (cm/sec)	WS	wood smoke
V _D	volume of the anatomic or	WT	wild type; White Top Mountain site
v D	physiological dead space	wt %	percent by weight
Ϋ́ _E	ventilation rate; minute ventilation;	WUS	western U.S.
. F	ventilatory volume	w/v	
VEGF	vascular endothelial growth factor	w/v Y	weight per volume
॑V _E max	maximum minute ventilation		three parameter Weibull model
Vmax	maximum velocity	yr Z	year
Vmax _{25%}	maximum expiratory flow at 25%		Airway generation
	of the vital capacity	ZAPS	Zonal Air Pollution System
Vmax _{50%}	maximum expiratory flow at 50% of the vital capacity	ZELIG	a forest succession simulation model
Vmax _{75%}	maximum expiratory flow at 75% of the vital capacity	Zn	zinc
VMD	volume median diameter		
Vn	nasal volume		
VO ₂	oxygen consumption		
VO₂max	maximum volume per time, of oxygen (maximal oxygen consumption, maximal oxygen uptake or aerobic capacity)		
VOC(s)	volatile organic compound(s)		
VP	volumetric penetration		
VP _{50%}	volume at which 50% of an inhaled bolus is absorbed		
VPD	vapor pressure deficit; Vehicles per day; Ventricular premature depolarization		
VT	tidal volume		
VTB	terminal bronchiole region volume		
VTmax	maximum tidal volume		
VUA	volume of the upper airways		
vWF	von Willebrand factor		
W	width; wilderness; week(s)		
W126	cumulative integrated exposure index with a sigmoidal weighting function		
W95	cumulative integrated exposure index with a sigmoidal weighting function		
WBC	white blood cell		
WBGT	wet bulb globe temperature		
WC	sigmoidal weighting of hourly O_3 concentration		
WOD			

warm conveyor belt

WCB

PREAMBLE

Process of ISA Development

1	This preamble outlines the general process for developing an Integrated Science
2	Assessment (ISA) including the framework for evaluating weight of evidence and
3	drawing scientific conclusions and causal judgments. The ISA provides a concise review,
4	synthesis, and evaluation of the most policy-relevant science to serve as a scientific
5	foundation for the review of the National Ambient Air Quality Standards (NAAQS). The
6	general process for NAAQS reviews is described at
7	http://www.epa.gov/ttn/naaqs/review.html. Figure I depicts the general NAAQS review
8	process and information for individual NAAQS reviews is available at
9	www.epa.gov/ttn/naaqs. This preamble is a general discussion of the basic steps and
10	criteria used in developing an ISA; for each ISA, specific details and considerations are
11	included in the introductory section for that assessment.
12	The fundamental process for developing an ISA includes:
13	 literature searches;
14	 study selection;
15	 evaluation and integration of the evidence; and
16	 development of scientific conclusions and causal judgments.
17	An initial step in this process is publication of a call for information in the Federal
18	Register that invites the public to provide information relevant to the assessment, such as
19	new publications on health or welfare ¹ effects of the pollutant, or from atmospheric and
20	exposure sciences fields. EPA maintains an ongoing literature search process for
21	identification of relevant scientific studies published since the last review of the NAAQS.
22	Search strategies are designed for pollutants and scientific disciplines and iteratively
23	modified to optimize identification of pertinent publications. Papers are identified for
24	inclusion in several additional ways: specialized searches on specific topics; independent
25	review of tables of contents for journals in which relevant papers may be published;
26	independent identification of relevant literature by expert scientists; review of citations in
27	previous assessments and identification by the public and CASAC during the external
28	review process. This literature search and study selection process is depicted in Figure II.
29	Publications considered for inclusion in the ISA are added to the Health and

¹ Welfare effects as defined in Clean Air Act section 302(h) [42 U.S.C. 7602(h)] include, but are not limited to, "effects on soils, water, crops, vegetation, man-made materials, animals, wildlife, weather, visibility and climate, damage to and deterioration of property, and hazards to transportation, as well as effects on economic values and on personal comfort and well-being."

1	Environmental Research Online (HERO) database developed by EPA
2	(<u>http://hero.epa.gov/</u>); the references in the ISA include a hyperlink to the database.
3	Studies that have undergone scientific peer review and have been published or accepted
4	for publication and reports that have undergone review are considered for inclusion in the
5	ISA. Analyses conducted by EPA using publicly available data are also considered for
6	inclusion in the ISA. All relevant epidemiologic, controlled human exposure,
7	toxicological, and ecological and welfare effects studies published since the last review
8	are considered, including those related to exposure-response relationships, mode(s) of
9	action (MOA), and potentially at-risk populations and lifestages. Studies on atmospheric
10	chemistry, environmental fate and transport, dosimetry, toxicokinetics and exposure are
11	also considered for inclusion in the document, as well as analyses of air quality and
12	emissions data. References that were considered for inclusion in a specific ISA can be
13	found using the HERO website (<u>http://hero.epa.gov</u>).

National Ambient Air Quality Standard Review Process

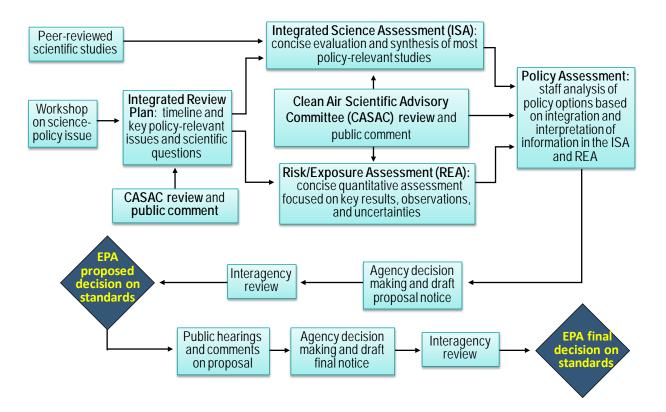
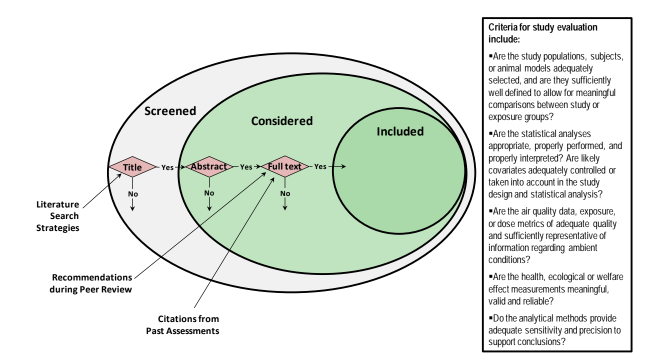
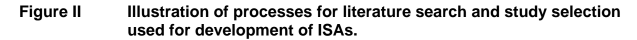


Figure I Illustration of the key steps in the process of the review of National Ambient Air Quality Standards.





1	Each ISA builds upon the conclusions of previous assessments for the pollutant under
2	review. EPA focuses on peer reviewed literature published following the completion of
3	the previous review and on any new interpretations of previous literature, integrating the
4	results of recent scientific studies with previous findings. Important earlier studies may
5	be discussed in detail to reinforce key concepts and conclusions or for reinterpretation in
6	light of newer data. Earlier studies also are the primary focus in some areas of the
7	document where research efforts have subsided, or if these earlier studies remain the
8	definitive works available in the literature.
9	Selection of studies for inclusion in the ISA is based on the general scientific quality of
10	the study, and consideration of the extent to which the study is informative and policy-
11	relevant. Policy relevant and informative studies include those that provide a basis for or
12	describe the relationship between the criteria pollutant and effects, including studies that
13	offer innovation in method or design and studies that reduce uncertainty on critical issues,

such as analyses of confounding or effect modification by copollutants or other variables,
analyses of concentration-response or dose-response relationships, or analyses related to
time between exposure and response. Emphasis is placed on studies that examine effects
associated with pollutant concentrations relevant to current population and ecosystem

1	exposures, and particularly those pertaining to concentrations currently found in ambient
2	air. Other studies are included if they contain unique data, such as a previously
3	unreported effect or MOA for an observed effect, or examine multiple concentrations to
4	elucidate exposure-response relationships. In general, in assessing the scientific quality
5	and relevance of health and welfare effects studies, the following considerations have
6	been taken into account when selecting studies for inclusion in the ISA.
7	 Are the study populations, subjects, or animal models adequately selected, and
8	are they sufficiently well defined to allow for meaningful comparisons
9	between study or exposure groups?
10	 Are the statistical analyses appropriate, properly performed, and properly
11	interpreted? Are likely covariates adequately controlled or taken into account
12	in the study design and statistical analysis?
13	Are the air quality data, exposure, or dose metrics of adequate quality and
14	sufficiently representative of information regarding ambient conditions?
15	 Are the health, ecological or welfare effect measurements meaningful, valid
16	and reliable?
17	 Do the analytical methods provide adequate sensitivity and precision to
18	support conclusions?
10	
 19 20 21 22 23 24 25 26 	Considerations specific to particular disciplines include the following. In selecting epidemiologic studies, EPA considers whether a given study: (1) presents information on associations with short- or long-term pollutant exposures at or near conditions relevant to ambient exposures; (2) addresses potential confounding by other pollutants; (3) assesses potential effect modifiers; (4) evaluates health endpoints and populations not previously extensively researched; and (5) evaluates important methodological issues related to interpretation of the health evidence (e.g., lag or time period between exposure and effects, model specifications, thresholds, mortality displacement).
20	epidemiologic studies, EPA considers whether a given study: (1) presents information on
21	associations with short- or long-term pollutant exposures at or near conditions relevant to
22	ambient exposures; (2) addresses potential confounding by other pollutants; (3) assesses
23	potential effect modifiers; (4) evaluates health endpoints and populations not previously
24	extensively researched; and (5) evaluates important methodological issues related to
25	interpretation of the health evidence (e.g., lag or time period between exposure and

- 1potentially at-risk populations and lifestages such as people with asthma or2cardiovascular diseases, children or older adults; (2) address issues such as concentration-3response or time-course of responses; and (3) have sufficient statistical power to assess4findings.
- 5 Review of the animal toxicological evidence focuses on studies that approximate 6 expected human dose conditions, which vary depending on the dosimetry, toxicokinetics 7 and biological sensitivity of the particular laboratory animal species or strains studied. 8 Emphasis is placed on studies that: (1) investigate animal models of disease that can 9 provide information on populations potentially at increased risk of effects; (2) address 10 issues such as concentration-response or time-course of responses; and (3) have sufficient statistical power to assess findings. Due to resource constraints on exposure duration and 11 12 numbers of animals tested, animal studies typically utilize high-concentration exposures 13 to acquire data relating to mechanisms and assure a measurable response. Emphasis is 14 placed on studies using doses or concentrations generally within 1-2 orders of magnitude 15 of current levels. Studies with higher concentration exposures or doses are considered to 16 the extent that they provide useful information to inform understanding of interspecies 17 differences and potential differences between healthy and potentially at-risk human 18 populations. Results from in vitro studies may also be included if they provide 19 mechanistic insight or further support for results demonstrated in vivo.
- 20 These criteria provide benchmarks for evaluating various studies and for focusing on the 21 policy-relevant studies in assessing the body of health, ecological and welfare effects 22 evidence. As stated initially, the intent of the ISA is to provide a concise review, 23 synthesis, and evaluation of the most policy-relevant science to serve as a scientific 24 foundation for the review of the NAAQS, not extensive summaries of all health, 25 ecological and welfare effects studies for a pollutant. Of most relevance for inclusion of 26 studies is whether they provide useful qualitative or quantitative information on 27 exposure-effect or exposure-response relationships for effects associated with pollutant 28 exposures at doses or concentrations relevant to ambient conditions that can inform 29 decisions on whether to retain or revise the standards.
- 30 In developing an ISA, EPA reviews and summarizes the evidence from: studies of 31 atmospheric sciences and exposure; the health effects evidence from toxicological, 32 controlled human exposure and epidemiologic studies; and ecological and welfare effects 33 evidence. In the process of developing the first draft ISA, EPA may convene a public 34 workshop in which EPA and non-EPA experts review the scientific content of 35 preliminary draft materials to ensure that the ISA is up to date and focused on the most 36 policy-relevant findings, and to assist EPA with integration of evidence within and across 37 disciplines. The general process for ISA development is illustrated in Figure III.

1	EPA integrates the evidence from across scientific disciplines or study types and
2	characterizes the weight of evidence for relationships between the pollutant and various
3	outcomes. The integration of evidence on health, and ecological or welfare effects,
4	involves collaboration between scientists from various disciplines. As an example, an
5	evaluation of health effects evidence would include the integration of the results from
6	epidemiologic, controlled human exposure, and toxicological studies, and application of
7	the causal framework (described below) to draw conclusions. Using the causal
8	framework described in the following section, EPA scientists consider aspects such as
9	strength, consistency, coherence, and biological plausibility of the evidence, and develop
10	causality determinations on the nature of the relationships. Causality determinations often
11	entail an iterative process of review and evaluation of the evidence. Two drafts of the ISA
12	are typically released for review by the CASAC and the public, and comments received
13	on the characterization of the science as well as the implementation of the causal
14	framework are carefully considered in revising and completing the final ISA.

Integrated Science Assessment Development Process

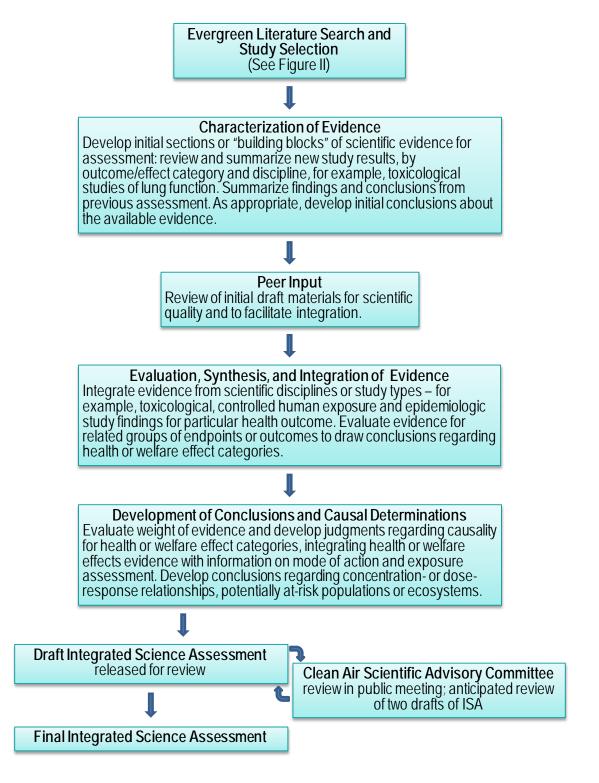


Figure III Characterization of the general process of ISA development.

EPA Framework for Causal Determination

1	EPA has developed a consistent and transparent basis to evaluate the causal nature of air
2	pollution-related health or welfare effects for use in developing ISAs. The framework
3	described below establishes uniform language concerning causality and brings more
4	specificity to the findings. This standardized language was drawn from sources across the
5	federal government and wider scientific community, especially the National Academy of
6	Sciences (NAS) Institute of Medicine (IOM) document, Improving the Presumptive
7	Disability Decision-Making Process for Veterans (2008), a comprehensive report on
8	evaluating causality. This framework:
9	 describes the kinds of scientific evidence used in establishing a general causal
10	relationship between exposure and health effects;
11	 characterizes the evidence necessary to reach a conclusion about the existence
12	of a causal relationship;
13	 identifies issues and approaches related to uncertainty; and
14	 provides a framework for classifying and characterizing the weight of
15	evidence in support of a general causal relationship.
16	Approaches to assessing the separate and combined lines of evidence
17	(e.g., epidemiologic, controlled human exposure, and animal toxicological studies) have
18	been formulated by a number of regulatory and science agencies, including the IOM of
19	the NAS (2008), International Agency for Research on Cancer (2006), U.S. EPA (2005),
20	and Centers for Disease Control and Prevention (2004). Causal inference criteria have
21	also been described for ecological effects evidence (U.S. EPA, 1998a; Fox, 1991). These
22	formalized approaches offer guidance for assessing causality. The frameworks are similar
23	in nature, although adapted to different purposes, and have proven effective in providing
24	a uniform structure and language for causal determinations.

Evaluating Evidence for Inferring Causation

25The 1964 Surgeon General's report defined "cause" as a "significant, effectual26relationship between an agent and an associated disorder or disease in the host" (HEW,271964); more generally, a cause is defined as an agent that brings about an effect or a28result. An association is the statistical relationship among variables; alone, however, it is29insufficient proof of a causal relationship between an exposure and a health outcome.30Unlike an association, a causal claim supports the creation of counterfactual claims, that

- is, a claim about what the world would have been like under different or changed
 circumstances (<u>Samet and Bodurow, 2008</u>).
- 3 Many of the health and environmental outcomes reported in these studies have complex 4 etiologies. Diseases such as asthma, coronary heart disease (CHD) or cancer are typically 5 initiated by multiple agents. Outcomes depend on a variety of factors, such as age, 6 genetic susceptibility, nutritional status, immune competence, and social factors (Samet 7 and Bodurow, 2008; Gee and Payne-Sturges, 2004). Effects on ecosystems are often also 8 multifactorial with a complex web of causation. Further, exposure to a combination of 9 agents could cause synergistic or antagonistic effects. Thus, the observed risk may 10 represent the net effect of many actions and counteractions.
- 11 In estimating the causal influence of an exposure on health or environmental effects, it is 12 recognized that scientific findings incorporate uncertainty. "Uncertainty" can be defined 13 as having limited knowledge to exactly describe an existing state or future outcome, 14 e.g., the lack of knowledge about the correct value for a specific measure or estimate. 15 Uncertainty analysis may be qualitative or quantitative in nature. In many cases, the 16 analysis is qualitative, and can include professional judgment or inferences based on 17 analogy with similar situations. Quantitative uncertainty analysis may include use of 18 simple measures (e.g., ranges) and analytical techniques. Quantitative uncertainty 19 analysis might progress to more complex measures and techniques, if needed for decision 20 support. Various approaches to evaluating uncertainty include classical statistical 21 methods, sensitivity analysis, or probabilistic uncertainty analysis, in order of increasing 22 complexity and data requirements. However, data may not be available for all aspects of 23 an assessment and those data that are available may be of questionable or unknown 24 quality. Ultimately, the assessment is based on a number of assumptions with varying 25 degrees of uncertainty. The ISA generally evaluates uncertainties qualitatively in 26 assessing the evidence from across studies; in some situations quantitative analysis 27 approaches, such as meta-regression, may be used.
- Publication bias is a source of uncertainty regarding the magnitude of health risk
 estimates. It is well understood that studies reporting non-null findings are more likely to
 be published than reports of null findings, and publication bias can also result in
 overestimation of effect estimate sizes (Ioannidis, 2008). For example, effect estimates
 from single-city epidemiologic studies have been found to be generally larger than those
 from multicity studies (Bell et al., 2005).

Consideration of Evidence from Scientific Disciplines

1 Moving from association to causation involves the elimination of alternative explanations 2 for the association. The ISA focuses on evaluation of the findings from the body of 3 evidence, drawing upon the results of all studies determined to meet the criteria described 4 previously. Causality determinations are based on the evaluation and synthesis of 5 evidence from across scientific disciplines. The relative importance of different types of 6 evidence varies by pollutant or assessment, as does the availability of different types of 7 evidence for causality determination. Three general types of studies inform consideration 8 of human health effects: controlled human exposure, epidemiologic and toxicological 9 studies. Evidence on ecological or welfare effects may be drawn from a variety of 10 experimental approaches (e.g., greenhouse, laboratory, field) and numerous disciplines 11 (e.g., community ecology, biogeochemistry and paleontological/historical 12 reconstructions).

13 Direct evidence of a relationship between pollutant exposures and human health effects 14 comes from controlled human exposure studies. Controlled human exposure studies 15 experimentally evaluate the health effects of administered exposures in human volunteers 16 under highly controlled laboratory conditions. Also referred to as human clinical studies, 17 these experiments allow investigators to expose subjects to known concentrations of air 18 pollutants under carefully regulated environmental conditions and activity levels. In some 19 instances, controlled human exposure studies can also be used to characterize 20 concentration-response relationships at pollutant concentrations relevant to ambient 21 conditions. Controlled human exposures are typically conducted using a randomized 22 crossover design, with subjects exposed both to the pollutant and a clean air control. In 23 this way, subjects serve as their own controls, effectively controlling for many potential 24 confounders. However, controlled human exposure studies are limited by a number of 25 factors, including small sample size and short exposure time. For example, exposure 26 patterns relevant to understanding real-world exposures, especially long-term exposures, 27 are generally not practical to replicate in a laboratory setting. In addition, although 28 subjects do serve as their own controls, personal exposure to pollutants in the hours and 29 days preceding the controlled exposures may vary significantly between and within 30 individuals. Finally, controlled human exposure studies require investigators to adhere to 31 stringent health criteria for subjects included in the study, and therefore the results often 32 cannot be generalized to an entire population. Although some controlled human exposure 33 studies have included health-compromised individuals such as those with respiratory or 34 cardiovascular disease, these individuals must also be relatively healthy and may not 35 represent the most sensitive individuals in the population. In addition, the study design is 36 limited to exposures and endpoints that are not expected to result in severe health 37 outcomes. Thus, not observing an effect in controlled human exposure studies does not

1necessarily mean that a causal relationship does not exist. While controlled human2exposure studies provide important information on the biological plausibility of3associations observed in epidemiologic studies, observed effects in these studies may4underestimate the response in certain populations.

- 5 Epidemiologic studies provide important information on the associations between health 6 effects and exposure of human populations to ambient air pollution. In epidemiologic or 7 observational studies of humans, the investigator generally does not control exposures or 8 intervene with the study population. Broadly, observational studies can describe 9 associations between exposures and effects. These studies fall into several categories: 10 e.g., cross-sectional, prospective cohort, panel and time-series studies. "Natural experiments" offer the opportunity to investigate changes in health related to a change in 11 12 exposure, such as closure of a pollution source.
- 13 In evaluating epidemiologic studies, consideration of many study design factors and 14 issues must be taken into account to properly inform their interpretation. One key 15 consideration is evaluation of the potential contribution of the pollutant to a health 16 outcome when it is a component of a complex air pollutant mixture. Reported effect 17 estimates in epidemiologic studies may reflect: independent effects on health outcomes; 18 effects of the pollutant acting as an indicator of a copollutant or a complex ambient air 19 pollution mixture; effects resulting from interactions between that pollutant and 20 copollutants.
- 21 In the evaluation of epidemiologic evidence, one important consideration is potential 22 confounding. Confounding is "... a confusion of effects. Specifically, the apparent effect 23 of the exposure of interest is distorted because the effect of an extraneous factor is 24 mistaken for or mixed with the actual exposure effect (which may be null)" (Rothman 25 and Greenland, 1998). One approach to remove spurious associations due to possible 26 confounders is to control for characteristics that may differ between exposed and 27 unexposed persons; this is frequently termed "adjustment." Scientific judgment is needed 28 to evaluate likely sources and extent of confounding, together with consideration of how 29 well the existing constellation of study designs, results, and analyses address this 30 potential threat to inferential validity. A confounder is associated with both the exposure 31 and the effect; for example, confounding can occur between correlated pollutants that are 32 associated with the same effect.
- Several statistical methods are available to detect and control for potential confounders,
 with none of them being completely satisfactory. Multivariable regression models
 constitute one tool for estimating the association between exposure and outcome after
 adjusting for characteristics of participants that might confound the results. The use of
 multipollutant regression models has been the prevailing approach for controlling

- 1 potential confounding by copollutants in air pollution health effects studies. Finding the 2 likely causal pollutant from multipollutant regression models is made difficult by the 3 possibility that one or more air pollutants may be acting as a surrogate for an unmeasured 4 or poorly measured pollutant or for a particular mixture of pollutants. In addition, more 5 than one pollutant may exert similar health effects, resulting in independently observed 6 associations for multiple pollutants. The number and degree of diversity of covariates, as 7 well as their relevance to the potential confounders, remain matters of scientific 8 judgment. Despite these limitations, the use of multipollutant models is still the 9 prevailing approach employed in most air pollution epidemiologic studies and provides 10 some insight into the potential for confounding or interaction among pollutants.
- 11Confidence that unmeasured confounders are not producing the findings is increased12when multiple studies are conducted in various settings using different subjects or13exposures, each of which might eliminate another source of confounding from14consideration. For example, multicity studies can provide insight on potential15confounding through the use of a consistent method to analyze data from across locations16with different levels of copollutants and other covariates . Intervention studies, because of17their quasi-experimental nature, can be particularly useful in characterizing causation.
- 18 Another important consideration in the evaluation of epidemiologic evidence is effect 19 modification, which occurs when the effect differs between subgroups or strata; for 20 example, effect estimates that vary by age group or potential risk factor. "Effect-measure 21 modification differs from confounding in several ways. The main difference is that, 22 whereas confounding is a bias that the investigator hopes to prevent or remove from the 23 effect estimate, effect-measure modification is a property of the effect under study ... In 24 epidemiologic analysis one tries to eliminate confounding but one tries to detect and 25 estimate effect-measure modification" (Rothman and Greenland, 1998). When a risk 26 factor is a confounder, it is the true cause of the association observed between the 27 exposure and the outcome; when a risk factor is an effect modifier, it changes the 28 magnitude of the association between the exposure and the outcome in stratified analyses. 29 For example, the presence of a preexisting disease or indicator of low socioeconomic 30 status may be an effect modifier in causing increased risk of effects related to air 31 pollution exposure. It is often possible to stratify the relationship between health outcome 32 and exposure by one or more of these potential effect modifiers. For variables that 33 modify the association, effect estimates in each stratum will be different from one another 34 and different from the overall estimate, indicating a different exposure-response 35 relationship may exist in populations represented by these variables.
- 36Exposure measurement error, which refers to the uncertainty associated with using37exposure metrics to represent the actual exposure of an individual or population, can be

1 an important contributor to variability in air pollution epidemiologic study results. 2 Exposure error can under- or over-estimate epidemiologic associations between ambient 3 pollutant concentrations and health outcomes by biasing effect estimates toward or away 4 from the null, and tends to widen confidence intervals around those estimates. There are 5 several components that contribute to exposure measurement error in air pollution 6 epidemiologic studies, including the difference between true and measured ambient 7 concentrations, the difference between average personal exposure to ambient pollutants 8 and ambient concentrations at central monitoring sites, and the use of average population 9 exposure rather than individual exposure estimates. Factors that could influence exposure 10 estimates include nonambient sources of exposure, topography of the natural and built 11 environment, meteorology, measurement errors, time-location-activity patterns and extent 12 to which ambient pollutants penetrate indoor environments. The importance of exposure 13 misclassification varies with study design and is dependent on the spatial and temporal 14 aspects of the design.

- 15 The third main type of health effects evidence, animal toxicological studies, provides 16 information on the pollutant's biological action under controlled and monitored exposure 17 circumstances. Taking into account physiological differences of the experimental species from humans, these studies inform characterization of health effects of concern, 18 19 exposure-response relationships and MOAs. Further, animal models can inform 20 determinations of at-risk populations. These studies evaluate the effects of exposures to a 21 variety of pollutants in a highly controlled laboratory setting and allow exploration of 22 toxicological pathways or mechanisms by which a pollutant may cause effects. 23 Understanding the biological mechanisms underlying various health outcomes can prove 24 crucial in establishing or negating causality. In the absence of human studies data, 25 extensive, well-conducted animal toxicological studies can support determinations of 26 causality, if the evidence base indicates that similar responses are expected in humans 27 under ambient exposure conditions.
- 28 Interpretations of animal toxicological studies are affected by limitations associated with 29 extrapolation between animal and human responses. The differences between humans 30 and other species have to be taken into consideration, including metabolism, hormonal 31 regulation, breathing pattern, and differences in lung structure and anatomy. Also, in spite 32 of a high degree of homology and the existence of a high percentage of orthologous 33 genes across humans and rodents (particularly mice), extrapolation of molecular 34 alterations at the gene level is complicated by species-specific differences in 35 transcriptional regulation. Given these differences, there are uncertainties associated with 36 quantitative extrapolations of observed pollutant-induced pathophysiological alterations 37 between laboratory animals and humans, as those alterations are under the control of 38 widely varying biochemical, endocrine, and neuronal factors.

1	For ecological effects assessment, both laboratory and field studies (including field
2	experiments and observational studies) can provide useful data for causality
3	determination. Because conditions can be controlled in laboratory studies, responses may
4	be less variable and smaller differences easier to detect. However, the control conditions
5	may limit the range of responses (e.g., animals may not be able to seek alternative food
6	sources), so they may not reflect responses that would occur in the natural environment.
7	In addition, larger-scale processes are difficult to reproduce in the laboratory.
8	Field observational studies measure biological changes in uncontrolled situations, and
9	describe an association between a disturbance and an ecological effect. Field data can
10	provide important information for assessments of multiple stressors or where site-specific
11	factors significantly influence exposure. They are also often useful for analyses of larger
12	geographic scales and higher levels of biological organization. However, because
13	conditions are not controlled, variability is expected to be higher and differences harder
14	to detect. Field surveys are most useful for linking stressors with effects when stressor
15	and effect levels are measured concurrently. The presence of confounding factors can
16	make it difficult to attribute observed effects to specific stressors.
17	Intermediate between laboratory and field are studies that use environmental media
18	collected from the field to examine response in the laboratory, and experiments that are
19	performed in the natural environment while controlling for some environmental
20	conditions (i.e., mesocosm studies). This type of study in manipulated natural
21	environments can be considered a hybrid between a field experiment and laboratory study
22	since some aspects are performed under controlled conditions but others are not. They
23	make it possible to observe community and/or ecosystem dynamics, and provide strong
24	evidence for causality when combined with findings of studies that have been made
25	under more controlled conditions.

Application of Framework for Causal Determination

26In its evaluation of the scientific evidence on health or welfare effects of criteria27pollutants, EPA determines the weight of evidence in support of causation and28characterizes the strength of any resulting causal classification. EPA also evaluates the29quantitative evidence and draws scientific conclusions, to the extent possible, regarding30the concentration-response relationships and the loads to ecosystems, exposure doses or31concentrations, duration and pattern of exposures at which effects are observed.

Table I	Aspects to aid in judging causality
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Consistency of the observed association	An inference of causality is strengthened when a pattern of elevated risks is observed across several independent studies. The reproducibility of findings constitutes one of the strongest arguments for causality. If there are discordant results among investigations, possible reasons such as differences in exposure, confounding factors, and the power of the study are considered.	
Coherence	An inference of causality from one line of evidence (e.g., epidemiologic, clinical or animal studies) may be strengthened by other lines of evidence that support a cause-and-effect interpretation of the association. Evidence on ecological or welfare effects may be drawn from a variety of experimental approaches (e.g., greenhouse, laboratory, and field) and subdisciplines of ecology (e.g., community ecology, biogeochemistry and paleontological/historical reconstructions). The coherence of evidence from various fields greatly adds to the strength of an inference of causality. In addition, there may be coherence in demonstrating effects across multiple study designs or related health endpoints within on scientific line of evidence.	
Biological plausibility.	An inference of causality tends to be strengthened by consistency with data from experimental studies or other sources demonstrating plausible biological mechanisms. A proposed mechanistic linking between an effect and exposure to the agent is an important source of support for causality, especially when data establishing the existence and functioning of those mechanistic links are available.	
Biological gradient (exposure-response relationship)	A well-characterized exposure-response relationship (e.g., increasing effects associated with greater exposure) strongly suggests cause and effect, especially when such relationships are also observed for duration of exposure (e.g., increasing effects observed following longer exposure times).	
Strength of the observed association	The finding of large, precise risks increases confidence that the association is not likely due to chance, bias, or other factors. However, it is noted that a small magnitude in an effect estimate may represent a substantial effect in a population.	
Experimental evidence	Strong evidence for causality can be provided through "natural experiments" when a change in exposure is found to result in a change in occurrence or frequency of health or welfare effects.	
Temporal relationship of the observed association		
Specificity of the observed association	Evidence linking a specific outcome to an exposure can provide a strong argument for causation. However, it must be recognized that rarely, if ever, does exposure to a pollutant invariably predict the occurrence of an outcome, and that a given outcome may have multiple causes.	
Analogy Structure activity relationships and information on the agent's structural analogs can provide in whether an association is causal. Similarly, information on mode of action for a chemical, as or many structural analogs, can inform decisions regarding likely causality.		

1	To aid judgment, various "aspects" ¹ of causality have been discussed by many
2	philosophers and scientists. The 1964 Surgeon General's report on tobacco smoking
3	discussed criteria for the evaluation of epidemiologic studies, focusing on consistency,
4	strength, specificity, temporal relationship, and coherence (HEW, 1964). Sir Austin
5	Bradford Hill (Hill, 1965) articulated aspects of causality in epidemiology and public
6	health that have been widely used (Samet and Bodurow, 2008; IARC, 2006; U.S. EPA,
7	2005; CDC, 2004). These aspects (Hill, 1965) have been modified (Table I) for use in
8	causal determinations specific to health and welfare effects for pollutant exposures (U.S.
9	EPA, 2009d). ² Although these aspects provide a framework for assessing the evidence,
10	they do not lend themselves to being considered in terms of simple formulas or fixed

¹ The "aspects" described by Sir Austin Bradford Hill (<u>Hill, 1965</u>) have become, in the subsequent literature, more commonly described as "criteria." The original term "aspects" is used here to avoid confusion with "criteria" as it is used, with different meaning, in the Clean Air Act.

² The Hill aspects were developed for interpretation of epidemiologic results. They have been modified here for use with a broader array of data, i.e., epidemiologic, controlled human exposure, ecological, and animal toxicological studies, as well as in vitro data, and to be more consistent with EPA's Guidelines for Carcinogen Risk Assessment.

1 rules of evidence leading to conclusions about causality (Hill, 1965). For example, one 2 cannot simply count the number of studies reporting statistically significant results or 3 statistically nonsignificant results and reach credible conclusions about the relative 4 weight of the evidence and the likelihood of causality. Rather, these aspects are taken into 5 account with the goal of producing an objective appraisal of the evidence, informed by 6 peer and public comment and advice, which includes weighing alternative views on 7 controversial issues. In addition, it is important to note that the aspects in Table I cannot 8 be used as a strict checklist, but rather to determine the weight of the evidence for 9 inferring causality. In particular, not meeting one or more of the principles does not automatically preclude a determination of causality [see discussion in (CDC, 2004)]. 10

Determination of Causality

11 In the ISA, EPA assesses the body of relevant literature, building upon evidence available 12 during previous NAAOS reviews, to draw conclusions on the causal relationships 13 between relevant pollutant exposures and health or environmental effects. ISAs use a 14 five-level hierarchy that classifies the weight of evidence for causation¹. In developing 15 this hierarchy, EPA has drawn on the work of previous evaluations, most prominently the 16 IOM's Improving the Presumptive Disability Decision-Making Process for Veterans 17 (Samet and Bodurow, 2008), EPA's Guidelines for Carcinogen Risk Assessment (U.S. 18 EPA, 2005), and the U.S. Surgeon General's smoking report (CDC, 2004). This weight 19 of evidence evaluation is based on various lines of evidence from across the health and 20 environmental effects disciplines. These separate judgments are integrated into a 21 qualitative statement about the overall weight of the evidence and causality. The five 22 descriptors for causal determination are described in Table II.

23 Determination of causality involves the evaluation of evidence for different types of 24 health, ecological or welfare effects associated with short- and long-term exposure 25 periods. In making determinations of causality, evidence is evaluated for major outcome 26 categories and then conclusions are drawn based upon the integration of evidence from 27 across disciplines and also across the spectrum of related endpoints. In making causal 28 judgments, the ISA focuses on major outcome categories (e.g., respiratory effects, 29 vegetation growth), by evaluating the coherence of evidence across a spectrum of related 30 endpoints (e.g., health effects ranging from inflammatory effects to respiratory mortality) 31 to draw conclusions regarding causality. In discussing the causal determination, EPA

¹ The Center for Disease Control (CDC) and IOM frameworks use a four-category hierarchy for the strength of the evidence. A five-level hierarchy is used here to be consistent with the EPA Guidelines for Carcinogen Risk Assessment and to provide a more nuanced set of categories.

- 1characterizes the evidence on which the judgment is based, including strength of2evidence for individual endpoints within the major outcome category.
- 3 In drawing judgments regarding causality for the criteria air pollutants, the ISA focuses 4 on evidence of effects in the range of relevant pollutant exposures or doses, and not on 5 determination of causality at any dose. Emphasis is placed on evidence of effects at doses 6 (e.g., blood lead concentration) or exposures (e.g., air concentrations) that are relevant to, 7 or somewhat above, those currently experienced by the population. The extent to which 8 studies of higher concentrations are considered varies by pollutant and major outcome 9 category, but generally includes those with doses or exposures in the range of one to two 10 orders of magnitude above current or ambient conditions. Studies that use higher doses or 11 exposures may also be considered to the extent that they provide useful information to 12 inform understanding of mode of action, interspecies differences, or factors that may 13 increase risk of effects for a population. Thus, a causality determination is based on 14 weight of evidence evaluation for health, ecological or welfare effects, focusing on the 15 evidence from exposures or doses generally ranging from current levels to one or two 16 orders of magnitude above current levels.
- 17In addition, EPA evaluates evidence relevant to understand the quantitative relationships18between pollutant exposures and health, ecological or welfare effects. This includes19evaluation of the form of concentration-response or dose-response relationships and, to20the extent possible, drawing conclusions on the levels at which effects are observed. The21ISA also draws scientific conclusions regarding important exposure conditions for effects22and populations that may be at greater risk for effects, as described in the following23section.

Table II Weight of evidence for causal determination

	Health Effects	Ecological and Welfare Effects
Causal relationship	Evidence is sufficient to conclude that there is a causal relationship with relevant pollutant exposures (i.e., doses or exposures generally within one to two orders of magnitude of current levels). That is, the pollutant has been shown to result in health effects in studies in which chance, bias, and confounding could be ruled out with reasonable confidence. For example: a) controlled human exposure studies that demonstrate consistent effects; or b) observational studies that cannot be explained by plausible alternatives or are supported by other lines of evidence (e.g., animal studies multiple high-quality studies	Evidence is sufficient to conclude that there is a causal relationship with relevant pollutant exposures i.e., doses or exposures generally within one to two orders of magnitude of current levels). That is, the pollutant has been shown to result in effects in studies in which chance, bias, and confounding could be ruled out with reasonable confidence. Controlled exposure studies (laboratory or small- to medium-scale field studies) provide the strongest evidence for causality, but the scope of inference may be limited. Generally, determination is based on multiple studies conducted by multiple research groups, and evidence that is considered sufficient to infer a causal relationship is usually obtained from the joint consideration of many lines of evidence that reinforce each other.
Likely to be a causal relationship	Evidence is sufficient to conclude that a causal relationship is likely to exist with relevant pollutant exposures, but important uncertainties remain. That is, the pollutant has been shown to result in health effects in studies in which chance and bias can be ruled out with reasonable confidence but potential issues remain. For example: a) observational studies show an association, but copollutant exposures are difficult to address and/or other lines of evidence (controlled human exposure, animal, or mode of action information) are limited or inconsistent; or b) animal toxicological evidence from multiple studies from different laboratories that demonstrate effects, but limited or no human data are available. Evidence generally includes multiple high-quality studies.	Evidence is sufficient to conclude that there is a likely causal association with relevant pollutant exposures. That is, an association has been observed between the pollutant and the outcome in studies in which chance, bias and confounding are minimized, but uncertainties remain. For example, field studies show a relationship, but suspected interacting factors cannot be controlled, and other lines of evidence are limited or inconsistent. Generally, determination is based on multiple studies in multiple research groups.
Suggestive of a causal relationship	Evidence is suggestive of a causal relationship with relevant pollutant exposures, but is limited. For example, (a) at least one high-quality epidemiologic study shows an association with a given health outcome but the results of other studies are inconsistent; or (b) a well-conducted toxicological study, such as those conducted in the National Toxicology Program (NTP), shows effects in animal species.	Evidence is suggestive of a causal relationship with relevant pollutant exposures, but chance, bias and confounding cannot be ruled out. For example, at least one high-quality study shows an effect, but the results of other studies are inconsistent.
Inadequate to infer a causal relationship	Evidence is inadequate to determine that a causal relationship exists with relevant pollutant exposures. The available studies are of insufficient quantity, quality, consistency, or statistical power to permit a conclusion regarding the presence or absence of an effect.	The available studies are of insufficient quality, consistency, or statistical power to permit a conclusion regarding the presence or absence of an effect.
Not likely to be a causal relationship	Evidence is suggestive of no causal relationship with relevant pollutant exposures. Several adequate studies, covering the full range of levels of exposure that human beings are known to encounter and considering at-risk populations, are mutually consistent in not showing an effect at any level of exposure.	Several adequate studies, examining relationships with relevant exposures, are consistent in failing to show an effect at any level of exposure.

Quantitative Relationships: Effects on Human Populations

1	Once a determination is made regarding the causal relationship between the pollutant and		
2	outcome category, important questions regarding quantitative relationships include:		
3	• What is the concentration-response, exposure-response, or dose-response		
4	relationship in the human population?		
5	What is the interrelationship between incidence and severity of effect?		
6 7	 What exposure conditions (dose or exposure, duration and pattern) are important? 		
8 9	 What populations and lifestages appear to be differentially affected (i.e., more at risk of experiencing effects)? 		
10	To address these questions, the entirety of quantitative evidence is evaluated to		
11	characterize pollutant concentrations and exposure durations at which effects were		
12	observed for exposed populations, including populations and lifestages potentially at		
13	increased risk. To accomplish this, evidence is considered from multiple and diverse		
14 15	types of studies, and a study or set of studies that best approximates the concentration-		
15 16	response relationships between health outcomes and the pollutant may be identified. Controlled human exposure studies provide the most direct and quantifiable exposure-		
10	response data on the human health effects of pollutant exposures. To the extent available,		
18	the ISA evaluates results from across epidemiologic studies that use various methods to		
19	characterize the form of relationships between the pollutant and health outcomes and		
20	draws conclusions on the shape of these relationships. Animal data may also inform		
21	evaluation of concentration-response relationships, particularly relative to MOAs and		
22	characteristics of at-risk populations.		
23	An important consideration in characterizing the public health impacts associated with		
24	exposure to a pollutant is whether the concentration-response relationship is linear across		
25	the range of concentrations or if nonlinear relationships exist along any part of this range.		
26	Of particular interest is the shape of the concentration-response curve at and below the		
27	level of the current standards. Various sources of variability and uncertainty, such as low		
28	data density in the lower concentration range, possible influence of exposure		
29	measurement error, and variability between individuals in susceptibility to air pollution		
30	health effects, tend to smooth and "linearize" the concentration-response function, and		
31	thus can obscure the existence of a threshold or nonlinear relationship. Since individual		
32	thresholds vary from person to person due to individual differences such as genetic level		
33	susceptibility or preexisting disease conditions (and even can vary from one time to		
34	another for a given person), it can be difficult to demonstrate that a threshold exists in a		
35	population study. These sources of variability and uncertainty may explain why the		

1	available human data at ambient concentrations for some environmental pollutants
2	(e.g., particulate matter [PM], O ₃ , lead [Pb], environmental tobacco smoke [ETS],
3	radiation) do not exhibit thresholds for cancer or noncancer health effects, even though
4	likely mechanisms include nonlinear processes for some key events. These attributes of
5	human population dose-response relationships have been extensively discussed in the
6	broader epidemiologic literature (Rothman and Greenland, 1998).
7	Finally, identification of the population groups or lifestages that may be at greater risk of
8	health effects from air pollutant exposures contributes to an understanding of the public
9	health impact of pollutant exposures. In the ISA, the term "at-risk population" is used to
10	encompass populations or lifestages that have a greater likelihood of experiencing health
11	effects related to exposure to an air pollutant due to a variety of factors; other terms used
12	in the literature include susceptible, vulnerable, and sensitive. These factors may be
13	intrinsic, such as genetic or developmental factors, race, gender, lifestage, or the presence
14	of preexisting diseases, or they may be extrinsic, such as socioeconomic status (SES),
15	activity pattern and exercise level, reduced access to health care, low educational
16	attainment, or increased pollutant exposures (e.g., near roadways). Epidemiologic studies
17	can help identify populations potentially at increased risk of effects by evaluating health
18	responses in the study population. Examples include testing for interactions or effect
19	modification by factors such as gender, age group, or health status. Experimental studies
20	using animal models of susceptibility or disease can also inform the extent to which
21	health risks are likely greater in specific population groups.

Quantitative Relationships: Effects on Ecosystems or Public Welfare

22 Key questions for understanding the quantitative relationships between exposure (or 23 concentration or deposition) to a pollutant and risk to ecosystems or the public welfare 24 include: 25 • What elements of the ecosystem (e.g., types, regions, taxonomic groups, 26 populations, functions, etc.) appear to be affected, or are more sensitive to 27 effects? Are there differences between locations or materials in welfare effects 28 responses, such as impaired visibility or materials damage? • Under what exposure conditions (amount deposited or concentration, duration 29 30 and pattern) are effects seen? 31 • What is the shape of the concentration-response or exposure-response 32 relationship? 33 Evaluations of causality generally consider the probability of quantitative changes in 34 ecological and welfare effects in response to exposure. A challenge to the quantification

of exposure-response relationships for ecological effects is the great regional and local
 spatial variability, as well as temporal variability, in ecosystems. Thus, exposure response relationships are often determined for a specific ecological system and scale,
 rather than at the national or even regional scale. Quantitative relationships therefore are
 available site by site and may differ greatly between ecosystems.

Concepts in Evaluating Adversity of Health Effects

6 In evaluating health evidence, a number of factors can be considered in delineating 7 between adverse and nonadverse health effects resulting from exposure to air pollution. 8 Some health outcomes, such as hospitalization for respiratory or cardiovascular diseases, 9 are clearly considered adverse. It is more difficult to determine the extent of change that 10 constitutes adversity in more subtle health measures. These include a wide variety of 11 responses, such as alterations in markers of inflammation or oxidative stress, changes in 12 pulmonary function or heart rate variability, or alterations in neurocognitive function 13 measures. The challenge is determining the magnitude of change in these measures when 14 there is no clear point at which a change become adverse; for example, what percentage 15 change in a lung function measure represents an adverse effect. What constitutes an 16 adverse health effect may vary between populations. Some changes that may not be 17 considered adverse in healthy individuals would be potentially adverse in more at-risk 18 individuals.

19 For example, the extent to which changes in lung function are adverse has been discussed 20 by the American Thoracic Society (ATS) in an official statement titled What Constitutes 21 an Adverse Health Effect of Air Pollution? (2000b). An air pollution-induced shift in the 22 population distribution of a given risk factor for a health outcome was viewed as adverse, 23 even though it may not increase the risk of any one individual to an unacceptable level. 24 For example, a population of asthmatics could have a distribution of lung function such 25 that no identifiable individual has a level associated with significant impairment. 26 Exposure to air pollution could shift the distribution such that no identifiable individual 27 experiences clinically relevant effects. This shift toward decreased lung function, 28 however, would be considered adverse because individuals within the population would 29 have diminished reserve function and therefore would be at increased risk to further environmental insult. The committee also observed that elevations of biomarkers, such as 30 31 cell number and types, cytokines and reactive oxygen species, may signal risk for ongoing 32 injury and clinical effects or may simply indicate transient responses that can provide 33 insights into mechanisms of injury, thus illustrating the lack of clear boundaries that 34 separate adverse from nonadverse effects.

1 The more subtle health outcomes may be connected mechanistically to health events that 2 are clearly adverse. For example, air pollution may affect markers of transient myocardial 3 ischemia such as ST-segment abnormalities and onset of exertional angina. These effects 4 may not be apparent to the individual, yet may still increase the risk of a number of 5 cardiac events, including myocardial infarction and sudden death. Thus, small changes in 6 physiological measures may not appear to be clearly adverse when considered alone, but 7 may be a part of a coherent and biologically plausible chain of related health outcomes 8 that range up to responses that are very clearly adverse, such as hospitalization or 9 mortality.

Concepts in Evaluating Adversity of Ecological Effects

- 10 Adversity of ecological effects can be understood in terms ranging in scale from the 11 cellular level to the individual organism and to the population, community, and 12 ecosystem levels. In the context of ecology, a population is a group of individuals of the 13 same species, and a community is an assemblage of populations of different species interacting with one another that inhabit an area. An ecosystem is the interactive system 14 15 formed from all living organisms and their abiotic (physical and chemical) environment 16 within a given area (IPCC, 2007a). The boundaries of what could be called an ecosystem 17 are somewhat arbitrary, depending on the focus of interest or study. Thus, the extent of an 18 ecosystem may range from very small spatial scales to, ultimately, the entire Earth 19 (IPCC, 2007a).
- 20 Effects on an individual organism are generally not considered to be adverse to public 21 welfare. However if effects occur to enough individuals within a population, then 22 communities and ecosystems may be disrupted. Changes to populations, communities 23 and ecosystems can in turn result in an alteration of ecosystem processes. Ecosystem 24 processes are defined as the metabolic functions of ecosystems including energy flow, 25 elemental cycling, and the production, consumption and decomposition of organic matter 26 (U.S. EPA, 2002). Growth, reproduction, and mortality are species-level endpoints that 27 can be clearly linked to community and ecosystem effects and are considered to be 28 adverse when negatively affected. Other endpoints such as changes in behavior and 29 physiological stress can decrease ecological fitness of an organism, but are harder to link 30 unequivocally to effects at the population, community, and ecosystem level. The degree 31 to which pollutant exposure is considered adverse may also depend on the location and its 32 intended use (i.e., city park, commercial, cropland). Support for consideration of 33 adversity beyond the species level by making explicit the linkages between stress-related 34 effects at the species and effects at the ecosystem level is found in A Framework for 35 Assessing and Reporting on Ecological Condition: an SAB report (U.S. EPA, 2002). 36 Additionally, the National Acid Precipitation Assessment Program (NAPAP) uses the

1following working definition of "adverse ecological effects" in the preparation of reports2to Congress mandated by the Clean Air Act: "any injury (i.e., loss of chemical or physical3quality or viability) to any ecological or ecosystem component, up to and including at the4regional level, over both long and short terms."

5 On a broader scale, ecosystem services may provide indicators for ecological impacts. 6 Ecosystem services are the benefits that people obtain from ecosystems (UNEP, 2003). 7 According to the Millennium Ecosystem Assessment, ecosystem services include: 8 "provisioning services such as food and water; regulating services such as regulation of 9 floods, drought, land degradation, and disease; supporting services such as soil formation 10 and nutrient cycling; and cultural services such as recreational, spiritual, religious and 11 other nonmaterial benefits." For example, a more subtle ecological effect of pollution 12 exposure may result in a clearly adverse impact on ecosystem services if it results in a 13 population decline in a species that is recreationally or culturally important.

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LEGISLATIVE AND HISTORICAL BACKGROUND

Legislative Requirements for the NAAQS Review

1	Two sections of the Clean Air Act (CAA) govern the establishment and revision of
2	the National Ambient Air Quality Standards (NAAQS). Section 108 (42 USC §7408)
3	directs the Administrator to identify and list certain air pollutants and then to issue air
4	quality criteria for those pollutants. The Administrator is to list those air pollutants
5	that in her "judgement; cause or contribute to air pollution which may reasonably be
6	anticipated to endanger public health or welfare;" "the presence of which in the
7	ambient air results from numerous or diverse mobile or stationary sources" and "for
8	which [the Administrator] plans to issue air quality criteria" (CAA, 1990a). Air
9	quality criteria are intended to "accurately reflect the latest scientific knowledge
10	useful in indicating the kind and extent of identifiable effects on public health or
11	welfare, which may be expected from the presence of [a] pollutant in ambient air"
12	[42 USC §7408(b)].
13	Section 109 (CAA, 1990b) directs the Administrator to propose and promulgate
14	"primary" and "secondary" NAAQS for pollutants for which air quality criteria have
15	been issued. Section 109(b)(1) defines a primary standard as one "the attainment and
16	maintenance of which in the judgment of the Administrator, based on such criteria
17	and allowing an adequate margin of safety, are requisite to protect the public
18	health." ¹ A secondary standard, as defined in section 109(b)(2), must "specify a level
19	of air quality the attainment and maintenance of which, in the judgment of the
20	Administrator, based on such criteria, is required to protect the public welfare from
21	any known or anticipated adverse effects associated with the presence of [the]
22	pollutant in the ambient air." ²
23	The requirement that primary standards include an adequate margin of safety was
24	intended to address uncertainties associated with inconclusive scientific and technical
25	information available at the time of standard setting. It was also intended to provide a
26	reasonable degree of protection against hazards that research has not yet identified.
27	See Lead Industries Association v. EPA, 647 F.2d 1130, 1154 (D.C. Cir 1980), cert.
28	denied, 449 U.S. 1042 (1980); American Petroleum Institute v. Costle, 665 F.2d
29	1176, 1186 (D.C. Cir. (1981), cert. denied, 455 U.S. 1034 (1982). Both kinds of

¹ The legislative history of section 109 indicates that a primary standard is to be set at "the maximum permissible ambient air level . . . which will protect the health of any [sensitive] group of the population," and that for this purpose "reference should be made to a representative sample of persons comprising the sensitive group rather than to a single person in such a group" [S. Rep. No. 91-1196, 91st Cong., 2d Sess. 10 (1970)].

² Welfare effects as defined in section 302(h) include, but are not limited to, "effects on soils, water, crops, vegetation, man-made materials, animals, wildlife, weather, visibility and climate, damage to and deterioration of property, and hazards to transportation, as well as effects on economic values and on personal comfort and well-being" (<u>CAA, 2005</u>).

1	uncertainties are components of the risk associated with pollution at levels below
2	those at which human health effects can be said to occur with reasonable scientific
3	certainty. Thus, in selecting primary standards that include an adequate margin of
4	safety, the Administrator is seeking not only to prevent pollution levels that have
5	been demonstrated to be harmful but also to prevent lower pollutant levels that may
6	pose an unacceptable risk of harm, even if the risk is not precisely identified as to
0 7	nature or degree. The CAA does not require the Administrator to establish a primary
8	NAAQS at a zero-risk level or at background concentration levels, see Lead
8 9	Industries v. EPA, 647 F.2d at 1156 n.51, but rather at a level that reduces risk
10	
10	sufficiently so as to protect public health with an adequate margin of safety.
11	In addressing the requirement for a margin of safety, EPA considers such factors as
12	the nature and severity of the health effects involved, the size of the sensitive
13	population(s) at risk, and the kind and degree of the uncertainties that must be
14	addressed. The selection of any particular approach to providing an adequate margin
15	of safety is a policy choice left specifically to the Administrator's judgment. See
16	Lead Industries Association v. EPA, supra, 647 F.2d at 1161-1162; Whitman v.
17	American Trucking Associations, 531 U.S. 457, 495 (2001).
18	In setting standards that are "requisite" to protect public health and welfare, as
19	provided in Section 109(b), EPA's task is to establish standards that are neither more
20	nor less stringent than necessary for these purposes. In so doing, EPA may not
21	consider the costs of implementing the standards. [See generally, Whitman v.
22	American Trucking Associations, 531 U.S. 457, 465-472, 475-76. (2001)]. Likewise,
23	"[a]ttainability and technological feasibility are not relevant considerations in the
24	promulgation of national ambient air quality standards." American Petroleum
25	Institute v. Costle, 665 F. 2d at 1185.
26	Section 109(d)(1) requires that "not later than December 31, 1980, and at 5-year
27	intervals thereafter, the Administrator shall complete a thorough review of the
28	criteria published under section 108 and the national ambient air quality standards
29	and shall make such revisions in such criteria and standards and promulgate such
30	new standards as may be appropriate" Section 109(d)(2) requires that an
31	independent scientific review committee "shall complete a review of the criteria
32	and the national primary and secondary ambient air quality standards and shall
33	recommend to the Administrator any new standards and revisions of existing
34	criteria and standards as may be appropriate" Since the early 1980s, this
35	independent review function has been performed by CASAC.

History of the NAAQS for Ozone

1	Tropospheric (ground-level) O_3 is the indicator for the mix of photochemical
2	oxidants (e.g., peroxyacetyl nitrate, hydrogen peroxide) formed from biogenic and
3	anthropogenic precursor emissions. Naturally occurring O_3 in the troposphere can
4	result from biogenic organic precursors reacting with naturally occurring nitrogen
5	oxides (NO _X) and by stratospheric O_3 intrusion into the troposphere. Anthropogenic
6	precursors of O_3 , especially NO_X , and volatile organic compounds (VOCs), originate
7	from a wide variety of stationary and mobile sources. Ambient O ₃ concentrations
8	produced by these emissions are directly affected by temperature, solar radiation,
9	wind speed, and other meteorological factors.
10	NAAQS are comprised of four basic elements: indicator, averaging time, level, and
11	form. The indicator defines the pollutant to be measured in the ambient air for the
12	purpose of determining compliance with the standard. The averaging time defines the
13	time period over which air quality measurements are to be obtained and averaged or
14	cumulated, considering evidence of effects associated with various time periods of
15	exposure. The level of a standard defines the air quality concentration used (i.e., an
16	ambient concentration of the indicator pollutant) in determining whether the standard
17	is achieved. The form of the standard specifies the air quality measurements that are
18	to be used for compliance purposes (e.g., the annual fourth-highest daily maximum
19	8-hour concentration, averaged over 3 years), and whether the statistic is to be
20	averaged across multiple years. These four elements taken together determine the
21	degree of public health and welfare protection afforded by the NAAQS.

Table IIISummary of primary and secondary NAAQS promulgated for ozone
during the period 1971-2008

Final Rule	Indicator	Avg Time	Level (ppm)	Form
1971 (36 FR 8186)	Total photochemical oxidants	1-h	0.08	Not to be exceeded more than 1 hour per year
1979 (44 FR 8202)	O ₃	1-h	0.12	Attainment is defined when the expected number of days per calendar year, with maximum hourly average concentration greater than 0.12 ppm, is ≤ 1
1993 (58 FR 13008)	EPA decided that	revisions to	the standards were	not warranted at the time.
1997 (62 FR 38856)	O ₃	8-h	0.08	Annual fourth-highest daily maximum 8-h concentration averaged over 3 years
2008 (73 FR 16483)	O ₃	8-h	0.075	Form of the standards remained unchanged relative to the 1997 standard

1	Table III summarizes the O_3 NAAQS that have been promulgated to date. In each
2	review, the secondary standard has been set to be identical to the primary standard.
3	These reviews are briefly described below.
4	EPA first established primary and secondary NAAQS for photochemical oxidants in
5	1971 . Both primary and secondary standards were set at a level of 0.08 parts per
6	million (ppm), 1-h avg, total photochemical oxidants, not to be exceeded more than
7	1 hour per year. The standards were based on scientific information contained in the
8	1970 AQCD.
9	In 1977, EPA announced the first periodic review of the 1970 AQCD in accordance
10	with Section 109(d)(1) of the Clean Air Act. In 1978, EPA published an AQCD.
11	Based on the 1978 AQCD, EPA published proposed revisions to the original
12	NAAQS in 1978 (<u>U.S. EPA, 1978b</u>) and final revisions in 1979 (<u>U.S. EPA, 1979a</u>).
13	The level of the primary and secondary standards was revised from 0.08 to 0.12 ppm;
14	the indicator was revised from photochemical oxidants to O_3 ; and the form of the
15	standards was revised from a deterministic to a statistical form, which defined
16	attainment of the standards as occurring when the expected number of days per
17	calendar year with maximum hourly average concentration greater than 0.12 ppm is
18	equal to or less than one.
19	In 1982, EPA announced plans to revise the 1978 AQCD (U.S. EPA, 1978a). In
19 20	In 1982, EPA announced plans to revise the 1978 AQCD (U.S. EPA, 1978a). In 1983, EPA announced that the second periodic review of the primary and secondary
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20 21 22	1983, EPA announced that the second periodic review of the primary and secondary standards for O_3 had been initiated (<u>U.S. EPA, 1983</u>). EPA subsequently published the 1986 O_3 AQCD (<u>U.S. EPA, 1986</u>) and 1989 Staff Paper (<u>U.S. EPA, 1989</u>).
20 21 22 23	1983, EPA announced that the second periodic review of the primary and secondary standards for O ₃ had been initiated (<u>U.S. EPA, 1983</u>). EPA subsequently published the 1986 O ₃ AQCD (<u>U.S. EPA, 1986</u>) and 1989 Staff Paper (<u>U.S. EPA, 1989</u>). Following publication of the 1986 O ₃ AQCD, a number of scientific abstracts and
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20 21 22 23 24 25 26 27 28 29 30 31 32 33 34	1983, EPA announced that the second periodic review of the primary and secondary standards for O_3 had been initiated (U.S. EPA, 1983). EPA subsequently published the 1986 O_3 AQCD (U.S. EPA, 1986) and 1989 Staff Paper (U.S. EPA, 1989). Following publication of the 1986 O_3 AQCD, a number of scientific abstracts and articles were published that appeared to be of sufficient importance concerning potential health and welfare effects of O_3 to warrant preparation of a Supplement to the 1986 O_3 AQCD (Costa et al., 1992). Under the terms of a court order, on August 10, 1992, EPA published a proposed decision (U.S. EPA, 1992) stating that revisions to the existing primary and secondary standards were not appropriate at the time (U.S. EPA, 1992). This notice explained that the proposed decision would complete EPA's review of information on health and welfare effects of O_3 assembled over a 7-year period and contained in the 1986 O_3 AQCD (U.S. EPA, 1986) and its Supplement to the 1986 O_3 AQCD (Costa et al., 1992). The proposal also announced EPA's intention to proceed as rapidly as possible with the next review of the air quality criteria and standards for O_3 in light of emerging evidence of health effects

1	In August 1992, EPA announced plans to initiate the third periodic review of the air
2	quality criteria and O_3 NAAQS (U.S. EPA, 1992). On the basis of the scientific
3	evidence contained in the 1996 O_3 AQCD and the 1996 Staff Paper (U.S. EPA,
4	<u>1996</u>), and related technical support documents, linking exposures to ambient O_3 to
5	adverse health and welfare effects at levels allowed by the then existing standards,
6	EPA proposed to revise the primary and secondary O_3 standards on December 13,
7	1996 (U.S. EPA, 1996d). The EPA proposed to replace the then existing 1-hour
8	primary and secondary standards with 8-h avg O_3 standards set at a level of 0.08 ppm
9	(equivalent to 0.084 ppm using standard rounding conventions). The EPA also
10	proposed, in the alternative, to establish a new distinct secondary standard using a
11	biologically based cumulative seasonal form. The EPA completed the review on July
12	18, 1997 by setting the primary standard at a level of 0.08 ppm, based on the annual
13	fourth-highest daily maximum 8-h avg concentration, averaged over 3 years, and
14	setting the secondary standard identical to the revised primary standard (U.S. EPA,
15	<u>1997</u>).
16	On May 14, 1999, in response to challenges to EPA's 1997 decision by industry and
17	others, the U.S. Court of Appeals for the District of Columbia Circuit (D.C. Cir.)
18	remanded the O_3 NAAQS to EPA, finding that Section 109 of the CAA, as
19	interpreted by EPA, effected an unconstitutional delegation of legislative authority.
20	In addition, the D.C. Cir. directed that, in responding to the remand, EPA should
21	consider the potential beneficial health effects of O ₃ pollution in shielding the public
22	from the effects of solar ultraviolet (UV) radiation, as well as adverse health effects.
23	On January 27, 2000, EPA petitioned the U.S. Supreme Court for certiorari on the
24	constitutional issue (and two other issues) but did not request review of the D.C. Cir.,
25	ruling regarding the potential beneficial health effects of O ₃ . On February 27, 2001,
26	the U.S. Supreme Court unanimously reversed the judgment of the D.C. Cir. on the
27	constitutional issue, holding that Section 109 of the CAA does not delegate
• •	

28	legislative power to the EPA in contravention of the Constitution, and remanded the
29	case to the D.C. Cir. to consider challenges to the O_3 NAAQS that had not been
30	addressed by that Court's earlier decisions. On March 26, 2002, the D.C. Cir. issued
31	its final decision, finding the 1997 O_3 NAAQS to be "neither arbitrary nor
32	capricious," and denied the remaining petitions for review. On November 14, 2001,
33	in response to the D.C. Cir. remand to consider the potential beneficial health effects
34	of O ₃ pollution in shielding the public from effects of solar (UV) radiation, EPA
35	proposed to leave the 1997 8-h O_3 NAAQS unchanged (U.S. EPA, 2001). After
36	considering public comment on the proposed decision, EPA published its final
37	response to this remand on January 6, 2003, reaffirming the 8-h O_3 NAAQS set in
38	1997 (U.S. EPA, 2003). On April 30, 2004, EPA announced the decision to make the
39	1-h O ₃ NAAQS no longer applicable to areas 1 year after the effective date of the

designation of those areas for the 8-h NAAQS (2004). For most areas, the date that
the 1-h NAAQS no longer applied was June 15, 2005.

- 3 EPA initiated the next periodic review if the air quality criteria and O₃ standards in
 4 September 2000 with a call for information (U.S. EPA, 2000). The schedule for
- completion of that rulemaking later became governed by a consent decree resolving a
 lawsuit filed in March 2003 by a group of plaintiffs representing national
- environmental and public health organizations. Based on the 2006 O₃ AQCD (U.S.
 EPA, 2006b) published in March 2006, the Staff Paper (U.S. EPA, 2007b) and
 related technical support documents, the proposed decision was published in the
 Federal Register on July 11, 2007 (U.S. EPA, 2007a). The EPA proposed to revise
 the level of the primary standard to a level within the range of 0.075 to 0.070 ppm.
 Two options were proposed for the secondary standard: (1) replacing the current
- 13 standard with a cumulative, seasonal standard, expressed as an index of the annual 14 sum of weighted hourly concentrations cumulated over 12 daylight hours during the 15 consecutive 3-month period within the O_3 season with the maximum index value, set 16 at a level within the range of 7 to 21 ppm-h; and (2) setting the secondary standard 17 identical to the revised primary standard. The EPA completed the rulemaking with 18 publication of a final decision on March 27, 2008 (U.S. EPA, 2008e), revising the 19 level of the 8-hour primary O_3 standard from 0.08 ppm to 0.075 ppm and revising the 20 secondary standard to be identical to the primary standard.
- 21 In May 2008, state, public health, environmental, and industry petitioners filed suit 22 against EPA regarding that final decision. At EPA's request the consolidated cases 23 were held in abeyance pending EPA's reconsideration of the 2008 decision. A notice 24 of proposed rulemaking to reconsider the 2008 final decision was issued by the 25 Administrator on January 6, 2010. Three public hearings were held. The Agency 26 solicited CASAC review of the proposed rule on January 25, 2010 and additional 27 CASAC advice on January 26, 2011. On September 2, 2011, the Office of 28 Management and Budget returned the draft final rule on reconsideration to EPA for 29 further consideration. EPA decided to coordinate further proceedings on its voluntary 30 rulemaking on reconsideration with the ongoing periodic review, by deferring the 31 completion of its voluntary rulemaking on reconsideration until it completes its 32 statutorily-required periodic review. In light of that, the litigation on the 2008 final 33 decision is no longer being held in abeyance and is proceeding. The 2008 ozone 34 standards remain in effect.

1 2

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1 EXECUTIVE SUMMARY

Introduction and Purpose

1	The purpose of this Integrated Science Assessment (ISA) is to provide a synthesis and
2	evaluation of the most policy-relevant science that forms the scientific foundation for the
3	review of the primary (health-based) and secondary (welfare-based) national ambient air
4	quality standard (NAAQS) for ozone (O ₃) and related photochemical oxidants. The ISA
5	is intended to inform the EPA Risk and Exposure Assessment and Policy Assessment and
6	thereby support decisions by the EPA Administrator on the NAAQS for O_3 (See Figure I
7	in Preamble). The current primary O_3 standard includes an 8-hour average standard set in
8	2008 at 75 parts per billion (ppb). The secondary standard for O_3 is equal to the primary
9	standard. The current primary NAAQS protects against respiratory health effects incurred
10	after short-term exposure to O ₃ , while the secondary NAAQS protects against damage to
11	vegetation and ecosystems.

Scope and Methods

12	EPA has developed an extensive and robust process for evaluating the scientific evidence
13	and drawing conclusions regarding air pollution-related health and welfare effects, which
14	is applied to the health and welfare effects resulting from current ambient concentrations
15	of O ₃ . Building upon the findings of previous assessments, this review includes
16	identification, selection, evaluation, and integration of the relevant results pertaining to
17	the atmospheric science of O_3 ; short- and long-term exposure to ambient O_3 ; health
18	effects due to relevant O3 concentrations as characterized in epidemiologic, controlled
19	human exposure, and toxicological studies; and ecological or welfare effects; as well as
20	O3 concentration-response relationships, mode(s) of action, and populations at increased
21	risk for O_3 -related health effects. The conclusions and key findings from previous
22	reviews provide the foundation for the consideration of evidence from recent studies (i.e.,
23	studies published since the completion of the 2006 O ₃ AQCD). Conclusions are drawn
24	based on the synthesis of evidence across disciplines from recent studies and building
25	upon the extensive evidence presented in previous reviews.
26	EPA has developed a consistent and transparent approach to evaluate the causal nature of
27	air pollution-related health and environmental effects for use in developing ISAs; the

framework for causal determinations is described in the Preamble to this document.
Causality determinations are based on the evaluation and synthesis of evidence across
scientific disciplines; however, the type of evidence that is most important for such
determinations will vary by pollutant or assessment. EPA assesses the entire body of

- 1 peer-reviewed literature, building upon evidence available during the previous NAAOS 2 reviews, to draw conclusions on the causal relationships between relevant pollutant 3 concentrations and health or welfare effects. EPA also evaluates the quantitative evidence 4 and draws scientific conclusions, to the extent possible, regarding the 5 concentration-response relationships and the loads to ecosystems, exposure doses or 6 concentrations, duration and pattern of exposures at which effects are observed. 7 A five-level hierarchy is used to classify the weight of evidence for causation, not just 8 association. This weight of evidence evaluation is based on various lines of evidence 9 from across the health and environmental effects disciplines. These separate judgments 10 are integrated into a qualitative statement about the overall weight of the evidence and 11 causality. The causal determinations are:
- 12 Causal relationship

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- Likely to be a causal relationship
- Suggestive of a causal relationship
 - Inadequate to infer a causal relationship
- Not likely to be a causal relationship

Ambient Ozone Concentrations

- 17 Ozone is naturally present in the stratosphere, where it serves the beneficial role of 18 blocking harmful ultraviolet radiation from the Sun, and preventing the majority of this 19 radiation from reaching the surface of the Earth. However, in the troposphere, O_3 acts as 20 a powerful oxidant and can harm living organisms and materials. Tropospheric O_3 is 21 present not only in polluted urban air, but throughout the globe. Ozone can be influenced 22 by local meteorological conditions, circulation patterns, emissions, and topographic 23 barriers, resulting in heterogeneous concentrations across an individual urban area. On a 24 larger scale, O_3 persists in the atmosphere long enough that it can be transported from 25 continent to continent. 26 Ozone in the troposphere originates from both anthropogenic (i.e., man-made) and 27 natural source categories. Ozone attributed to anthropogenic sources is formed in the 28 atmosphere by photochemical reactions involving sunlight and precursor pollutants 29 including volatile organic compounds, nitrogen oxides, and carbon monoxide. Ozone 30 attributed to natural sources is formed through the same photochemical reactions 31 involving natural emissions of precursor pollutants from vegetation, microbes, animals, 32 biomass burning, and lightning. Because O_3 is produced downwind of urban source areas
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and O_3 tends to persist longer in rural than in urban areas as a result of lower chemical

- scavenge, resulting in concentrations in rural areas that are often higher than those in
 urban areas.
- 3 In the context of a review of the NAAQS, it is useful to define background O3 4 concentrations in a way that distinguishes between concentrations that result from 5 precursor emissions that are relatively less controllable from those that are relatively 6 more controllable through U.S. policies. North American (NA) background can be 7 defined as those concentrations resulting from natural sources everywhere in the world 8 and from anthropogenic sources outside the U.S., Canada and Mexico. Since NA 9 background is a construct that cannot be measured, NA background O₃ concentrations are 10 estimated using chemistry transport models. Seasonal mean daily maximum 8-h average 11 NA background O₃ concentrations are generally higher in spring than in summer across 12 the U.S. The highest estimates are found in the Intermountain West during the spring and 13 in the Southwest during the summer. The lowest estimates occur over the East in the 14 spring and over the Northeast in the summer (See Section 3.4).

Human Exposure to Ozone

15 The widespread presence of O_3 in the environment results in exposure as people 16 participate in normal daily activities. The relationship between personal exposure and 17 ambient concentration measured at fixed-site monitors can be described in terms of 18 correlation, or how they covary in time, and ratio, which describes their relative 19 mangnitude. Personal-ambient O_3 correlations are generally moderate (0.3-0.8), although 20 low correlations have been observed with increased time spent indoors, low air exchange 21 rate, and concentrations below the personal sampler detection limit (See Section 4.3). 22 Ratios of 0.1-0.3 between personal exposure and ambient concentration have been 23 observed for the general population, with ratios of up to 0.9 observed for outdoor 24 workers. Evidence suggests that some groups, particularly children, older adults, and 25 those with respiratory problems, change their behavior on high-O₃ days to reduce 26 exposure (See Section 4.4.2). Such behavioral changes may result in reduced effect 27 estimates in epidemiologic studies that do not account for averting behavior on high-O₃ 28 days. Variation in O_3 concentrations occurs over multiple spatial and temporal scales, and 29 this introduces exposure error into epidemiologic results (See Section 4.6.2). However, 30 epidemiologic studies evaluating the influence of spatial scale and monitor selection find 31 little difference among effect estimates, and comparable risk estimates have been 32 reported in studies using a variety of exposure assessment techniques expected to produce 33 different levels of personal-ambient associations. This suggests that there is no clear 34 indication that a particular method of exposure assessment produces stronger 35 epidemiologic results.

Dosimetry and Modes of Action

1	When O_3 is inhaled, the amount of O_3 that is absorbed is affected by a number of factors
2	including the shape and size of the respiratory tract, route of breathing (nose or mouth),
3	as well as how quickly and deeply a person is breathing. The site of the greatest O_3 dose
4	to the lung tissue is the junction of the conducting airway and the gas exchange region, in
5	the deeper portion of the respiratory tract. Additionally, the primary site of O_3 uptake
6	moves deeper into the respiratory tract during exercise when breathing becomes faster
7	and the breathing route changes from the nose only to oronasal breathing (i.e., through
8	the nose and mouth) (See Section 5.2).
9	Once O_3 has been absorbed, there are several key events in the toxicity pathway of O_3 in
10	the respiratory tract that lead to O_3 -induced health effects (See Section 5.3). These
11	include formation of secondary oxidation products in the lung, activation of neural
12	reflexes, initiation of inflammation, alterations of epithelial barrier function, sensitization
13	of bronchial smooth muscle, modification of innate and adaptive immunity, and airway
14	remodeling. Another key event, systemic inflammation and vascular oxidative/nitrosative
15	stress, may be critical to the extrapulmonary effects of O_3 .
16	Overall, biological responses to O_3 exposure are common across many species (See
17	Section 5.5). Thus, animal studies are used to add to the understanding of the full range
18	of potential O ₃ -mediated health effects.

Integration of Ozone Health Effects

19	The body of evidence from short-term (i.e., hours, days, weeks) or long-term
20	(i.e., months to years) exposure studies is evaluated and integrated across scientific
21	disciplines (i.e., controlled human exposure studies, toxicology, and epidemiology) and
22	interpreted for the health effects evidence that spans all lifestages, and which vary in
23	severity from minor subclinical effects to death. The results from the health studies,
24	supported by the evidence from atmospheric chemistry and exposure assessment studies,
25	contribute to the causal determinations made for the health outcomes. The conclusions
26	from the previous NAAQS review and the causality determinations from this review are
27	summarized in Table 1-1. Additional details are provided here for respiratory health
28	effects and mortality, for which there is the strongest evidence of an effect from O_3 ;
29	details for a wider range of health effects are provided subsequent chapters.

Table 1-1Summary of ozone causal determinations by exposure duration
and health outcome

Health Outcome	Conclusions from Previous Review	Conclusions from 2012 3rd Draft ISA
Short-Term Exposure to O ₃		
Respiratory effects	The overall evidence supports a causal relationship between acute ambient O_3 exposures and increased respiratory morbidity outcomes.	Causal Relationship
Cardiovascular effects	The limited evidence is highly suggestive that O_3 directly and/or indirectly contributes to cardiovascular-related morbidity, but much remains to be done to more fully substantiate the association.	Suggestive of a Causal Relationship
Central nervous system effects	Toxicological studies report that acute exposures to O_3 are associated with alterations in neurotransmitters, motor activity, short and long term memory, sleep patterns, and histological signs of neurodegeneration.	Suggestive of a Causal Relationship
Total Mortality	The evidence is highly suggestive that O_3 directly or indirectly contributes to non-accidental and cardiopulmonary-related mortality.	Likely to be a Causal Relationship
Long-term Exposur	e to O ₃	
Respiratory effects	The current evidence is suggestive but inconclusive for respiratory health effects from long-term O_3 exposure.	Likely to be a Causal Relationship
Cardiovascular effects	No studies from previous review.	Suggestive of a Causal Relationship
Reproductive and developmental effects	Limited evidence for a relationship between air pollution and birth-related health outcomes, including mortality, premature births, low birth weights, and birth defects, with little evidence being found for O_3 effects.	Suggestive of a Causal Relationship
Central nervous system effects	Evidence regarding chronic exposure and neurobehavioral effects was not available.	Suggestive of a Causal Relationship
Cancer	Little evidence for a relationship between chronic O_3 exposure and increased risk of lung cancer.	Inadequate to infer a Causal Relationship
Total Mortality	There is little evidence to suggest a causal relationship between chronic O_3 exposure and increased risk for mortality in humans.	Suggestive of a Causal Relationship

Respiratory Effects

1	The clearest evidence for health effects associated with exposure to O_3 is provided by
2	studies of respiratory effects. Collectively, a very large amount of evidence spanning
3	several decades supports a relationship between exposure to O_3 and a broad range of
4	respiratory effects (See Section $6.2.9$ and Section $7.2.8$). The majority of this evidence is
5	derived from studies investigating short-term exposure (i.e., hours to weeks) to O_3 ,
6	although animal toxicological studies and recent epidemiologic evidence demonstrate
7	that long-term exposure (i.e., months to years) may also be detrimental to the respiratory
8	system.
9	The last review concluded that there was clear, consistent evidence of a causal
10	relationship between short-term exposure to O3 and respiratory health effects. This causal
11	association is substantiated by the coherence of effects observed across recent controlled
12	human exposure, epidemiologic, and toxicological studies indicating associations of

1	short-term O ₃ exposures with a range of respiratory health endpoints from respiratory
2	tract inflammation to respiratory-related emergency department (ED) visits and hospital
3	admissions. Across disciplines, short-term O3 exposures induced or were associated with
4	statistically significant declines in lung function. An equally strong body of evidence
5	from controlled human exposure and toxicological studies demonstrated O_3 -induced
6	inflammatory responses, increased epithelial permeability, and airway
7	hyperresponsiveness. Toxicological studies provided additional evidence for O ₃ -induced
8	impairment of host defenses. Combined, these findings from experimental studies
9	provided support for epidemiologic evidence, in which short-term increases in O_3
10	concentration were consistently associated with increases in respiratory symptoms and
11	asthma medication use in children with asthma, respiratory-related hospital admissions,
12	and ED visits for COPD and asthma. Additionally, recent evidence supports the range of
13	respiratory effects induced by O ₃ by demonstrating that short-term increases in ambient
14	O3 concentrations can lead to respiratory mortality. The combined evidence across
15	disciplines supports a causal relationship between short-term O ₃ exposure and
16	respiratory effects.

17 Taken together, the recent epidemiologic studies of respiratory health effects (including 18 respiratory symptoms, new-onset asthma and respiratory mortality) combined with 19 toxicological studies in rodents and nonhuman primates, provide biologically plausible 20 evidence that there is likely to be a causal relationship between long-term exposure 21 to O_3 and respiratory effects. The strongest epidemiologic evidence for a relationship 22 between long-term O₃ exposure and respiratory effects is provided by studies that 23 demonstrate interactions between exercise or different genetic variants and long-term 24 measures of O₃ exposure on new-onset asthma in children; and increased respiratory 25 symptom effects in asthmatics. Additional studies of respiratory health effects and a 26 study of respiratory mortality provide a collective body of evidence supporting these 27 relationships. Studies considering other pollutants provide data suggesting that the effects 28 related to O_3 are independent from potential effects of the other pollutants. Some studies 29 provide evidence for a positive concentration-response relationship. Short-term studies 30 provide supportive evidence with increases in respiratory symptoms and asthma 31 medication use, hospital admissions and ED visits for all respiratory outcomes and 32 asthma, and decrements in lung function in children. The recent epidemiologic and 33 toxicological data base provides a compelling case to support the hypothesis that a 34 relationship exists between long-term exposure to ambient O₃ and measures of 35 respiratory health effects.

Mortality Effects

1	The last review concluded that the overall body of evidence was highly suggestive that
2	short-term exposure to O_3 directly or indirectly contributes to non-accidental and
3	cardiopulmonary-related mortality; but that additional research was needed to more fully
4	establish underlying mechanisms by which such effects occur. The evaluation of recent
5	multicity studies and a multicontinent study that have examined the association between
6	short-term O3 exposure and mortality found evidence that supports the conclusions of the
7	last review (See Section 6.6). These recent studies reported consistent positive
8	associations between short-term O_3 exposure and total (nonaccidental) mortality, with
9	associations being stronger during the warm season. They also added support for
10	associations between O_3 exposure and cardiovascular mortality being similar to or
11	stronger than those between O_3 exposure and respiratory mortality. Additionally, these
12	recent studies examined previously identified areas of uncertainty in the O3-mortality
13	relationship, and provide evidence that continues to support an association between short-
14	term O_3 exposure and mortality. The body of evidence indicates that there is likely to be
15	a causal relationship between short-term exposures to ${\sf O}_3$ and total mortality.

Populations Potentially at Increased Risk

17populations that are at increased risk for O_3 -related health effects; these studies do so b18examining groups within the study population, such as those with an underlying health	-
18 examining groups within the study population, such as those with an underlying health	
19 condition or genetic variant; categories of age, race, or sex; or by developing animal	
20 models that mimic the conditions associated with a health effect. Such studies have	
21 identified a multitude of factors that could potentially contribute to whether an individu	ıal
22 is at increased risk for O_3 -related health effects (See Chapter 8). The populations	
23 identified as having adequate evidence for increased risk of O ₃ -related health effects ar	e
24 individuals with asthma, younger and older age groups, individuals with reduced intake	•
25 of certain nutrients (i.e., Vitamins C and E), and outdoor workers. The evidence for oth	er
26 potential factors, including variations in multiple genes related to oxidative metabolism	1
27 or inflammation, sex, socioeconomic status, and obesity is suggestive of an increased	
risk, but further evidence is needed.	

Integration of Effects on Vegetation and Ecosystems

29The most policy-relevant information pertaining to the review of the NAAQS for the30effects of O3 on vegetation and ecosystems are evaluated and synthesized, integrating key31findings about plant physiology, biochemistry, whole plant biology, ecosystems and

1	exposure-response relationships. The welfare effects of O_3 can be observed across spatial
2	scales, starting at the cellular and subcellular level, then the whole plant and finally,
3	ecosystem-level processes. Ozone effects at small spatial scales, such as the leaf of an
4	individual plant, can result in effects at a continuum of larger spatial scales. These effects
5	include altered rates of leaf gas exchange, growth and reproduction at the individual plant
6	level and can result in changes in ecosystems, such as productivity, carbon storage, water
7	cycling, nutrient cycling, and community composition. The conclusions from the
8	previous NAAQS review and the causality determinations from this review are
9	summarized in the table below. Further discussion of these conclusions is provided below
10	for visible foliar injury, growth, productivity, and carbon storage, reduced yield and
11	quality of agricultural crops, water cycling, below-ground processing, community
12	composition, and O3 exposure-response relationships; discussion for all relevant welfare
13	effects is provided in Chapter <u>9</u> .

Table 1-2	Summary of ozone causal determination for welfare effects
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Vegetation and Ecosystem Effects	Conclusions from Previous Review	Conclusions from 2012 3rd Draft ISA
Visible Foliar Injury Effects on Vegetation	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause impaired aesthetic quality of many native plants and trees by increasing foliar injury.	Causal Relationship
Reduced Vegetation Growth	Data published since the 1996 O ₃ AQCD strengthen previous conclusions that there is strong evidence that current ambient O ₃ concentrations cause decreased growth and biomass accumulation in annual, perennial and woody plants, including agronomic crops, annuals, shrubs, grasses, and trees.	Causal Relationship
Reduced Productivity in Terrestrial Ecosystems	There is evidence that O₃ is an important stressor of ecosystems and that the effects of O_3 on individual plants and processes are scaled up through the ecosystem, affecting net primary productivity.	Causal Relationship
Reduced Carbon (C) Sequestration in Terrestrial Ecosystems	Limited studies from previous review	Likely to be a Causal Relationship
Reduced Yield and Quality of Agricultural Crops	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased yield and/or nutritive quality in a large number of agronomic and forage crops.	Causal Relationship
Alteration of Terrestrial Ecosystem Water Cycling	Ecosystem water quantity may be affected by O ₃ exposure at the landscape level.	Likely to be a Causal Relationship
Alteration of Below- ground Biogeochemical Cycles	Ozone-sensitive species have well known responses to O₃ exposure , including altered C allocation to below-ground tissues, and altered rates of leaf and root production, turnover, and decomposition. These shifts can affect overall C and N loss from the ecosystem in terms of respired C, and leached aqueous dissolved organic and inorganic C and N.	Causal Relationship
Alteration of Terrestrial Community Composition	Ozone may be affecting above- and below -ground community composition through impacts on both growth and reproduction. Significant changes in plant community composition resulting directly from O_3 exposure have been demonstrated.	Likely to be a Causal Relationship

Visible Foliar Injury

1	Visible foliar injury resulting from exposure to O ₃ has been well characterized and
2	documented over several decades on many tree, shrub, herbaceous and crop species.
3	Ozone-induced visible foliar injury symptoms on certain plant species are considered
4	diagnostic of exposure to O ₃ , as experimental evidence has clearly established a
5	consistent association, with greater exposure often resulting in greater and more prevalent
6	injury. Additional sensitive species showing visible foliar injury continue to be identified
7	from field surveys and verified in controlled exposure studies (See Section 9.4.2).
8	Overall, evidence is sufficient to conclude that there is a causal relationship between
9	ambient O_3 exposure and the occurrence of O_3 induced visible foliar injury on
10	sensitive vegetation across the U.S.

Growth, Productivity, Carbon Storage and Agriculture

11	Ambient O ₃ concentrations have long been known to cause decreases in photosynthetic
12	rates and plant growth. The O ₃ -induced effects at the plant scale may translate to the
13	ecosystem scale, and cause changes in productivity and C storage. The effects of O_3
14	exposure on photosynthesis, growth, biomass allocation, ecosystem production and
15	ecosystem C sequestration were reviewed for natural ecosystems (See Section 9.4.3), and
16	crop productivity and crop quality were reviewed for agricultural ecosystems (See
17	Section <u>9.4.4</u>). There is strong and consistent evidence that ambient concentrations of O_3
18	decrease plant photosynthesis and growth in numerous plant species across the U.S.
19	Studies conducted during the past four decades have also demonstrated unequivocally
20	that O_3 alters biomass allocation and plant reproduction. Studies at the leaf and plant
21	scales showed that O ₃ reduced photosynthesis and plant growth, providing coherence and
22	biological plausibility for the reported decreases in ecosystem productivity. In addition to
23	primary productivity, other indicators such as net ecosystem CO ₂ exchange and
24	C sequestration were often assessed by modeling studies. Model simulations consistently
25	found that O ₃ exposure caused negative impacts on those indicators, but the severity of
26	these impacts was influenced by multiple interactions of biological and environmental
27	factors. Although O_3 generally causes negative effects on ecosystem productivity, the
28	magnitude of the response varies among plant communities. Overall, evidence is
29	sufficient to conclude that there is a causal relationship between ambient O_3
30	exposure and reduced native plant growth and productivity, and a likely causal
31	relationship between O_3 exposure and reduced carbon sequestration in terrestrial
32	ecosystems.
33	The detrimental effect of O_3 on crop production has been recognized since the 1960's.

33The detrimental effect of O_3 on crop production has been recognized since the 1960's,34and current O_3 concentrations across the U.S. are high enough to cause yield loss for a

1	variety of agricultural crops including, but not limited to, soybean, wheat, potato,
2	watermelon, beans, turnip, onion, lettuce, and tomato. Continued increases in O ₃
3	concentration may further decrease yield in these sensitive crops while also initiating
4	yield losses in less sensitive crops. Research has linked increasing O ₃ concentration to
5	decreased photosynthetic rates and accelerated senescence, which are related to yield
6	(See Section $9.4.4$). Evidence is sufficient to conclude that there is a causal relationship
7	between O_3 exposure and reduced yield and quality of agricultural crops.

Water Cycling

8	Ozone can affect water use in plants and ecosystems through several mechanisms
9	including damage to stomatal functioning and loss of leaf area. Possible mechanisms for
10	O_3 exposure effects on stomatal functioning include the build-up of CO_2 in the
11	substomatal cavity, impacts on signal transduction pathways and direct O_3 impact on
12	guard cells. Regardless of the mechanism, O_3 exposure has been shown to alter stomatal
13	performance, which may affect plant and stand transpiration and therefore may affect
14	hydrological cycling (See Section $9.4.5$). Although the direction of the response differed
15	among studies, the evidence is sufficient to conclude that there is likely to be a causal
16	relationship between O_3 exposure and the alteration of ecosystem water cycling.

Below Ground Processes

17	Below-ground processes are tightly linked with above-ground processes. The responses
18	of above-ground process to O_3 exposure, such as reduced photosynthetic rates, increased
19	metabolic cost, and reduced root C allocation, have provided biologically plausible
20	mechanisms for the alteration of below-ground processes. These include altered quality
21	and quantity of C input to soil, microbial community composition, and C and nutrient
22	cycling (See Section $9.4.6$). The evidence is sufficient to conclude that there is a causal
23	relationship between O_3 exposure and the alteration of below-ground
24	biogeochemical cycles.

Community Composition

25	Ozone exposure changes competitive interactions and leads to loss of O ₃ -sensitive
26	species or genotypes. Studies at the plant level found that the severity of O_3 damage to
27	growth, reproduction, and foliar injury varied among species, which provided the
28	biological plausibility for the alteration of community composition (See Section $9.4.3$ and
29	Section <u>9.4.7</u>). For example, there is a tendency for O_3 exposure to shift the biomass of
30	grass-legume mixtures in favor of grass species. Ozone exposure not only altered

1	community composition of plant species, but also microorganisms: research since the last
2	review has shown that O_3 can also alter community composition and diversity of soil
3	microbial communities. Shifts in community composition of bacteria and fungi have been
4	observed in both natural and agricultural ecosystems, although no general patterns could
5	be identified. The evidence is sufficient to conclude that there is likely to be a causal
6	relationship between O_3 exposure and the alteration of community composition of
7	some ecosystems.

Ozone Exposure-Response Relationships

8	Previous reviews of the NAAQs have included exposure-response functions for the yield
9	of many crop species, and for the biomass accumulation of tree species. They were based
10	on large-scale experiments designed to obtain clear exposure-response data, and are
11	updated by using the W126 metric to quantify exposure. In recent years, extensive
12	exposure-response data obtained in more naturalistic settings have become available for
13	yield of soybean and growth of aspen. The exposure-response median functions are
14	validated based on previous data by comparing their predictions with the newer
15	observations (See Section 9.6). The functions supply very accurate predictions of effects
16	in naturalistic settings. Recent meta-analyses of large sets of crop and tree studies do not
17	give rise to exposure-response functions, but their results are consistent with the
18	functions presented in Section 9.6 . It is important to note that although these median
19	functions provide reliable models for groups of species or group of genotypes within a
20	species, the original data and recent results consistently show that some species, and
21	some genotypes within species are much more severely affected by exposure to O_3 .

The Role of Tropospheric Ozone in Climate Change and UV-B Effects

22Atmospheric O_3 plays an important role in the Earth's energy budget by interacting with23incoming solar radiation and outgoing infrared radiation. Tropospheric O_3 makes up only24a small portion of the total column of O_3 , but it has important incremental effects on the25overall radiation budget. Perturbations in tropospheric O_3 concentrations can have direct26effects on climate and indirect effects on health, ecology, and welfare by shielding the27earth's surface from solar ultraviolet (UV) radiation.

Radiative Forcing and Climate Change

28	Tropospheric O_3 is a major greenhouse gas, third in importance after CO_2 and CH_4
29	according to the IPCC (See Section 10.3). Models calculate that the global average

1	concentration of tropospheric O ₃ has doubled since the preindustrial era, while
2	observations indicate that in some regions O_3 may have increased by factors as great as 4
3	or 5. These increases are tied to the rise in emissions of O_3 precursors from human
4	activity, mainly fossil fuel consumption and agricultural processes. There are large
5	uncertainties in the radiative forcing estimate attributed to tropospheric O ₃ , making the
6	effect of tropospheric O_3 on climate more uncertain than the effect of the long-lived
7	greenhouse gases. Overall, the evidence supports a causal relationship between
8	changes in tropospheric O_3 concentrations and radiative forcing.
9	Radiative forcing does not take into account the climate feedbacks that could amplify or
9 10	Radiative forcing does not take into account the climate feedbacks that could amplify or dampen the actual surface temperature response. Quantifying the change in surface
10	dampen the actual surface temperature response. Quantifying the change in surface
10 11	dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and
10 11 12	dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily
10 11 12 13	dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given radiative forcing is highly uncertain
10 11 12 13 14	dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given radiative forcing is highly uncertain and can vary greatly among models and from region to region within the same model. In

UV-B Effects

18	UV radiation emitted from the Sun contains sufficient energy when it reaches the Earth to
19	have damaging effects on living organisms and materials (see Section 10.4). Atmospheric
20	O_3 plays a crucial role in reducing exposure to UV radiation at the Earth's surface. Ozone
21	in the stratosphere is responsible for the majority of this shielding, but O ₃ in the
22	troposphere provides supplemental shielding of UV radiation in the mid-wavelength
23	range (UV-B), thereby influencing human and ecosystem health and materials damage.
24	There is a lack of published studies that critically examine the incremental health or
25	welfare effects (adverse or beneficial) attributable specifically to changes in UV-B
26	exposure resulting from perturbations in tropospheric O_3 concentrations. While the
27	effects are expected to be small, they cannot yet be critically assessed within reasonable
28	uncertainty. Overall, the evidence is inadequate to determine if a causal relationship
29	exists between changes in tropospheric O_3 concentrations and effects on health and
30	welfare related to UV-B shielding.
31	The conclusions from the previous NAAQS review and the causality determinations from

31The conclusions from the previous NAAQS review and the causanty determinations from32this review relating climate change and UV-B effects are summarized in the table below33(Table 1-3), with details provided in Chapter 10.

Table 1-3Summary of ozone causal determination for climate change and
UV-B effects.

Effects	Conclusions from Previous Review	Conclusions from 2012 3rd Draft ISA
Radiative Forcing	Climate forcing by O_3 at the regional scale may be its most important impact on climate.	Causal Relationship
Climate Change	While more certain estimates of the overall importance of global-scale forcing due to tropospheric O_3 await further advances in monitoring and chemical transport modeling, the overall body of scientific evidence suggests that high concentrations of O_3 on the regional scale could have a discernible influence on climate, leading to surface temperature and hydrological cycle changes.	Likely to be a Causal Relationship
Health and Welfare Effects Related to UV-B Shielding	UV-B has not been studied in sufficient detail to allow for a credible health benefits assessment. In conclusion, the effect of changes in surface-level O_3 concentrations on UV-induced health outcomes cannot yet be critically assessed within reasonable uncertainty.	Inadequate to Determine if a Causal Relationship Exists

Conclusion

1	The clearest evidence for human health effects associated with exposure to O ₃ is provided
2	by studies of respiratory effects. Collectively, there is a very large amount of evidence
3	spanning several decades in support of a causal association between exposure to O_3 and a
4	broad range of respiratory effects. The majority of this evidence is derived from studies
5	investigating short-term O3 exposure (i.e., hours to weeks), although animal toxicological
6	studies and recent epidemiologic evidence demonstrate that long-term exposure
7	(i.e., months to years) may also be detrimental to the respiratory system. Additionally,
8	consistent positive associations between short-term O ₃ exposure and total (nonaccidental)
9	mortality have helped to resolve previously identified areas of uncertainty in the
10	O_3 -mortality relationship, indicating that there is likely to be a causal relationship
11	between short-term exposures to O_3 and total mortality. Taken together, the recent
12	epidemiologic studies of respiratory health effects (including respiratory symptoms, new-
13	onset asthma and respiratory mortality) combined with toxicological studies in rodents
14	and nonhuman primates, provide biologically plausible evidence that there is likely to be
15	a causal relationship between long-term exposure to O_3 and respiratory effects.
16	Recent evidence is suggestive of a causal relationship between long-term O_3
17	exposures and total mortality. The evidence for these health effects indicates that the
18	relationship between concentration and response is linear along the range of O_3
19	concentrations observed in the U.S., with no indication of a threshold. However, there is
20	less certainty in the shape of the concentration-response curve at O_3 concentrations
21	generally below 20 ppb. The populations identified as having increased risk of O ₃ -related
22	health effects are individuals with asthma, younger and older age groups, individuals with
23	certain dietary deficiencies, and outdoor workers.

1	There has been over 40 years of research on the effects of O_3 exposure on vegetation and
2	ecosystems. The best evidence for effects is from controlled exposure studies. These
3	studies have clearly shown that exposure to O_3 is causally linked to visible foliar injury,
4	decreased photosynthesis, changes in reproduction, and decreased growth. Recently,
5	studies at larger spatial scales support the results from controlled studies and indicate that
6	ambient O3 exposures can affect ecosystem productivity, crop yield, water cycling, and
7	ecosystem community composition. And on a global scale, tropospheric O_3 is the third
8	most important greenhouse gas, playing an important role in climate change.

2 INTEGRATIVE SUMMARY

1	This Integrated Science Assessment (ISA) forms the scientific foundation for the review
2	of the national ambient air quality standards (NAAQS) for ozone (O_3) . The ISA is a
3	concise evaluation and synthesis of the most policy-relevant science—and it
4	communicates critical science judgments relevant to the review of the NAAQS for O_3 .
5	The ISA accurately reflects "the latest scientific knowledge useful in indicating the kind
6	and extent of identifiable effects on public health or welfare which may be expected from
7	the presence of [a] pollutant in ambient air" (CAA, 1990a). Key information and
8	judgments contained in prior Air Quality Criteria Documents (AQCD) for O3 are
9	incorporated into this assessment. Additional details of the pertinent scientific literature
10	published since the last review, as well as selected earlier studies of particular interest,
11	are included. This ISA thus serves to update and revise the evaluation of the scientific
12	evidence available at the time of the completion of the 2006 O_3 AQCD. The current
13	primary O_3 standard includes an 8-hour (h) average (avg) standard set at 75 parts per
14	billion (ppb). The secondary standard for O_3 is set equal to the primary standard. Further
15	information on the legislative and historical background for the O3 NAAQS is contained
16	in the Preface to this ISA.
17	This chapter summarizes and synthesizes the available scientific evidence and is intended
17 18	This chapter summarizes and synthesizes the available scientific evidence and is intended to provide a concise synopsis of the ISA conclusions and findings that best inform
18	to provide a concise synopsis of the ISA conclusions and findings that best inform
18 19	to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA,
18 19 20	to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes:
18 19 20 21	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and
18 19 20 21 22	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in
18 19 20 21 22 23	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations.
18 19 20 21 22 23 24	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations. An integration of the evidence on the welfare effects associated with exposure
 18 19 20 21 22 23 24 25 	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations.
 18 19 20 21 22 23 24 25 26 	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations. An integration of the evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and
 18 19 20 21 22 23 24 25 26 27 	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations. An integration of the evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and discussion of important uncertainties identified in the interpretation of the scientific evidence in the interpretation of the scientific evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and discussion of important uncertainties identified in the interpretation of the scientific evidence.
 18 19 20 21 22 23 24 25 26 27 28 	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations. An integration of the evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and discussion of important uncertainties identified in the interpretation of the scientific evidencies in the interpretation of the scientific evidencies. Discussion of policy-relevant considerations, such as potentially at-risk
 18 19 20 21 22 23 24 25 26 27 28 29 	 to provide a concise synopsis of the ISA conclusions and findings that best inform consideration of the policy-relevant questions that frame this assessment (U.S. EPA, 2009c). It includes: An integration of the evidence on the health effects associated with short- and long-term exposure to O₃, discussion of important uncertainties identified in the interpretation of the scientific evidence, and an integration of health evidence from the different scientific disciplines and exposure durations. An integration of the evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and discussion of important uncertainties identified in the interpretation of the scientific evidence in the interpretation of the scientific evidence on the welfare effects associated with exposure to O₃, including those associated with vegetation and ecosystems, and discussion of important uncertainties identified in the interpretation of the scientific evidence.

2.1 ISA Development and Scope

1 2 3 4 5 6 7 8	EPA has a developed a robust, consistent, and transparent process for evaluating the scientific evidence and drawing conclusions and causal judgments regarding air pollution-related health and environmental effects. The ISA development process includes literature search strategies, criteria for selecting and evaluating studies, approaches for evaluating weight of the evidence, and a framework for making causality determinations. The process and causality framework are described in more detail in the Preamble to the ISA. This section provides a brief overview of the process for development of this ISA.
9	EPA initiated the current review of the NAAQS for O_3 on September 29, 2008, with a
10	call for information from the public (U.S. EPA, 2008f). Literature searches were
11	conducted routinely to identify studies published since the last review, focusing on
12	studies published from 2005 (close of previous scientific assessment) through July 2011.
13	References that were considered for inclusion in this ISA can be found using the HERO
14	website (http://hero.epa.gov/ozone). This site contains HERO links to lists of references
15	that are cited in the ISA, as well as those that were considered for inclusion, but not cited
16	in the ISA, with bibliographic information and abstracts.
17	This review has endeavored to evaluate all relevant data published since the last review;
18	this includes studies pertaining to the atmospheric science of O ₃ , human exposure to
19	ambient O_3 , and health, ecological, climate and UV-B effects studies. These include
20	studies that are related to concentration-response relationships, mode(s) of action (MOA),
21	and understanding of at-risk populations for effects of O_3 exposure. Added to the body of
22	research were EPA's analyses of air quality and emissions data, studies on atmospheric
23	chemistry, transport, and fate of these emissions.
24	Previous AQCDs (U.S. EPA, 2006b, 1996a, b, 1984, 1978a) have included an extensive
25	body of evidence on both health and welfare effects of O_3 exposure, as well as an
26	understanding of the atmospheric chemistry of O_3 (U.S. EPA, 2006b). In this ISA, the
27	conclusions and key findings from previous reviews are summarized at the beginning of
28	each section, to provide the foundation for consideration of evidence from recent studies.
29	Results of key studies from previous reviews are included in discussions or tables and
30	figures, as appropriate, and conclusions are drawn based on the synthesis of evidence
31	from recent studies with the extensive literature summarized in previous reviews.
32	The Preamble discusses the general framework for conducting the science assessment
33	and developing an ISA, including criteria for evaluating studies and developing scientific
34	conclusions. For selection of epidemiologic studies in the O_3 ISA, particular emphasis is
35	placed on those studies most relevant to the review of the NAAQS. Studies conducted in

1	the United States (U.S.) or Canada are discussed in more detail than those from other
2	geographical regions, and in regard to human health, particular emphasis is placed on:
3	(1) recent multicity studies that employ standardized analysis methods for evaluating
4	effects of O_3 and that provide overall estimates for effects, based on combined analyses
5	of information pooled across multiple cities; (2) studies that help understand quantitative
6	relationships between exposure concentrations and effects; (3) new studies that provide
7	evidence on effects in at-risk populations; and (4) studies that consider and report O_3 as a
8	component of a complex mixture of air pollutants. In evaluating toxicological and
9	controlled human exposure studies, emphasis is placed on studies using concentrations
10	that are within about an order of magnitude of ambient O_3 concentrations. Consideration
10	of studies important for evaluation of human exposure to ambient O_3 places emphasis on
12	those evaluating the relationship between O_3 measured at central site monitors and
12	personal exposure to ambient O_3 . Important factors affecting this relationship include
14	spatial and temporal variations in ambient O_3 concentration, and time spent outdoors,
15	spatial and temporal variations in amolent O_3 concentration, and time spent outdoors, since penetrations of O_3 into indoor environments may be limited.
15	since penetrations of O ₃ into indoor environments may be ininted.
16	Epidemiologic studies generally present O3-related effect estimates for mortality and
17	morbidity health outcomes based on an incremental change in exposure, traditionally
18	equal to the interquartile range in O_3 concentrations or some other arbitrary value
19	(e.g., 10 ppb). Additionally, various averaging times are used in O ₃ epidemiologic studies,
20	with the three most common being the maximum 1-hour average within a 24-hour period
21	(1-h max), the maximum 8-hour average within a 24-h period (8-h max), and 24-hour
22	average (24-h avg). For the purpose of presenting results from studies that use different
23	exposure metrics, EPA consistently applies the same O ₃ increments to facilitate
24	comparisons between the results of various studies that may use different indices. These
25	increments were derived using the nationwide distributional data for O ₃ monitors in U.S.
26	Metropolitan Statistical Areas and are representative of a low-to-high change in O_3
27	concentrations and were approximated on the basis of annual mean to 95th percentile
28	differences (Langstaff, 2003). Therefore, throughout Chapter 6, efforts were made to
29	standardize O ₃ -related effect estimates using the increments of 20 ppb for 24-h avg,
30	30 ppb for 8-h max, and 40 ppb for 1-h max O ₃ concentrations, except as noted. In long-
31	term exposure studies, typically, O ₃ concentrations are lower and less variable when
32	averaged across longer exposure periods, and differences due to the use of varying
33	averaging times (e.g., 1-h max, 24-h avg) become less apparent. As such, in the long-term
34	exposure chapter (Chapter 7) an increment of 10 ppb was consistently applied across
35	studies, regardless of averaging time, to facilitate comparisons between the results from
36	these studies.

1	This ISA uses a five-level hierarchy that classifies the weight of evidence for causation:
2	 Causal relationship
3	 Likely to be a causal relationship
4	 Suggestive of a causal relationship
5	 Inadequate to infer a causal relationship
6	 Not likely to be a causal relationship
7	Beyond judgments regarding causality are questions relevant to quantifying health or
8	environmental risks based on the understanding of the quantitative relationships between
9	pollutant exposures and health or welfare effects. Once a determination is made regarding
10	the causal relationship between the pollutant and outcome category, important questions
11	regarding quantitative relationships include:
12	• What is the concentration-response, exposure-response, or dose-response
13	relationship?
14	 Under what exposure conditions (dose or concentration, duration and pattern)
15	are effects observed?
16	What populations appear to be differentially affected i.e., at-risk to effects?
17	 What elements of the ecosystem (e.g., types, regions, taxonomic groups,
18	populations, functions, etc.) appear to be affected or are more sensitive to
19	effects?
20	This chapter summarizes and integrates the newly available scientific evidence that best
21	informs consideration of the policy-relevant questions that frame this assessment.
22	Section 2.2 discusses the trends in ambient concentrations and sources of O_3 and provides
23	a brief summary of ambient air quality for short- and long-term exposure durations.
24	Section 2.3 presents the evidence regarding personal exposure to ambient O_3 in outdoor
25	and indoor microenvironments, and it discusses the relationship between ambient O ₃
26	concentrations and personal exposure to ambient O_3 . Section <u>2.4</u> provides a discussion of
27	the dosimetry and mode of action evidence for O_3 exposure. Section <u>2.5</u> integrates the
28	evidence for studies that examine the health effects associated with short- and long-term
29	exposure to O ₃ and discusses important uncertainties identified in the interpretation of the
30	scientific evidence. A discussion of policy-relevant considerations, such as potentially at-
31	risk populations, lag structure, and the O_3 concentration-response relationship is also
32	included in Section 2.5. Finally, Section 2.6 summarizes the evidence for welfare effects
33	related to O_3 exposure, and Section <u>2.7</u> reviews the literature on the influence of
34	tropospheric O_3 on climate and exposure to solar ultraviolet radiation.

2.2 Atmospheric Chemistry and Ambient Concentrations

2.2.1 Physical and Chemical Processes

1	Ozone in the troposphere is a secondary pollutant; it is formed by photochemical
2	reactions of precursor gases and is not directly emitted from specific sources. Ozone
3	precursor gases originate from both anthropogenic (i.e., man-made) and natural source
4	categories. Ozone attributed to anthropogenic sources is formed in the atmosphere by
5	photochemical reactions involving sunlight and precursor pollutants including volatile
6	organic compounds (VOCs), nitrogen oxides (NO _X), and carbon monoxide (CO). Ozone
7	attributed to natural sources is formed through similar photochemical reactions involving
8	natural emissions of precursor pollutants from vegetation, microbes, animals, biomass
9	burning, lightning, and geogenic sources. The distinction between natural and
10	anthropogenic sources of O3 precursors is often difficult to make in practice, as human
11	activities affect directly or indirectly emissions from what would have been considered
12	natural sources during the pre-industrial era. A schematic overview of the major
13	photochemical cycles influencing O_3 in the troposphere and the stratosphere is shown in
14	Figure 2-1. The processes depicted in this figure are fairly well understood, and were
15	covered in detail in the previous O ₃ AQCD. The formation of O ₃ , other oxidants, and
16	oxidation products from these precursors is a complex, nonlinear function of many
17	factors including: (1) the intensity and spectral distribution of sunlight reaching the lower
18	troposphere; (2) atmospheric mixing; (3) concentrations of precursors in the ambient air
19	and the rates of chemical reactions of these precursors; and (4) processing on cloud and
20	aerosol particles.

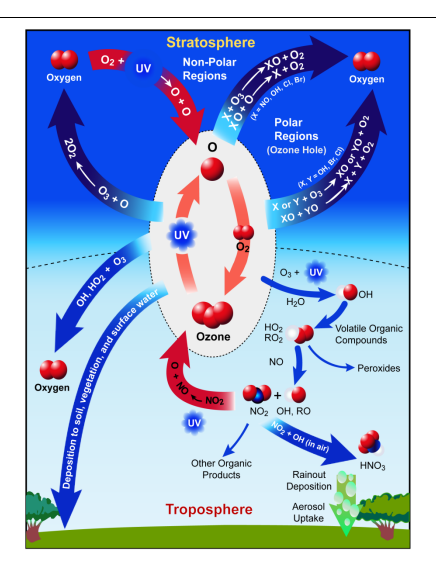


Figure 2-1 Schematic overview of photochemical processes influencing stratospheric and tropospheric ozone.

1	Ozone is present not only in polluted urban atmospheres but throughout the troposphere,
2	even in remote areas of the globe. Similar basic processes involving sunlight-driven
3	reactions of NO_X , VOCs and CO contribute to O_3 formation throughout the troposphere.
4	These processes also lead to the formation of other photochemical products, such as
5	peroxyacetyl nitrate, nitric acid, and sulfuric acid, and to other compounds, such as
6	formaldehyde and other carbonyl compounds. In urban areas, NO_X , VOCs, and CO are
7	all important for O_3 formation. In non-urban vegetated areas, biogenic VOCs emitted
8	from vegetation tend to be the most important precursor to O_3 formation. In the remote
9	troposphere, methane-structurally the simplest VOC-and CO are the main carbon-
10	containing precursors to O_3 formation. Ozone is subsequently removed from the

1 2	troposphere through a number of gas phase reactions and deposition to surfaces as shown in Figure 2-1.
3	Convective processes and turbulence transport O_3 and other pollutants both upward and
4	downward throughout the planetary boundary layer and the free troposphere. In many
5	areas of the U.S., O ₃ and its precursors can be transported over long distances, aided by
6	vertical mixing. The transport of pollutants downwind of major urban centers is
7	characterized by the development of urban plumes. Meteorological conditions, small-
8	scale circulation patterns, localized chemistry, and mountain barriers can influence
9	mixing on a smaller scale, resulting in frequent heterogeneous O3 concentrations across
10	individual urban areas.
11	Furthermore, because the mean tropospheric lifetime of O_3 is a few weeks, O_3 can be
12	transported from continent to continent. The degree of influence from intercontinental
13	transport varies greatly by location and time. For instance, high elevation sites are most
14	susceptible to the intercontinental transport of pollution, particularly during spring.
15	However, because the atmospheric chemistry of O_3 is quite complex and can be highly
16	non-linear in environments close to sources of precursors, isolating the influence of
17	intercontinental transport of O_3 and O_3 precursors on urban air quality is particularly
18	problematic.

2.2.2 Atmospheric Modeling of Background Ozone Concentrations

19	A number of recent studies indicate that natural sources such as wildfires and
20	stratospheric intrusions and the intercontinental transport of pollution can affect O_3 air
21	quality at specific times and in specific locations in the United States. These contributions
22	are in addition to contributions from dominant local pollution sources. To gain a broader
23	perspective and to isolate the influence of natural or transported O ₃ , estimates from
24	chemical transport models (CTMs) must be used. This is because observations within the
25	U.S.—even at relatively remote monitoring sites—are impacted by transport from
26	anthropogenic source regions within U.S. borders.
27	In the context of a review of the NAAQS, it is useful to define background O_3
27 28	In the context of a review of the NAAQS, it is useful to define background O_3 concentrations in a way that distinguishes between concentrations that result from
28	concentrations in a way that distinguishes between concentrations that result from
28 29	concentrations in a way that distinguishes between concentrations that result from precursor emissions that are relatively less controllable from those that are relatively
28 29 30	concentrations in a way that distinguishes between concentrations that result from precursor emissions that are relatively less controllable from those that are relatively more controllable through U.S. policies. For this assessment, three definitions of
28 29 30 31	concentrations in a way that distinguishes between concentrations that result from precursor emissions that are relatively less controllable from those that are relatively more controllable through U.S. policies. For this assessment, three definitions of background O_3 concentrations are considered, including (1) North American (NA)
28 29 30 31 32	concentrations in a way that distinguishes between concentrations that result from precursor emissions that are relatively less controllable from those that are relatively more controllable through U.S. policies. For this assessment, three definitions of background O_3 concentrations are considered, including (1) North American (NA) background (simulated O_3 concentrations that would exist in the absence of

1	
1	anthropogenic emissions from the U.S.), and (3) natural background (simulated O_3
2	concentrations in the absence of all anthropogenic emissions globally). Each definition of
3	background O ₃ includes contributions resulting from emissions from natural sources
4	(e.g., stratospheric intrusion, wildfires, biogenic methane and more short-lived VOC
5	emissions) throughout the globe. There is no chemical difference between background O_3
6	and O ₃ attributable to U.S. or North American anthropogenic sources. However, to
7	inform policy considerations regarding the current or potential alternative standards, it is
8	useful to understand how total O ₃ concentrations (i.e., O ₃ from all sources) can be
9	attributed to different sources.
10	Since background O ₃ concentrations as defined above are a construct that cannot be
11	directly measured, the range of background O ₃ concentrations is estimated using CTMs.
12	For the current assessment, the GEOS-Chem model at $0.5^{\circ} \times 0.667^{\circ}$ (~50 km×50 km)
13	horizontal resolution and a nested, hybrid GEOS-Chem/CAMx model at finer horizontal
14	resolution (12 km \times 12 km) were used. Results from these two models represent the latest
15	estimates for background O3 concentrations documented in the peer-reviewed literature
16	and are shown in <u>Table 2-1</u> . The R^2 for both models are generally <0.5, with CAMx
17	showing generally higher values than GEOS-Chem (Table 3-1). The GEOS-Chem
18	model-predicted seasonal mean daily maximum 8-h average O_3 concentrations for the
19	base case (i.e., including all anthropogenic and natural sources globally), U.S.
20	background, and NA background simulations during spring and summer 2006 are shown
21	in <u>Figure 2-2</u> .

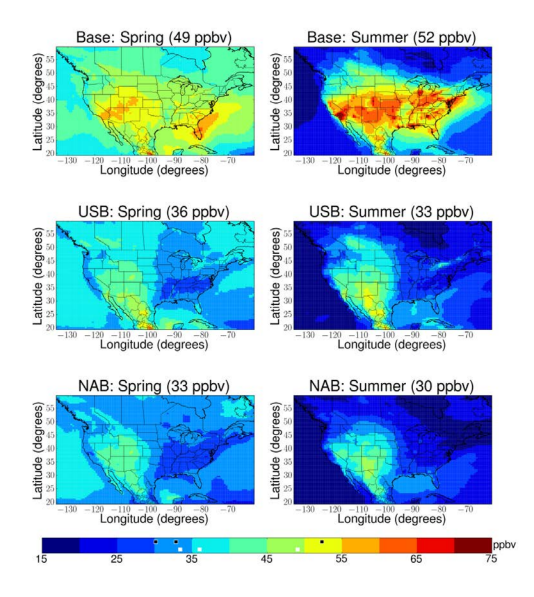
Table 2-1Comparison of seasonal mean MDA8 ozone concentrations
simulated by the GEOS-Chem and CAMx base case models for
2006, with measurements at CASTNET sites.

Region	CASTNET		GEOS-Chem		GEOS-Chem/CAMx	
	Spring	Summer	Spring	Summer	Spring	Summer
	58 ± 12 ^b	00 44	52 ± 11	66 ± 18	50 ± 10	66 ± 13
California (5) ^a		69 ± 14	$38 \pm 7^{\circ}$	37 ± 9	39 ± 6	42 ± 6
	54 0	55 44	53 ± 7	55 ± 11	49 ± 8	57 ± 10
West (14)	54 ± 9	55 ± 11	42 ± 6	40 ± 9	40 ± 7	41 ± 8
	47 ± 10 50		47 ± 8	51 ± 14	45 ± 11	54 ± 13
North Central (6)		50 ± 12	33 ± 6	27 ± 7	30 ± 6	31 ± 5
	40 40		45 ± 7	45 ± 13	46 ± 11	53 ± 14
Northeast (5)	48 ± 10 45 ± 14	45 ± 14	33 ± 7	24 ± 7	30 ± 5	27 ± 6
		/-	51 ± 7	54 ± 9	54 ± 9	61 ± 12
Southeast (9)	52 ± 11	52 ± 16	32 ± 7	29 ± 10	33 ± 6	30 ± 6

^aValues in parentheses after each region name refer to the number of sites.

^bShown are seasonal (spring, summer) mean daily maximum 8-h avg O₃ concentrations in ppb ± standard deviation.

^cNorth American background mean daily maximum 8-h avg O₃ concentrations (ppb ± standard deviation) are shown beneath the



Note: Mean daily average 8-h O₃ concentrations were calculated by GEOS-Chem for the base case (top, Base), United States background (middle, USB) and North American Background (lower, NAB). Values in parentheses (above each map) refer to continental U.S. means, and are shown in the color bar as black squares for summer and white squares for spring.

Source: Adapted from Zhang et al. (2011).

Figure 2-2 Mean daily average 8-hour ozone concentrations in surface air, for spring and summer 2006.

1	The main results from these modeling efforts can be summarized as follows.
2	 Simulated regional and seasonal means of base-case O₃ using both models
3	generally agree to within a few ppb with observations throughout the western
4	and central U.S., except in California; but GEOS-Chem shows better
5	agreement than CAMx in the eastern U.S. However, these results are likely to
6	change with updates to model chemistry and physics.
7	 Both models show background concentrations vary spatially and temporally.
8	NA background concentrations are generally higher in spring than in summer
9	across the U.S. Simulated mean NA background concentrations are highest in
10	the Intermountain West (i.e., at high altitude) in spring and in the Southwest in
11	summer. Lowest estimates of NA background occur in the East in the spring
12	and the Northeast in summer.
13	 NA background concentrations tend to increase with total (i.e., base case) O₃
14	concentrations at high elevation, but that tendency is not as clear at low
15	elevations.
16	 Comparison of NA background and natural background indicate that methane
17	is a major contributor to NA background O ₃ , accounting for slightly less than
18	half of the increase in background since the preindustrial era and whose
19	relative contribution is projected to grow in the future.
20	 U.S. background concentrations are on average 2.6 ppb higher than NA
21	background concentrations during spring and 2.7 ppb during summer across
22	the U.S. with highest increases above NA background over the Northern Tier
23	of New York State (19.1 ppb higher than NA background) in summer. High
24	values for U.S. background are also found in other areas bordering Canada
25	and Mexico.
26	 Contributions to background O₃ can be episodic or non-episodic; high
27	background concentrations are driven primarily by the episodic events such as
28	stratospheric intrusions and wildfires. The most pronounced differences
29	between these model results and observations are at the upper end of the
30	concentration distribution, particularly at high elevations.
31	Note that the calculations of background concentrations presented in this chapter were
32	formulated to answer the question, "what would O ₃ concentrations be if there were no
33	anthropogenic sources". This is different from asking, "how much of the O ₃ measured or
34	simulated in a given area is due to background contributions". Because of potentially
35	strong non-linearities—particularly in many urban areas—these estimates should not be
36	used by themselves to answer the second question posed above. The extent of these non-
37	linearities will generally depend on location and time, the strength of concentrated

1	sources, and the nature of the chemical regime. Further work is needed on how these
2	estimates of background concentrations can be used to help determine the contributions
3	of background sources of O ₃ to urban concentrations.

2.2.3 Monitoring

4	The federal reference method (FRM) for O_3 measurement is based on the detection of
5	chemiluminescence resulting from the reaction of O_3 with ethylene gas. However, almost
6	all of the state and local air monitoring stations (SLAMS) that reported data to the EPA's
7	Air Quality System (AQS) database from 2005 to 2009 used the federal equivalence
8	method (FEM) UV absorption photometer. More than 96% of O_3 monitors met precision
9	and bias goals during this period.
10	In 2010, there were 1250 SLAMS O ₃ monitors reporting data to AQS. Ozone monitoring
11	is required at SLAMS sites during the local "ozone season" which varies by state. In
12	addition, National Core (NCore) is a new multipollutant monitoring network
13	implemented to meet multiple monitoring objectives and each state is required to operate
14	at least one NCore site. The NCore network consists of 60 urban and 20 rural sites
15	nationwide (See Figure 3-21 and Figure 3-22). The densest concentrations of O_3 sites are
16	located in California and the eastern half of the U.S.
17	The Clean Air Status and Trends Network (CASTNET) is a regional monitoring network
18	established to assess trends in acidic deposition and also provides concentration
19	measurements of O ₃ . CASTNET O ₃ monitors operate year round and are primarily
20	located in rural areas; in 2010, there were 80 CASTNET sites reporting O_3 data to AQS.
21	The National Park Service (NPS) operates 23 CASTNET sites in national parks and other
22	Class-i areas, and provided data to AQS from 20 additional Portable Ozone Monitoring
23	Systems (POMS) in 2010 (See Figure 3-22). Compared to urban-focused monitors, rural-
24	focused monitors are relatively scarce across the U.S.

2.2.4 Ambient Concentrations

25Ozone is the only photochemical oxidant other than NO2 that is routinely monitored and26for which a comprehensive database exists. Other photochemical oxidants are typically27only measured during special field studies. The concentration analyses in Chapter 3 are28limited to widely available O3 data obtained directly from AQS for the period from 200729to 2009. The median 24-h average, 8-h daily maximum, and 1-h daily maximum O330concentrations across all U.S. sites reporting data to AQS between 2007 and 2009 were3129, 40, and 44 ppb, respectively.

1	To investigate O_3 variability in urban areas across the U.S., 20 combined statistical areas
2	(CSAs) were selected for closer analysis based on their importance in O ₃ epidemiology
3	studies and on their location. Several CSAs had relatively little spatial variability in
4	8-hour daily maximum O ₃ concentrations (e.g., inter-monitor correlations ranging from
5	0.61 to 0.96 in the Atlanta CSA) while other CSAs exhibited considerably more
6	variability in O ₃ concentrations (e.g., inter-monitor correlations ranging from -0.06 to
7	0.97 in the Los Angeles CSA). Uncertainties resulting from the observed variability in O_3
8	concentration fields should be considered when using data from the network of ambient
9	O ₃ monitors to approximate community-scale exposures.
10	To investigate O_3 variability in rural settings across the U.S., six focus areas were
11	selected for closer analysis based on the impact of O ₃ or O ₃ precursor transport from
12	upwind urban areas. The selected rural focus area with the largest number of available
13	AQS monitors was Great Smoky Mountains National Park where the May-September
14	median 8-h daily maximum O_3 concentration ranged from 47 ppb at the lowest elevation
15	(564 m) site to 60 ppb at the highest elevation (2,021 m) site. Correlations between sites
16	within each rural focus area ranged from 0.78 to 0.02 . Ozona in rural areas is produced

- within each rural focus area ranged from 0.78 to 0.92. Ozone in rural areas is produced 16 from emissions of O_3 precursors emitted directly within the rural areas, from emissions in 17 18 urban areas that are processed during transport, and from occasional stratospheric 19 intrusions. Factors contributing to variations observed within these rural focus areas 20 include proximity to local O₃ precursor emissions, local scale circulations related to 21 topography, and possibly stratospheric intrusions as a function of elevation. In addition, 22 O_3 tends to persist longer in rural than in urban areas as a result of less chemical 23 scavenging. This results in a more uniform O_3 concentration throughout the day and night 24 without the typical nocturnal decrease in O₃ concentration observed in urban areas. 25 Persistently high O₃ concentrations observed at many of the rural sites investigated here 26 indicate that cumulative exposures for humans and vegetation in rural areas can 27 frequently exceed cumulative exposures in urban areas.
- 28 Nation-wide surface level O_3 concentrations have declined over the last decade, with a 29 particularly noticeable decrease between 2003 and 2004 coinciding with NO_x emissions 30 reductions resulting from implementation of the NO_X SIP Call rule, which began in 2003 31 and was fully implemented in 2004. This rule was designed to reduce NO_x emissions 32 from power plants and other large combustion sources in the eastern U.S. The largest 33 density of individual monitors showing downward trends in O₃ concentrations over the 34 last decade occur in the Northeast where this rule was focused. In addition to a downward 35 trend, the nation-wide surface level O_3 concentration data also show a general tightening 36 of the distribution across sites. In contrast to the majority of U.S. surface level monitors 37 reporting downward trends, a few surface-level monitors and elevated observations along 38 the Pacific Coast have shown increases in O_3 concentrations in recent years, possibly

1	resulting from intercontinental transport from Asia. As noted in the 2006 O ₃ AQCD,
2	trends in national parks and rural areas are similar to nearby urban areas, reflecting the
3	regional nature of O ₃ pollution.
4	Since O_3 is a secondary pollutant, it is not expected to be highly correlated with primary
5	pollutants such as CO and NO _X . Furthermore, O ₃ formation is strongly influenced by
6	meteorology, entrainment, and transport of both O3 and O3 precursors, resulting in a
7	broad range in correlations with other pollutants which can vary substantially with
8	season. Correlations between 8-h daily maximum O ₃ and other criteria pollutants exhibit
9	mostly negative correlations in the winter and mostly positive correlations in the summer.
10	The median seasonal correlations are modest at best with the highest positive correlation
11	at 0.52 for $PM_{2.5}$ in the summer and the highest negative correlation at -0.38 for $PM_{2.5}$ in
12	the winter. As a result, statistical analyses that may be sensitive to correlations between
13	copollutants need to take seasonality into consideration, especially when O ₃ is being
14	investigated.

2.3 Human Exposure

15	The widespread presence of O_3 in the environment results in exposure as people
16	participate in normal daily activities. Personal exposure measurements have been found
17	to be moderately associated with fixed-site ambient O ₃ concentrations, although a number
18	of factors affect the relationship between ambient concentration and personal exposure.
19	These include: infiltration of ambient O ₃ into indoor microenvironments, which is driven
20	by air exchange rate; time spent outdoors and activity pattern, which includes changes in
21	personal behavior by some populations to avoid exposure to O_3 ; and the variation in O_3
22	concentrations at various spatial and temporal scales. Personal exposure to O ₃ is
23	moderately correlated with ambient O ₃ concentration, as indicated by studies reporting
24	correlations generally in the range of 0.3-0.8 (Table 4-2). This suggests that ambient
25	monitor concentrations are representative of day-to-day changes in personal exposure to
26	ambient O3. Some studies report lower personal-ambient correlations, a result attributable
27	in part to low building air exchange rates and O_3 concentrations below the personal
28	sampler detection limit. Low correlations may also occur for individuals or populations
29	spending increased time indoors. In contrast to correlation, which represents the temporal
30	association between exposure and concentration, the magnitude of exposure can be
31	represented as the ratio between personal exposure and ambient concentration. This ratio
32	varies widely depending on activity patterns, housing characteristics, and season.
33	Personal-ambient ratios are typically 0.1-0.3 for sampling durations of several hours to
34	several days, although individuals spending substantial time outdoors (e.g., outdoor
35	workers) have shown much higher ratios (0.5-0.9) (<u>Table 4-3</u>). Since there are relatively

- 1few indoor sources of O3, and because of reactions of O3 with indoor surfaces and2airborne constituents, indoor O3 concentrations are often substantially lower than outdoor3concentrations (Section 4.3.2). The lack of indoor sources also suggests that fluctuations4in ambient O3 may be primarily responsible for changes in personal exposure, even under5low-ventilation, low-concentration conditions.
- 6 Another factor that may influence the pattern of exposure is the tendency for people to 7 avoid O_3 exposure by altering their behavior (e.g., reducing outdoor activity levels or 8 time spent being active outdoors) on high-O₃ days. Activity pattern has a substantial 9 effect on ambient O₃ exposure, with time spent outdoors contributing to increased 10 exposure (Section 4.4.2). Air quality alerts and public health recommendations induce reductions in time spent outdoors on high-O₃ days among some populations, particularly 11 12 for children, older adults, and people with respiratory problems. Such effects are less 13 pronounced in the general population. Limited evidence from an epidemiologic study 14 conducted in the 1990's in Los Angeles, CA reports increased asthma hospital 15 admissions among children and older adults when O₃ alert days (1-h max O₃ 16 concentration >200 ppb) were excluded from the analysis of daily hospital admissions 17 and O_3 concentrations (presumably thereby eliminating averting behavior based on high 18 O₃ forecasts). The lower rate of admissions observed when alert days were included in 19 the analysis suggests that estimates of health effects based on concentration-response 20 functions that do not account for averting behavior may be biased towards the null.
- 21 Variations in O₃ concentrations occur over multiple spatial and temporal scales. Near 22 roadways, O₃ concentrations are reduced due to reaction with NO and other species 23 (Section 4.3.4.2). Over spatial scales of a few kilometers and away from roads, O_3 may 24 be somewhat more homogeneous due to its formation as a secondary pollutant, while 25 over scales of tens of kilometers, additional atmospheric processing can result in higher 26 concentrations downwind of an urban area. Although local-scale variability impacts the 27 magnitude of O_3 concentrations, O_3 formation rates are influenced by factors that vary 28 over larger spatial scales, such as temperature (Section 3.2), suggesting that urban 29 monitors may track one another temporally, but miss small-scale variability. This 30 variation in concentrations changes the pattern of exposure people experience as they 31 move through different microenvironments and affects the magnitude of exposures in 32 different locations within an urban area. The various factors affecting exposure patterns 33 and quantification of exposure result in uncertainty which may contribute to exposure 34 measurement error in epidemiologic studies, which typically use fixed-site monitor data 35 as an indicator of exposure. Low personal-ambient correlations are a source of exposure 36 error for epidemiologic studies, tending to obscure the presence of potential thresholds, 37 bias effect estimates toward the null, and widen confidence intervals, and this impact may 38 be more pronounced among populations spending substantial time indoors. The impact of

1	this exposure error may tend more toward widening confidence intervals than biasing
2	effect estimates, since epidemiologic studies evaluating the influence of monitor selection
3	indicate that effect estimates are similar across different spatial averaging scales and
4	monitoring sites. In addition, in examinations of respiratory endpoints in epidemiologic
5	studies, associations were similar in magnitude across analyses using several different
6	exposure assessment methods that likely vary in how well ambient O_3 concentrations
7	represent personal exposures and between-subject variability in exposures. Respiratory
8	effects were observed with ambient O3 concentrations found to have stronger personal-
9	ambient relationships, including those measured on-site during long periods of outdoor
10	activity. However, such effects were also found with ambient O3 measurements expected
11	to have weaker personal-ambient relationships, including those measured at home or
12	school, measured at the closest site, averaged from multiple community sites, and
13	measured at a single site. Overall, there was no clear indication that a particular method
14	of exposure assessment produced stronger findings.

2.4 Dosimetry and Mode of Action

15	Upon inspiration, O_3 uptake in the respiratory tract is affected by a number of factors
16	including respiratory tract morphology, and breathing route, frequency, and volume.
17	Additionally, physicochemical properties of O_3 itself and how it is transported, as well as
18	the physical and chemical properties of the extracellular lining fluid (ELF) and tissue
19	layers in the respiratory tract can influence O3 uptake. Experimental studies and models
20	have suggested that there are differences between the total absorption of O_3 from the
21	inhaled air and the O_3 dose reaching the respiratory tract tissues. The total O_3 absorption
22	gradually decreases with distal progression into the respiratory tract. In contrast, the
23	primary site of O ₃ delivery to the lung epithelium is believed to be the centriacinar region
24	or the junction of the conducting airways with the gas exchange region.
25	Ozone uptake is sensitive to a number of factors including tidal volume, breathing
25 26	Ozone uptake is sensitive to a number of factors including tidal volume, breathing frequency, O_3 concentration, and exposure time. Interindividual variability also accounts
26	frequency, O ₃ concentration, and exposure time. Interindividual variability also accounts
26 27	frequency, O_3 concentration, and exposure time. Interindividual variability also accounts for a large amount of the variability in local dose due to differences in pulmonary
26 27 28	frequency, O_3 concentration, and exposure time. Interindividual variability also accounts for a large amount of the variability in local dose due to differences in pulmonary physiology, anatomy, and biochemistry. An increase in tidal volume and breathing
26 27 28 29	frequency, O_3 concentration, and exposure time. Interindividual variability also accounts for a large amount of the variability in local dose due to differences in pulmonary physiology, anatomy, and biochemistry. An increase in tidal volume and breathing frequency are both associated with increased physical activity. These changes and a
26 27 28 29 30	frequency, O_3 concentration, and exposure time. Interindividual variability also accounts for a large amount of the variability in local dose due to differences in pulmonary physiology, anatomy, and biochemistry. An increase in tidal volume and breathing frequency are both associated with increased physical activity. These changes and a switch to oronasal breathing during exercise result in deeper penetration of O_3 into the
26 27 28 29 30 31	frequency, O_3 concentration, and exposure time. Interindividual variability also accounts for a large amount of the variability in local dose due to differences in pulmonary physiology, anatomy, and biochemistry. An increase in tidal volume and breathing frequency are both associated with increased physical activity. These changes and a switch to oronasal breathing during exercise result in deeper penetration of O_3 into the lower respiratory tract in part due to less oral versus nasal uptake efficiency. For these

1	The ELE is a complex minture of livids, matrice, and entionidents that some as the first
1	The ELF is a complex mixture of lipids, proteins, and antioxidants that serves as the first harries and target for inhold O_{1} (see Figure 5.7). Distinct and had which diverge machinity
2 3	barrier and target for inhaled O_3 (see Figure 5-7). Distinct products with diverse reactivity
	(i.e., secondary oxidation products), are mainly formed by reactions of O_3 with soluble
4	ELF components. The thickness of the ELF and that of the mucus layer, within the ELF,
5	are important determinants of the dose of O_3 to the tissues; a thicker ELF generally
6	results in a lower dose of O_3 to the tissues. Additionally, the quenching ability and the
7	concentrations of antioxidants and other ELF components are determinants of the
8	formation of secondary oxidation products. These reactions appear to limit interaction of
9	O_3 with underlying tissues and to reduce penetration of O_3 distally into the respiratory
10	tract.
11	In addition to contributing to the driving force for O_3 uptake, formation of secondary
12	oxidation products contributes to oxidative stress which may lead to cellular injury and
13	altered cell signaling in the respiratory tract. Secondary oxidation products initiate
14	pathways (See Figure 5-8) that provide the mechanistic basis for short- and long-term
15	health effects described in detail in Chapters 6 and 7. Other key events involved in the
16	mode of action of O_3 in the respiratory tract include the activation of neural reflexes,
17	initiation of inflammation, alterations of epithelial barrier function, sensitization of
18	bronchial smooth muscle, modification of innate and adaptive immunity, and airways
19	remodeling. Another key event, systemic inflammation and vascular oxidative/nitrosative
20	stress, may be critical to the extrapulmonary effects of O_3 .
21	Secondary oxidation products can transmit signals to respiratory tract cells resulting in
22	the activation of neural reflexes. Nociceptive sensory nerves mediate the involuntary
23	truncation of respiration, resulting in decreases in lung function (i.e., FVC, FEV ₁ , and
24	tidal volume), and pain upon deep inspiration. Studies implicate TRPA1 receptors on
25	bronchial C-fibers in this reflex. Another neural reflex involves vagal sensory nerves,
26	which mediate a mild increase in airways obstruction (i.e., bronchoconstriction)
27	following exposure to O ₃ via parasympathetic pathways. Substance P release from
28	bronchial C-fibers and the SP-NK receptor pathway may also contribute to this response.
29	Secondary oxidation products also initiate the inflammatory cascade following exposure
30	to O_3 . Studies have implicated eicosanoids, chemokines and cytokines, vascular
31	endothelial adhesion molecules, and tachykinins in mediating this response.
32	Inflammation is characterized by airways neutrophilia as well as the influx of other
33	inflammatory cell types. Recent studies demonstrate a later phase of inflammation
34	characterized by increased numbers of macrophages, which is mediated by hyaluronan.
35	Inflammation further contributes to O_3 -induced oxidative stress.
55	inflation further contributes to 03-induced oxidative success.
36	Alteration of the epithelial barrier function of the respiratory tract also occurs as a result
37	of O ₃ -induced secondary oxidation product formation. Increased epithelial permeability

1 2 3 4 5 6	may lead to enhanced sensitization of bronchial smooth muscle, resulting in airways hyperresponsiveness (AHR). Neurally-mediated sensitization also occurs and is mediated by cholinergic postganglionic pathways and bronchial C-fiber release of substance P. Recent studies implicate hyaluronan and Toll-like receptor 4 (TLR4) signaling in bronchial smooth muscle sensitization, while earlier studies demonstrate roles for eicosanoids, cytokines, and chemokines.
7 8 9 10 11 12	Evidence is accumulating that exposure to O ₃ modifies innate and adaptive immunity through effects on macrophages, monocytes, and dendritic cells. Enhanced antigen presentation, adjuvant activity, and altered responses to endotoxin have been demonstrated. TLR4 signaling contributes to some of these responses. Effects on innate and adaptive immunity may result in both short- and longer-term consequences related to the exacerbation and/or induction of asthma and to alterations in host defense.
13 14 15 16 17	Airways remodeling has been demonstrated following chronic and/or intermittent exposure to O_3 by mechanisms that are not well understood. However, the TGF- β signaling pathway has recently been implicated in O_3 -induced deposition of collagen in the airways wall. These studies were conducted in adult animal models and their relevance to effects in humans is unknown.
18 19 20 21 22	Evidence is also accumulating that O_3 exposure results in systemic inflammation and vascular oxidative/nitrosative stress. The release of diffusible mediators from the O_3 -exposed lung into the circulation may initiate or propagate inflammatory responses in the vascular or in systemic compartments. This may provide a mechanistic basis for extrapulmonary effects, such as vascular dysfunction.
23 24 25 26 27 28 29	Both dosimetric and mechanistic factors contribute to the understanding of inter-individual variability in response. Inter-individual variability is influenced by variability in respiratory tract volume and thus surface area, breathing route, certain genetic polymorphisms, pre-existing conditions and disease, nutritional status, lifestages, attenuation, and co-exposures. In particular, very young individuals may be sensitive to developmental effects of O_3 since studies in animal models demonstrated altered development of lung and immune system.
30 31 32 33 34 35 36	Some of these factors are also influential in understanding species homology and sensitivity. Qualitatively, animal models exhibit a similar pattern of tissue dose distribution for O_3 with the largest tissue dose delivered to the centriacinar region. However, due to anatomical and biochemical respiratory tract differences, the actual O_3 dose delivered differs between humans and animal models. Animal data obtained in resting conditions underestimates the dose to the respiratory tract tissue relative to exercising humans. Further, it should be noted that, with the exception of airways

1remodeling, the mechanistic pathways discussed above have been demonstrated in both2animals and human subjects in response to the inhalation of O3. Even though interspecies3differences limit quantitative comparison between species, the short- and long-term4functional responses of laboratory animals to O3 appear qualitatively homologous to5those of the human making them a useful tool in determining mechanistic and6cause-effect relationships with O3 exposure. Furthermore, animal studies add to a better7understanding of the full range of potential O3-mediated effects.

2.5 Integration of Ozone Health Effects

8 This section evaluates the evidence from toxicological, controlled hum	an exposure, and
9 epidemiologic studies (which examined the health effects associated w	ith short- and
10 long-term exposure to O ₃ ,) and summarizes the main conclusions of th	is assessment
11 regarding the health effects of O ₃ and the concentrations at which those	e effects are
12 observed. The results from the health studies, supported by the synthes	sis of atmospheric
13 chemistry (See Section <u>2.2</u>) and exposure assessment (See Section <u>2.3</u>)) studies, contribute
14 to the causal determinations made for the health outcomes discussed in	1 this assessment
15 (See Preamble to this document for details on the causal framework).	

2.5.1 Conclusions from Previous Ozone AQCDs

16	The 2006 O_3 AQCD concluded that there was clear, consistent evidence of a causal
17	relationship between short-term O_3 exposure and respiratory health effects (U.S. EPA,
18	2006b). The causal relationship for respiratory health effects was substantiated by the
19	coherence of effects observed across controlled human exposure, epidemiologic, and
20	toxicological studies indicating effects of short-term O ₃ exposures on a range of
21	respiratory health endpoints from respiratory tract inflammation to respiratory-related
22	emergency department (ED) visits and hospital admissions.
23	Across disciplines, short-term O_3 exposures induced or were associated with statistically
24	significant declines in lung function. An equally strong body of evidence from controlled
24	significant decimes in fung function. An equany strong body of evidence from controlled
25	human exposure and toxicological studies demonstrated O ₃ -induced inflammatory
26	responses, increased epithelial permeability, and airway hyperresponsiveness (both
27	specific and nonspecific). Toxicological studies provided additional evidence for
28	O ₃ -induced impairment of host defenses. Combined, these findings from experimental
29	studies provided support for epidemiologic evidence, in which short-term increases in
30	ambient O ₃ concentration were consistently associated with increases in respiratory
31	symptoms and asthma medication use in children with asthma, respiratory-related

1 2 3	hospital admissions, and asthma-related ED visits. Although O ₃ was consistently associated with nonaccidental and cardiopulmonary mortality, the contribution of respiratory causes to these findings was uncertain.
4	Collectively, there is a vast amount of evidence spanning several decades that
5	demonstrated that exposure to O ₃ induces a range of respiratory effects. The majority of
6	this evidence was derived from studies investigating short-term exposure (i.e., hours to
7	weeks) to O ₃ . The combined evidence across disciplines led to the causal relationship
8	between short-term O_3 exposure and respiratory effects reported in the 2006 O_3 AQCD.
9	Mechanistic evidence for the effects of O ₃ on the respiratory system was characterized in
10	the 1996 O ₃ AQCD, which identified O ₃ -induced changes in a variety of lung lipid
11	species whose numerous biologically active metabolites, in turn, can affect host defenses,
12	lung function, and the immune system. As summarized in Section 2.4 and fully
13	characterized in Chapter 5 , key events in the toxicity pathway of O ₃ have been identified
14	in humans and animal models. They include formation of secondary oxidation products,
15	activation of neural reflexes, initiation of inflammation, alteration of epithelial barrier
16	function, sensitization of bronchial smooth muscle, modification of innate/adaptive
17	immunity, airways remodeling, and systemic inflammation and oxidative/nitrosative
18	stress.

2.5.2 Summary of Causal Determinations

19	Recent studies support or build upon the strong body of evidence presented in the 1996
20	and 2006 O_3 AQCDs that short-term O_3 exposure is causally associated with respiratory
21	health effects. Recent controlled human exposure studies demonstrate statistically
22	significant group mean decreases in pulmonary function to exposures as low as
23	60-70 ppb O_3 in young, healthy adults, and are supported by the strong, cumulative
24	evidence from epidemiologic studies. Equally strong evidence demonstrated associations
25	of ambient O ₃ with respiratory hospital admissions and ED visits across the U.S., Europe,
26	and Canada. Most effect estimates ranged from a 1.6 to 5.4% increase in daily all
27	respiratory-related ED visits or hospital admissions in all-year analyses for unit increases ¹
28	in ambient O ₃ concentrations. Several multicity studies and a multicontinent study
29	reported associations between short-term increases in ambient O ₃ concentrations and
30	increases in respiratory mortality. This evidence is supported by a large body of
31	individual-level epidemiologic panel studies that demonstrate associations of ambient O ₃
32	with respiratory symptoms in children with asthma. Further support is provided by recent
33	studies that found O3-associated increases in indicators of airway inflammation and

¹ Effect estimates were standardized to a 40-, 30-, and 20-ppb unit increase for 1-h max, 8-h max, and 24-h avg O₃.

1 oxidative stress in children with asthma. Across respiratory endpoints, evidence indicates 2 antioxidant capacity may modify the risk of respiratory morbidity associated with O_3 3 exposure. The potentially elevated risk of populations with diminished antioxidant 4 capacity and the reduced risk of populations with enhanced antioxidant capacity 5 identified in epidemiologic studies is strongly supported by similar findings from 6 controlled human exposure studies and by evidence that characterizes O_3 -induced 7 decreases in intracellular antioxidant levels as a mode of action for downstream effects. 8 By demonstrating O₃-induced airway hyperresponsiveness, decreased pulmonary 9 function, allergic responses, lung injury, impaired host defense, and airway inflammation, 10 toxicological studies have characterized O₃ modes of action and provided biological 11 plausibility for epidemiologic associations of ambient O₃ concentrations with lung 12 function and respiratory symptoms, hospital admissions, ED visits, and mortality. 13 Together, the evidence integrated across controlled human exposure, epidemiologic, and 14 toxicological studies and across the spectrum of respiratory health endpoints continues to 15 demonstrate that there is a causal relationship between short-term O₃ exposure and 16 respiratory health effects.

17 The strongest evidence for a relationship between long-term O_3 exposure and respiratory 18 health effects (including respiratory symptoms, new-onset asthma, and repiratory 19 mortality) is contributed by recent studies that demonstrated associations between long-20 term measures of O₃ exposure and both new-onset asthma in children and increased 21 respiratory symptom effects in individuals with asthma. While the evidence is limited, a 22 U.S. multicommunity prospective cohort demonstrates that asthma risk is affected by 23 interactions among genetic variability, environmental O_3 exposure, and behavior. The 24 evidence relating new-onset asthma to long-term O₃ exposure is supported by 25 toxicological studies of asthma in monkeys. This nonhuman primate evidence of 26 O_3 -induced changes supports the biologic plausibility of long-term exposure to O_3 27 contributing to the effects of asthma in children. Early life O_3 exposure may alter airway 28 development and lead to the development of asthma. Other recent epidemiologic studies 29 provide coherent evidence for long-term O_3 exposure and respiratory effects such as first 30 asthma hospitalization, respiratory symptoms in asthmatics, and respiratory mortality. 31 Generally, the epidemiologic and toxicological evidence provides a compelling case that 32 supports the hypothesis that a relationship exists between long-term exposure to ambient 33 O_3 and measures of respiratory health effects and mortality. The evidence for short-term 34 exposure to O_3 and effects on respiratory endpoints provides coherence and biological 35 plausibility for the effects of long-term exposure to O_3 . Building upon that evidence, the 36 more recent epidemiologic evidence, combined with toxicological studies in rodents and 37 nonhuman primates, provides biologically plausible evidence that there is likely to be a 38 causal relationship between long-term exposure to O₃ and respiratory health 39 effects.

2that short-term exposure to O3 directly or indirectly contributes to nonaccidental and3cardiopulmonary-related mortality, but additional research was needed to more fully4establish underlying mechanisms by which such effects occur. The evaluation of recent5multicity studies and a multicontinent study that examined the association between <i>short-</i> 6 <i>term increases in ambient O3 concentration and mortality</i> found evidence that supports7the conclusions of the 2006 O3 AQCD. These recent studies reported consistent positive8associations between short-term increases in ambient O3 concentration and total9(nonaccidental) mortality, with associations being stronger during the warm season, as10well as provided additional support for associations between O3 concentrations and11cardiovascular mortality being similar or larger in magnitude compared to respiratory12mortality. Additionally, these new studies examined previously identified areas of13uncertainty in the O3-mortality relationship, and provide additional evidence supporting14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	1	The 2006 O ₃ AQCD concluded that the overall body of evidence was highly suggestive
 establish underlying mechanisms by which such effects occur. The evaluation of recent multicity studies and a multicontinent study that examined the association between <i>short-</i> <i>term increases in ambient O₃ concentration and mortality</i> found evidence that supports the conclusions of the 2006 O₃ AQCD. These recent studies reported consistent positive associations between short-term increases in ambient O₃ concentration and total (nonaccidental) mortality, with associations being stronger during the warm season, as well as provided additional support for associations between O₃ concentrations and cardiovascular mortality being similar or larger in magnitude compared to respiratory mortality. Additionally, these new studies examined previously identified areas of uncertainty in the O₃-mortality relationship, and provide additional evidence supporting an association between short-term O₃ exposure and mortality. Taken together, the body of evidence indicates that there is likely to be a causal relationship between short-term 	2	that short-term exposure to O_3 directly or indirectly contributes to nonaccidental and
5 multicity studies and a multicontinent study that examined the association between <i>short-</i> 6 <i>term increases in ambient</i> O_3 <i>concentration and mortality</i> found evidence that supports 7 the conclusions of the 2006 O_3 AQCD. These recent studies reported consistent positive 8 associations between short-term increases in ambient O_3 concentration and total 9 (nonaccidental) mortality, with associations being stronger during the warm season, as 10 well as provided additional support for associations between O_3 concentrations and 11 cardiovascular mortality being similar or larger in magnitude compared to respiratory 12 mortality. Additionally, these new studies examined previously identified areas of 13 uncertainty in the O_3 -mortality relationship, and provide additional evidence supporting 14 an association between short-term O_3 exposure and mortality. Taken together, the body of 15 evidence indicates that there is likely to be a causal relationship between short-term	3	cardiopulmonary-related mortality, but additional research was needed to more fully
$\begin{array}{ccc} & & term increases in ambient O_3 \ concentration \ and \ mortality \ found \ evidence \ that \ supports \ the \ conclusions \ of \ the \ 2006 \ O_3 \ AQCD. \ These \ recent \ studies \ reported \ consistent \ positive \ associations \ between \ short-term \ increases \ in \ ambient \ O_3 \ concentration \ and \ total \ (nonaccidental) \ mortality, \ with \ associations \ being \ stronger \ during \ the \ warm \ season, \ as \ well \ as \ provided \ additional \ support \ for \ associations \ between \ O_3 \ concentrations \ and \ total \ (nonaccidental) \ mortality \ with \ associations \ between \ O_3 \ concentrations \ and \ total \ (nonaccidental) \ mortality, \ with \ associations \ between \ O_3 \ concentrations \ and \ total \ (nonaccidental) \ mortality \ being \ similar \ or \ larger \ in \ magnitude \ compared \ to \ respiratory \ mortality. \ Additionally, \ these \ new \ studies \ examined \ previously \ identified \ areas \ of \ uncertainty \ in \ the \ O_3 \ mortality \ relationship, \ and \ provide \ additional \ evidence \ supporting \ an \ association \ between \ short-term \ O_3 \ exposure \ and \ mortality. \ Taken \ together, \ the \ body \ of \ evidence \ indicates \ that \ there \ is \ likely \ to \ be \ a \ causal \ relationship \ between \ short-term \$	4	establish underlying mechanisms by which such effects occur. The evaluation of recent
7the conclusions of the 2006 O3 AQCD. These recent studies reported consistent positive8associations between short-term increases in ambient O3 concentration and total9(nonaccidental) mortality, with associations being stronger during the warm season, as10well as provided additional support for associations between O3 concentrations and11cardiovascular mortality being similar or larger in magnitude compared to respiratory12mortality. Additionally, these new studies examined previously identified areas of13uncertainty in the O3-mortality relationship, and provide additional evidence supporting14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	5	multicity studies and a multicontinent study that examined the association between short-
8 associations between short-term increases in ambient O ₃ concentration and total 9 (nonaccidental) mortality, with associations being stronger during the warm season, as 10 well as provided additional support for associations between O ₃ concentrations and 11 cardiovascular mortality being similar or larger in magnitude compared to respiratory 12 mortality. Additionally, these new studies examined previously identified areas of 13 uncertainty in the O ₃ -mortality relationship, and provide additional evidence supporting 14 an association between short-term O ₃ exposure and mortality. Taken together, the body of 15 evidence indicates that there is likely to be a causal relationship between short-term	6	term increases in ambient O_3 concentration and mortality found evidence that supports
 9 (nonaccidental) mortality, with associations being stronger during the warm season, as 10 well as provided additional support for associations between O₃ concentrations and 11 cardiovascular mortality being similar or larger in magnitude compared to respiratory 12 mortality. Additionally, these new studies examined previously identified areas of 13 uncertainty in the O₃-mortality relationship, and provide additional evidence supporting 14 an association between short-term O₃ exposure and mortality. Taken together, the body of 15 evidence indicates that there is likely to be a causal relationship between short-term 	7	the conclusions of the 2006 O ₃ AQCD. These recent studies reported consistent positive
10well as provided additional support for associations between O3 concentrations and11cardiovascular mortality being similar or larger in magnitude compared to respiratory12mortality. Additionally, these new studies examined previously identified areas of13uncertainty in the O3-mortality relationship, and provide additional evidence supporting14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	8	associations between short-term increases in ambient O3 concentration and total
11cardiovascular mortality being similar or larger in magnitude compared to respiratory12mortality. Additionally, these new studies examined previously identified areas of13uncertainty in the O3-mortality relationship, and provide additional evidence supporting14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	9	(nonaccidental) mortality, with associations being stronger during the warm season, as
12mortality. Additionally, these new studies examined previously identified areas of13uncertainty in the O3-mortality relationship, and provide additional evidence supporting14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	10	well as provided additional support for associations between O_3 concentrations and
 uncertainty in the O₃-mortality relationship, and provide additional evidence supporting an association between short-term O₃ exposure and mortality. Taken together, the body of evidence indicates that there is likely to be a causal relationship between short-term 	11	cardiovascular mortality being similar or larger in magnitude compared to respiratory
14an association between short-term O3 exposure and mortality. Taken together, the body of15evidence indicates that there is likely to be a causal relationship between short-term	12	mortality. Additionally, these new studies examined previously identified areas of
15 evidence indicates that there is likely to be a causal relationship between short-term	13	uncertainty in the O ₃ -mortality relationship, and provide additional evidence supporting
	14	an association between short-term O_3 exposure and mortality. Taken together, the body of
	15	evidence indicates that there is likely to be a causal relationship between short-term
16 O_3 exposures and total mortality.	16	O_3 exposures and total mortality.

- 17 The 2006 O₃ AQCD concluded that an insufficient amount of evidence existed to suggest 18 a causal relationship between long-term O_3 exposure and mortality (U.S. EPA, 2006b). 19 A synthesis of the recent and earlier evidence reveals that the strongest evidence for an 20 association between long-term exposure to ambient O_3 concentrations and mortality is 21 derived from associations for respiratory mortality that remained robust after adjusting 22 for PM_{2.5} concentrations. There is inconsistent evidence for an association between long-23 term exposure to ambient O_3 and cardiopulomary mortality, with several analyses from 24 the ACS cohort reporting some positive associations, while other studies reported no 25 association. There is generally limited evidence for an association with long-term 26 exposure to ambient O_3 and total mortality. The findings for respiratory mortality are 27 consistent and coherent with the evidence from epidemiologic, controlled human 28 exposure, and animal toxicological studies for the effects of short- and long-term 29 exposure to O_3 on respiratory effects. Respiratory mortality is a relatively small portion 30 of total mortality [about 7.6% of all deaths in 2010 were due to respiratory causes 31 (Murphy et al., 2012)], thus it is not surprising that the respiratory mortality signal may 32 be difficult to detect in studies of cardiopulmonary or total mortality. Based on the recent 33 evidence for respiratory mortality along with limited evidence for total and 34 cardiopulmonary mortality, the evidence is suggestive of a causal relationship 35 between long-term O3 exposures and total mortality.
- In past O₃ AQCDs the effects of *short- and long-term exposure to O₃ on the cardiovascular system* could not be thoroughly evaluated due to the paucity of
 information available. However, studies investigating O₃-induced cardiovascular events

- 1 have advanced in the last two decades. Animal toxicological studies provide evidence for 2 both short- and long-term O_3 exposure leading to cardiovascular morbidity. The 3 toxicological studies demonstrate O_3 -induced cardiovascular effects, specifically 4 enhanced atherosclerosis and ischemia/reperfusion injury with or without the 5 corresponding development of a systemic oxidative, pro-inflammatory environment, 6 disrupted NO-induced vascular reactivity, decreased cardiac function, and increased heart 7 rate variability (HRV). The observed increase in HRV is supported by a recent controlled 8 human exposure study that also found increased high frequency HRV, but not altered 9 blood pressure, following O_3 exposure. It is still uncertain how O_3 inhalation may cause 10 systemic toxicity; however the cardiovascular effects of O₃ found in animals correspond 11 to the development and maintenance of an extrapulmonary oxidative, proinflammatory 12 environment that may result from pulmonary inflammation.
- 13 There is limited, inconsistent evidence for cardiovascular morbidity in epidemiologic 14 studies examining both short- and long-term exposure to O_3 . This is highlighted by the 15 multiple studies that examined the association between short-term increases in ambient 16 O₃ concentration and cardiovascular-related hospital admissions and ED visits and other 17 various cardiovascular effects and found no evidence of a consistent relationship with O_3 18 exposure. Positive associations between short-term increases in O₃ concentration and 19 cardiovascular mortality have been consistently reported in multiple epidemiologic 20 studies. However, the lack of coherence between the results from studies that examined 21 associations between short-term increases in O₃ concentration and cardiovascular 22 morbidity and subsequently cardiovascular mortality, complicate the interpretation of the 23 evidence for O₃-induced cardiovascular mortality.
- 24 Overall, animal toxicological studies provide some evidence for O₃-induced 25 cardiovascular effects, but the effects observed were not consistently supported by 26 controlled human exposure studies or epidemiologic studies. Although the toxicological 27 evidence provides initial support to the relatively strong body of evidence indicating 28 O₃-induced cardiovascular mortality, there is a lack of coherence with controlled human 29 exposure and epidemiologic studies of cardiovascular morbidity which together do not 30 support O_3 -induced cardiovascular effects. Thus, the overall body of evidence across 31 disciplines is suggestive of a causal relationship for both relevant short- and long-32 term exposures to O₃ and cardiovascular effects.
- 33In the 2006 O_3 AQCD, there were a number of health effects for which an insufficient34amount of evidence existed to adequately characterize the relationships with exposure to35 O_3 . However, recent evidence suggests that O_3 may impart health effects through36exposure durations and biological mechanisms not previously considered. For example,37recent toxicological studies add to earlier evidence that *short- and long-term exposures to*38 O_3 can produce a range of effects on the central nervous system and behavior.

1	Additionally, an epidemiologic study demonstrated that long-term exposure to O_3 affects
2	memory in humans as well. Together the evidence from studies of short- and long-term
3	exposure to O_3 is suggestive of a causal relationship between O_3 exposure and
4	central nervous system effects. There is also limited though positive toxicological
5	evidence for O_3 -induced developmental effects. Limited epidemiologic evidence exists
6	for an association of O ₃ concentration with decreased sperm concentration and
7	associations with reduced birth weight and restricted fetal growth. Overall, the evidence
8	is suggestive of a causal relationship between long-term exposures to O_3 and
9	reproductive and developmental effects.
10	These causal determinations are summarized in Table 2-2, along with the conclusions
11	from the previous NAAQS review. Special emphasis and additional details are provided
12	in Table 2-2 for respiratory health outcomes, for which there is the strongest body of
13	evidence.

Table 2-2Summary of evidence from epidemiologic, controlled human
exposure, and animal toxicological studies on the health effects
associated with short- and long-term exposure to ozone.

Health Outcome	Conclusions from 2006 O ₃ AQCD	Conclusions from 2012 3rd Draft ISA
Short-Term Exposure t	o O ₃	
Respiratory effects	The overall evidence supports a causal relationship between acute ambient O_3 exposures and increased respiratory morbidity outcomes.	Evidence integrated across controlled human exposure, epidemiologic, and toxicological studies and across the spectrum of respiratory health endpoints continues to demonstrate that there is a causal relationship between short-term O_3 exposure and respiratory health effects.
Lung function	Results from controlled human exposure studies and animal toxicological studies provide clear evidence of causality for the associations observed between acute (\leq 24 h) O ₃ exposure and relatively small, but statistically significant declines in lung function observed in numerous recent epidemiologic studies. Declines in lung function are particularly noted in children, asthmatics, and adults who work or exercise outdoors.	Recent controlled human exposure studies demonstrate group mean decreases in FEV ₁ in the range of 2 to 3% with 6.6 hour exposures to as low as 60 ppb O ₃ . The collective body of epidemiologic evidence demonstrates associations between short- term ambient O ₃ exposure and decrements in lung function, particularly in children with asthma, children, and adults who work or exercise outdoors.
Airway hyperresponsiveness	Evidence from human clinical and animal toxicological studies clearly indicate that acute exposure to O_3 can induce airway hyperreactivity, thus likely placing atopic asthmatics at greater risk for more prolonged bouts of breathing difficulties due to airway constriction in response to various airborne allergens or other triggering stimuli.	A limited number of studies have observed airway hyperresponsiveness in rodents and guinea pigs after exposure to less than 300 ppb O ₃ . As previously reported in the 2006 O ₃ AQCD, increased airway responsiveness has been demonstrated at 80 ppb in young, healthy adults, and at 50 ppb in certain strains of rats.

Health Outcome	Conclusions from 2006 O_3 AQCD	Conclusions from 2012 3rd Draft ISA
Pulmonary inflammation, injury and oxidative stress	The extensive human clinical and animal toxicological evidence, together with the limited available epidemiologic evidence, is clearly indicative of a causal role for O ₃ in inflammatory responses in the airways.	Epidemiologic studies provided new evidence for associations of ambient O_3 with mediators of airway inflammation and oxidative stress and indicate that higher antioxidant levels may reduce pulmonary inflammation associated with O_3 exposure. Generally, these studies had mean 8-h max O_3 concentrations less than 73 ppb . Recent controlled human exposure studies show O_3 -induced inflammatory responses at 60 ppb, the lowest concentration evaluated.
Respiratory symptoms and medication use	Young healthy adult subjects exposed in clinical studies to O_3 concentrations ≥ 80 ppb for 6 to 8 h during moderate exercise exhibit symptoms of cough and pain on deep inspiration. The epidemiologic evidence shows significant associations between acute exposure to ambient O_3 and increases in a wide variety of respiratory symptoms (e.g., cough, wheeze, production of phlegm, and shortness of breath) and medication use in asthmatic children.	The collective body of epidemiologic evidence demonstrates positive associations between short-term exposure to ambient O_3 and respiratory symptoms (e.g., cough, wheeze, and shortness of breath) in children with asthma. Generally, these studies had mean 8-h max O_3 concentrations less than 69 ppb .
Lung host defenses	Toxicological studies provided extensive evidence that acute O ₃ exposures as low as 80 to 500 ppb can cause increases in susceptibility to infectious diseases due to modulation of lung host defenses. A single controlled human exposure study found decrements in the ability of alveolar macrophages to phagocytize microorganisms upon exposure to 80 to 100 ppb O ₃ .	Recent controlled human exposure studies demonstrate the increased expression of cell surface markers and alterations in sputum leukocyte markers related to innate adaptive immunity with short-term O_3 exposures of 80-400 ppb . Recent studies demonstrating altered immune responses and natural killer cell function build on prior evidence that O_3 can affect multiple aspects of innate and acquired immunity with short- term O_3 exposures as low as 80 ppb .
Allergic and asthma related responses	Previous toxicological evidence indicated that O_3 exposure skews immune responses toward an allergic phenotype, and enhances the development and severity of asthma-related responses such as AHR.	Recent controlled human exposure studies demonstrate enhanced allergic cytokine production in atopic individuals and asthmatics, increased IgE receptors in atopic asthmatics, and enhanced markers of innate immunity and antigen presentation in health subjects or atopic asthmatics with short-term exposure to 80-400 ppb O ₃ , all of which may enhance allergy and/or asthma. Further evidence for O ₃ -induced allergic skewing is provided by a few recent studies in rodents using exposure concentrations as low as 200 ppb .
Respiratory Hospital admissions, ED visits, and physician visits	Aggregate population time-series studies observed that ambient O_3 concentrations are positively and robustly associated with respiratory-related hospitalizations and asthma ED visits during the warm season.	Consistent, positive associations of ambient O_3 with respiratory hospital admissions and ED visits in the U.S., Europe, and Canada with supporting evidence from single city studies. Generally, these studies had mean 8-h max O_3 concentrations less than 60 ppb .
Respiratory Mortality	Aggregate population time-series studies specifically examining mortality from respiratory causes were limited in number and showed inconsistent associations between acute exposure to ambient O ₃ exposure and respiratory mortality.	Recent multicity time-series studies and a multicontinent study consistently demonstrated associations between ambient O_3 and respiratory-related mortality visits across the U.S., Europe, and Canada with supporting evidence from single city studies. Generally, these studies had mean 8-h max O_3 concentrations less than 63 ppb .
Cardiovascular effects	The limited evidence is highly suggestive that O_3 directly and/or indirectly contributes to cardiovascular-related morbidity, but much remains to be done to more fully substantiate the association.	The overall body of evidence across disciplines is suggestive of a causal relationship for relevant short-term exposures to O_3 and cardiovascular effects.

Health Outcome	Conclusions from 2006 O ₃ AQCD	Conclusions from 2012 3rd Draft ISA
Central nervous system effects	Toxicological studies report that acute exposures to O_3 are associated with alterations in neurotransmitters, motor activity, short- and long-term memory, sleep patterns, and histological signs of neurodegeneration.	Together the evidence from studies of short-term exposure to O_3 is suggestive of a causal relationship between O_3 exposure and CNS effects.
Total Mortality	The evidence is highly suggestive that O_3 directly or indirectly contributes to non- accidental and cardiopulmonary-related mortality.	Taken together, the body of evidence indicates that there is likely to be a causal relationship between short-term exposures to O_3 and all-cause total mortality.
Long-term Exposure to (D ₃	
Respiratory effects	The current evidence is suggestive but inconclusive for respiratory health effects from long-term O_3 exposure.	Recent epidemiologic evidence, combined with toxicological studies in rodents and non-human primates, provides biologically plausible evidence that there is likely to be a causal relationship between long-term exposure to O_3 and respiratory health effects.
New onset asthma	No studies examining this outcome were evaluated in the 2006 O_3 AQCD.	Evidence that different genetic variants (HMOX, GST, ARG), in combination with O_3 exposure, are related to new onset asthma. These associations were observed when subjects living in areas where the mean annual 8-h max O_3 concentration was 55.2 ppb , compared to those who lived where it was 38.4 ppb .
Asthma hospital admissions	No studies examining this outcome were evaluated in the 2006 O_3 AQCD.	Chronic O_3 exposure was related to first childhood asthma hospital admissions in a positive concentration-response relationship. Generally, these studies had mean annual 8-h max O_3 concentrations less than 41 ppb .
Pulmonary structure and function	Epidemiologic studies observed that reduced lung function growth in children was associated with seasonal exposure to O_3 ; however, cohort studies of annual or multiyear O_3 exposure observed little clear evidence for impacts of longer-term, relatively low-level O_3 exposure on lung function development in children. Animal toxicological studies reported chronic O_3 -induced structural alterations, some of which were irreversible, in several regions of the respiratory tract including the centriacinar region. Morphologic evidence from studies using exposure regimens that mimic seasonal exposure patterns report increased lung injury compared to conventional chronic stable exposures.	Evidence for pulmonary function effects is inconclusive, with some new epidemiologic studies observing positive associations (mean annual 8-h max O ₃ concentrations less than 65 ppb). Information from toxicological studies indicates that long-term maternal exposure during gestation (100 ppb) or development (500 ppb) can result in irreversible morphological changes in the lung, which in turn can influence pulmonary function.
Pulmonary inflammation, injury and oxidative stress	Extensive human clinical and animal toxicological evidence, together with limited epidemiologic evidence available, suggests a causal role for O_3 in inflammatory responses in the airways.	Several epidemiologic studies (mean 8-h max O_3 concentrations less than 69 ppb) and toxicology studies (as low as 500 ppb) add to observations of O_3 -induced inflammation and injury.
Lung host defenses	Toxicological studies provided evidence that chronic O_3 exposure as low as 100 ppb can cause increases in susceptibility to infectious diseases due to modulation of lung host defenses, but do not cause greater effects on infectivity than short exposures.	Consistent with decrements in host defenses observed in rodents exposed to 100 ppb O ₃ , recen evidence demonstrates a decreased ability to respond to pathogenic signals in infant monkeys exposed to 500 ppb O ₃ .
Allergic responses	Limited epidemiologic evidence supported an association between ambient O_3 and allergic symptoms. Little if any information was available from toxicological studies.	Evidence relates positive outcomes of allergic response and O_3 exposure but with variable strength for the effect estimates; exposure to O_3 may increase total IgE in adult asthmatics. Allergic indicators in monkeys were increased by exposure to O_3 concentrations of 500 ppb .

Health Outcome	Conclusions from 2006 O_3 AQCD	Conclusions from 2012 3rd Draft ISA
Respiratory mortality	Studies of cardio-pulmonary mortality were insufficient to suggest a causal relationship between chronic O_3 exposure and increased risk for mortality in humans.	A single study demonstrated that exposure to O_3 (long-term mean O_3 less than 104 ppb) elevated the risk of death from respiratory causes and this effect was robust to the inclusion of PM _{2.5} .
Cardiovascular Effects	No studies examining this outcome were evaluated in the 2006 O_3 AQCD.	The overall body of evidence across disciplines is suggestive of a causal relationship for relevant long-term exposures to O ₃ and cardiovascular effects.
Reproductive and developmental effects	Limited evidence for a relationship between air pollution and birth-related health outcomes, including mortality, premature births, low birth weights, and birth defects, with little evidence being found for O_3 effects.	Overall, the evidence is suggestive of a causal relationship between long-term exposures to O_3 and reproductive and developmental effects.
Central nervous system effects	Toxicological studies reported that acute exposures to O_3 are associated with alterations in neurotransmitters, motor activity, short and long term memory, sleep patterns, and histological signs of neurodegeneration. Evidence regarding chronic exposure and neurobehavioral effects was not available.	Together the evidence from studies of long-term exposure to O_3 is suggestive of a causal relationship between O_3 exposure and CNS effects.
Cancer	Little evidence for a relationship between chronic O_3 exposure and increased risk of lung cancer.	Overall, the evidence is inadequate to determine if a causal relationship exists between ambient O_3 exposures and cancer.
Total Mortality	There is little evidence to suggest a causal relationship between chronic O ₃ exposure and increased risk for mortality in humans.	Collectively, the evidence is suggestive of a causal relationship between long-term O_3 exposures and total mortality.

2.5.3 Integrated Synthesis of Evidence for Health Effects

1	Building off that evaluated in previous O ₃ AQCDs, recent evidence demonstrates that O ₃
2	is associated with a broad range of respiratory effects, including altered development of
3	the respiratory tract. Recent animal toxicological studies of long-term exposure to O ₃
4	occurring throughout various lifestages in monkeys, beginning with prenatal and early
5	life exposures, provide novel evidence for effects on the development of the respiratory
6	system, including ultrastructural changes in bronchiole development, effects on the
7	developing immune system, and increased offspring airway hyper-reactivity
8	(Section $\underline{7.4.7}$). The strongest evidence for O ₃ -induced effects on the developing lung
9	comes from a series of experiments using infant rhesus monkeys episodically exposed to
10	500 ppb O_3 for approximately 5 months, starting at one month of age. Functional changes
11	in the conducting airways of infant rhesus monkeys exposed to either O_3 alone or O_3 +
12	antigen were accompanied by a number of cellular and morphological changes. In
13	addition to these functional and cellular changes, substantial structural changes in the
14	respiratory tract were observed. Importantly, the O3-induced structural pathway changes
15	persisted after recovery in filtered air for six months after cessation of the O ₃ exposures.
16	Exposure to O_3 has also been associated with similar types of alterations in pulmonary
17	structure, including airways remodeling and pulmonary injury and increased

permeability, in all adult laboratory animal species studied, from rats to monkeys (U.S. EPA, 1996a).

- 3 In addition to effects on the development and structure of the respiratory tract, there is 4 extensive evidence for the effects of short-term exposure to O_3 on pulmonary 5 inflammation and oxidative stress. Previous evidence from controlled human exposure 6 studies indicated that O₃ causes an inflammatory response in the lungs (U.S. EPA, 7 <u>1996a</u>). This inflammatory response to O_3 was detected after a single 1-h exposure with 8 exercise to O_3 concentrations of 300 ppb; the increased levels of some inflammatory cells 9 and mediators persisted for at least 18 hours. Toxicological studies provided additional 10 evidence for increases in permeability and inflammation in rabbits at levels as low as 11 100 ppb O₃. Evidence summarized in the 2006 O₃ AQCD demonstrated that 12 inflammatory responses were observed subsequent to 6.6 hours O_3 exposure to the lowest 13 tested level of 80 ppb in healthy human adults, while toxicological studies provided 14 extensive evidence that short-term $(1-3 \text{ hours}) O_3$ exposure in the range of 100-500 ppb 15 could cause lung inflammatory responses. The limited epidemiologic evidence reviewed 16 in the 2006 O₃ AQCD demonstrated an association between short-term increases in 17 ambient O₃ concentration and airways inflammation in children (1-h max O₃ of 18 approximately 100 ppb). Recent studies in animals and in vitro models described 19 inflammatory and injury responses mediated by Toll-like receptors (e.g., TLR4, TLR2), 20 receptors for TNF or IL-1, multiple signaling pathways (e.g., p38, JNK, NFkB, 21 MAPK/AP-1), and oxidative stress (Section 6.2.3.3). Recent epidemiologic studies 22 provide additional supporting evidence by demonstrating associations of ambient O_3 with 23 mediators of airways inflammation and oxidative stress.
- 24 The normal inflammatory response in lung tissue is part of host defense that aids in 25 removing microorganisms or particles that have reached the distal airways and alveolar 26 surface. The 1996 O₃ AQCD concluded that short-term exposure to elevated 27 concentrations of O₃ resulted in alterations in these host defense mechanisms in the 28 respiratory system. Specifically, toxicological studies of short-term exposures as low as 29 100 ppb O_3 for 2 hours were shown to decrease the ability of alveolar macrophages to 30 ingest particles, and short-term exposures as low as 80 ppb for 3 hours prevented mice 31 from resisting infection with streptococcal bacteria and resulted in infection-related 32 mortality. Similarly, alveolar macrophages removed from the lungs of human subjects 33 after 6.6 hours of exposure to 80 and 100 ppb O_3 had decreased ability to ingest 34 microorganisms, indicating some impairment of host defense capability. These altered 35 host defense mechanisms can lead to increased risk of respiratory infections, which can 36 often predispose individuals to developing asthma when occurring in early life. Despite 37 the strong toxicological evidence, in the limited body of epidemiologic evidence, ambient 38 O3 concentrations have not been consistently associated with hospital admissions or ED

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visits for respiratory infection, pneumonia, or influenza (Section 6.2.7.2 and Section 6.2.7.3).

- 3 The most commonly observed and strongest evidence for respiratory effects associated 4 with short-term exposure to O₃ is transient *decrements in pulmonary function*. Controlled 5 human exposure studies reviewed in previous assessments demonstrated O₃-induced 6 decrements in pulmonary function, characterized by alterations in lung volumes and flow 7 and airway resistance and responsiveness for multihour exposures (up to 8 hours) to O_3 8 concentrations as low as 80 ppb (U.S. EPA, 1996a). A series of mobile laboratory studies 9 of lung function and respiratory symptoms reported pulmonary function decrements at 10 mean ambient O₃ concentrations of 140 ppb in exercising healthy adolescents and 11 increased respiratory symptoms and pulmonary function decrements at 150 ppb in 12 heavily exercising athletes and at 170 ppb in lightly exercising healthy and asthmatic 13 subjects. Epidemiologic and animal toxicological evidence is coherent with the results of 14 the controlled human exposure studies, both indicating decrements in lung function upon 15 O₃ exposure. A combined statistical analysis of epidemiologic studies in children at 16 summer camp with particularly strong exposure assessment demonstrated decrements in 17 FEV_1 of 0.50 mL/ppb with an increase in previous hour O₃ concentration. For 18 preadolescent children exposed to 120 ppb ambient O_3 , this estimated volume decrease 19 corresponded to an average decrement of 2.4-3.0% in FEV₁. Key studies of lung function 20 (FEV₁) measured before and after well-defined outdoor exercise events in adults yielded 21 concentration-response slopes of 0.40 and 1.35 mL/ppb ambient O_3 after exposure for up 22 to 1 hour. Animal toxicological studies reported similar respiratory effects in rats at 23 exposures as low as 200 ppb O₃ for 3 hours. The 2006 O₃ AQCD characterized the 24 controlled human exposure and animal toxicological studies as providing clear evidence 25 of causality for the associations observed between short-term (≤ 24 hours) increases in O₃ 26 concentration and relatively small, but statistically significant declines in lung function 27 observed in numerous recent epidemiologic studies. In epidemiologic studies, declines in 28 lung function were particularly noted in children with and without asthma, and adults 29 who work or exercise outdoors. 30 Recent controlled human exposure studies examined lower concentration O₃ exposures
- 31 (40-80 ppb) and demonstrated that FEV₁, respiratory symptoms, and inflammatory 32 responses were affected by O₃ exposures of 6.6 hours as low as 60 to 70 ppb 33 (Section 6.2.1.1 and Section 6.2.3.1). These studies demonstrated average O_3 -induced 34 decreases in FEV₁ in the range of 2.8 to 3.6% with O_3 exposures to 60 ppb for 6.6 hours. 35 Further, in the controlled human exposure studies evaluating effects of 60 ppb O_3 , on 36 average, 10% of the exposed individuals experienced >10% FEV₁ decrements following 37 6.6 hours of exposure. Considerable intersubject variability has also been reported in 38 studies at higher exposure concentrations (\geq 70 ppb) with some subjects experiencing

1

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1	considerably greater decrements than average. Recent epidemiologic studies provide
2	greater insight into individual- and population-level factors that may increase for the risk
3	of O ₃ -associated respiratory morbidity. In addition to lung function decrements
4	consistently reported in healthy children at summer camp, O3-associated decreases in
5	lung function were consistently observed in epidemiologic studies that included
6	potentially at-risk populations (e.g., individuals with asthma with concurrent respiratory
7	infection, older adults with AHR or elevated body mass index, or groups with diminished
8	antioxidant capacity).

9 Exposure to O_3 may also result in *respiratory symptoms* (e.g., coughing, wheezing, shortness of breath). The 1996 O₃ AQCD identified an association between respiratory 10 11 symptoms and increasing ambient O_3 , particularly among children with asthma. In the 12 2006 O₃ AQCD, symptoms of cough and pain on deep inspiration were well documented 13 in young healthy adult subjects after exposure of ≥ 80 ppb O₃ for 6-8 hours during 14 moderate exercise. Limited data suggested an increase in respiratory symptoms down to 15 60 ppb. More recently, these effects have been observed at 70 ppb in healthy adults. 16 Controlled human exposure studies of healthy adults, have also reported an increased 17 incidence of cough with O_3 exposures as low as 120 ppb and 1-3 hours in duration with very heavy exercise. The controlled human exposure studies also demonstrated lesser 18 19 respiratory symptom responses in children and older adults relative to young healthy 20 adults. Cumulative epidemiologic evidence adds to the findings from controlled human 21 exposure studies for healthy adults by demonstrating the effects of ambient O_3 exposure 22 on respiratory symptoms in children with asthma. Increases in ambient O_3 concentration 23 were associated with a wide variety of respiratory symptoms (e.g., cough, wheeze, and 24 shortness of breath) in children with asthma. Epidemiologic studies also indicated that 25 short-term increases in O₃ concentration are likely associated with increased asthma 26 medication use in children with asthma. Additionally, epidemiologic studies provide 27 evidence for an association between long-term exposure to O_3 and respiratory symptoms 28 (Section 7.2.2).

29 Ozone exposure has been shown to result in both specific and non-specific airway 30 hyperresponsiveness (AHR). Increased AHR is an important consequence of exposure to 31 O_3 because its presence represents a change in airway smooth muscle reactivity and 32 implies that the airways are predisposed to narrowing on inhalation of a variety of stimuli 33 (e.g., specific allergens, SO₂, cold air). Specifically, short-term (2 or 3 hours) exposure to 34 250 or 400 ppb O₃ was found to cause increases in AHR in response to allergen 35 challenges among allergic asthmatic subjects who characteristically already had 36 somewhat increased AHR at baseline. Increased non-specific AHR has been 37 demonstrated in healthy young adults down to 80 ppb O₃ following 6.6 hours of exposure 38 during moderate exercise. While AHR has not been widely examined in epidemiologic

1	studies, findings for O ₃ -induced increases in AHR in controlled human exposure
2	(Section $6.2.2.1$) and toxicological (Section $6.2.2.2$) studies provide biological
3	plausibility for associations observed between increases in ambient O3 concentration and
4	increases in respiratory symptoms in subjects with asthma.

- 5 In addition to asthma exacerbations, recent epidemiologic evidence has indicated that 6 long-term ambient O_3 concentrations may contribute to new onset asthma (Section 7.2.1, 7 Table 7-2). The new epidemiologic evidence base consists of studies using a variety of 8 designs and analysis methods evaluating the relationship between long-term annual 9 measures of exposure to ambient O₃ and measures of respiratory morbidity. Studies from 10 two California cohorts have provided evidence for different variants in genes related to 11 oxidative or nitrosative stress (e.g., HMOX, GSTs, ARG) that, depending on community 12 long-term O_3 concentrations, are related to new onset asthma. This is the first time that 13 evidence has extended beyond the association of short-term exposure to O_3 and asthma 14 exacerbations to suggest that long-term exposure to O_3 may play a role in the 15 development of the disease and contribute to incident cases of asthma.
- 16 The frequency of *ED visits and hospital admissions* due to respiratory symptoms, asthma 17 exacerbations and other respiratory diseases is associated with short- and long-term 18 exposure to ambient O₃ concentrations. Summertime daily hospital admissions for 19 respiratory causes in various locations of eastern North America were consistently 20 associated with ambient concentrations of O_3 in studies reviewed in the 1996 O_3 AQCD. 21 This association remained even with examination of only concentrations below 120 ppb 22 O₃. The 2006 O₃ AQCD concluded that aggregate population time-series studies 23 demonstrate a positive and robust association between ambient O₃ concentrations and 24 respiratory-related hospitalizations and asthma ED visits during the warm season. Recent 25 epidemiologic time-series studies that include additional multicity studies and a 26 multicontinent study further demonstrate that short-term exposures to ambient O₃ 27 concentrations are consistently associated with increases in respiratory hospital 28 admissions and ED visits specifically during the warm/summer months across a range of 29 O_3 concentrations (Section 6.2.7). There is also recent evidence for an association 30 between respiratory hospital admissions and long-term exposure to O_3 (Section 7.2.2).
- 31Finally, O_3 exposure may contribute to *death from respiratory causes*. Recent evidence32from several multicity studies and a multicontinent study demonstrate consistent positive33associations between short-term exposure to ambient O_3 concentrations and increases in34respiratory mortality (Section <u>6.6.2.5</u>). Similarly, a study of long-term exposure to35ambient O_3 concentrations also demonstrated an association between O_3 and increases in36respiratory mortality (Section <u>7.7.1</u>). Evidence from these recent mortality studies is37consistent and coherent with the evidence from epidemiologic, controlled human

1exposure, and animal toxicological studies for the effects of short- and long-term2exposure to O3 on respiratory effects. Additionally, the evidence for respiratory morbidity3after short- and long-term exposure provides biological plausibility for mortality due to4respiratory disease.

- 5 There is similar evidence for a positive association between short-term exposure to O_3 6 and mortality. This evidence has been substantiated by single-city studies reviewed in the 7 2006 O₃ AQCD and recent multicity and multicontinent studies. When examining 8 mortality due to cardiovascular disease, epidemiologic studies consistently observe 9 positive associations with short-term exposure to O₃. Additionally, there is some 10 evidence for an association between long-term exposure to O₃ and mortality. However, 11 the association between long-term ambient O₃ concentrations and cardiovascular 12 mortality may be confounded by other pollutants as evident by a study of cardiovascular 13 mortality that reported no association after adjustment for $PM_{2.5}$ concentrations. The lack 14 of coherence between the results from studies that examined associations between short-15 and long-term O₃ concentrations and cardiovascular morbidity, and results from studies of 16 cardiovascular mortality, complicate the interpretation of the evidence for O_3 -induced 17 cardiovascular mortality.
- 18 Epidemiologic studies evaluating cardiovascular morbidity and short- and long-term 19 exposure to O_3 provide no consistent evidence for an association. This is highlighted by 20 the multiple studies that examined the association between short- and long-term O_3 21 concentrations and cardiovascular-related hospital admissions and ED visits and 22 cardiovascular disease-related biomarkers. Additionally, a single controlled human 23 exposure study reported no statistically significant O₃-induced differences in 24 electrocardiogram (ECG), heart rate, or blood pressure in normal or hypertensive subjects 25 (0.3 ppm for 3 h with intermittent exercise), however an overall increase in myocardial 26 work and impairment in pulmonary gas exchange was observed.
- 27 There is an emerging body of animal toxicological evidence suggesting that autonomic 28 nervous system alterations (in heart rate and/or heart rate variability) and 29 proinflammatory signals may mediate cardiovascular effects. Interactions of O₃ with ELF 30 components result in secondary oxidation products and inflammatory mediators that have 31 the potential to penetrate the epithelial barrier and to initiate toxic effects on the 32 cardiovascular system. Animal toxicological studies of long-term exposure to O₃ provide 33 evidence enhanced atherosclerosis and ischemia/reperfusion (I/R) injury, corresponding 34 with development of a systemic oxidative, proinflammatory environment.
- 35Overall, animal toxicological studies provide some evidence for O3-induced36cardiovascular effects, but the effects were not consistently supported by controlled37human exposure studies or epidemiologic evidence. Although the toxicological evidence

1	provides initial support to the body of evidence indicating an association between short-
2	term exposure to O_3 and cardiovascular mortality, there is a lack of coherence with
3	controlled human exposure and epidemiologic studies of cardiovascular morbidity.
4	Together, these findings are suggestive of O ₃ -induced cardiovascular effects.

2.5.4 Policy Relevant Considerations

2.5.4.1 Populations Potentially at Increased Risk

5 Studies were conducted to identify populations that are at increased risk for O₃-related 6 health effects. These studies have investigated factors that can cause populations to be at 7 increased risk for O₃-related health effects by conducting stratified epidemiologic 8 analyses; by examining individuals with an underlying health condition, genetic 9 polymorphism, or categorized by age, race, or sex in controlled human exposure studies; 10 or by developing animal models that mimic the pathophysiological conditions associated 11 with a health effect. These studies have identified a multitude of factors that could 12 potentially contribute to whether a population is at increased risk for O_3 -related health 13 effects.

14 The populations identified in Chapter 8 that were examined for their potential for 15 increased risk of O₃-related health effects are listed in Table 8-5 and are classified as 16 providing adequate, suggestive, inadequate, or no evidence of being an at-risk factor. The 17 factors that have adequate evidence to be classified as an at-risk factor for O₃-related 18 health effects are individuals with asthma, younger and older age groups, individuals with 19 reduced intake of certain nutrients (i.e., vitamins C and E), and outdoor workers, based 20 on consistency in findings across studies and evidence of coherence in results from 21 different scientific disciplines. Asthma as a factor affecting risk was supported by 22 controlled human exposure and toxicological studies, as well as some evidence from 23 epidemiologic studies. Generally, studies comparing age groups also reported greater 24 associations for respiratory hospital admissions and ED visits among children than for 25 adults. Biological plausibility for this increased risk is supported by toxicological and 26 controlled human exposure studies. Also, children have higher exposure and dose due to 27 increased time spent outdoors and ventilation rate, and childrens' respiratory systems are 28 also still undergrowing lung growth. Most studies comparing age groups reported greater 29 effects of short-term O₃ exposure on mortality among older adults, although studies of 30 other health outcomes had inconsistent findings regarding whether older adults were at 31 increased risk. Multiple epidemiologic, controlled human exposure, and toxicological 32 studies reported that diets lower in vitamins E and C are associated with increased risk of

1	O_3 -related health effects. Previous studies have shown that increased exposure to O_3 due
2	to outdoor work leads to increased risk of O3-related health effects and it is clear that
3	outdoor workers have higher exposures, and possibly greater internal doses, of O ₃ , which
4	may lead to increased risk of O_3 -related health effects.
5	Other potential factors [genetic variants (such as those in GSTM1, HMOX-1, NQO1, and
6	<i>TNF</i> - α), obesity, sex, and SES] provided some suggestive evidence of increased risk, but
7	further investigation is needed. Similarly, many factors had inadequate evidence to
8	determine if they increased the risk of O ₃ -related health effects, including
9	influenza/infection, COPD, CVD, diabetes, hyperthyroidism, smoking, race/ethnicity,
10	and air conditioning use.

2.5.4.2 Exposure Metrics in Epidemiologic Studies

11Some epidemiologic studies have conducted analyses between O3 concentration and12health effects (i.e., mortality, respiratory or cardiovascular) using various exposure13metrics (i.e., 1-h max, 8-h max, and 24-h avg). No studies of long-term exposure14(i.e., months to years) to O3 have compared the use of different exposure metrics on risk15estimation.

- 16 Among time-series studies, the limited evidence suggests comparable risk estimates
- 17across exposure metrics with some evidence for smaller O3 risk estimates when using a1824-hour average exposure metric. Several panel studies examined whether associations of
- 19 lung function and respiratory symptoms varied depending on the O₃ exposure metric
- 20 used. Although differences in effect estimates across exposure metrics were found within 21 some studies, collectively, there was no indication that the consistency or magnitude of 22 the observed association was stronger for a particular O₃ exposure metric. Comparisons 23 of lung function decrements among O_3 exposure metrics were similarly inconsistent in 24 populations without increased outdoor exposures. It is important to note in these studies, 25 the degree of exposure measurement error associated with use of central site ambient O_3 26 concentrations may vary among O_3 averaging times, depending on time spent outdoors. 27 Among studies that examined associations of multiple respiratory symptoms in children 28 with multiple O_3 exposure metrics, most did not find higher odds ratios for any particular
- exposure metric. Overall, the evidence from time-series and panel epidemiologic studies
 does not indicate that one exposure metric is more consistently or strongly associated
 with mortality or respiratory-related health effects.

2.5.4.3 Lag Structure in Epidemiologic Studies

1 2 3 4 5	Epidemiologic studies have attempted to identify the time-frame in which exposure to O_3 can impart a health effect. The time period between O_3 exposure and health effects can potentially be influenced by a multitude of factors, such as age or existence of pre-existing diseases. Different lag times have been evaluated for specific health outcomes.
6 7 8 9 10 11	The epidemiologic evidence evaluated in the 2006 O_3 AQCD indicated that one of the remaining uncertainties in characterizing the O_3 -mortality relationship was identifying the appropriate lag structure (e.g., single-day lags versus distributed lag model). An examination of lag times used in the epidemiologic studies evaluated in this assessment can provide further insight on the characterization of the relationship between O_3 exposure and morbidity and mortality outcomes from epidemiologic studies.
12 13 14 15 16 17	The majority of epidemiologic studies that focused on the association between short-term O_3 exposure and mortality (i.e., all-cause, respiratory and cardiovascular) examined the average of multiday lags with some studies examining single-day lags. Across a range of multiday lags (i.e., average of 0-1 to 0-6 days), the studies evaluated consistently demonstrate that the O_3 effects on mortality occur within a few days of exposure (Figure 6-28).
18 19 20 21 22 23 24 25 26 27 28	Epidemiologic studies of lung function, respiratory symptoms, and biological markers of airway inflammation and oxidative stress examined associations with single-day ambient O_3 concentrations (using various averaging times) lagged from 0 to 7 days as well as concentrations averaged over 2 to 19 days. Lags of 0 and 1 day ambient O_3 concentrations were associated with decreases in lung function and increases in respiratory symptoms, airway inflammation, and oxidative stress. Additionally, several studies found that multiday averages of O_3 concentration were associated with these endpoints, indicating that not only single day, but exposures accumulated over several days led to a respiratory health effect. In studies of respiratory hospital admissions and ED visits, investigators either examined the lag structure of associations by including both single-day and the average of multiday lags, or selecting lags a priori. The collective
29 30 31 32	evidence indicates a rather immediate response within the first few days of O_3 exposure (i.e., for lags days averaged at 0-1, 0-2, and 0-3 days) for hospital admissions and ED visits for all respiratory outcomes, asthma, and chronic obstructive pulmonary disease in all-year and seasonal analyses.

2.5.4.4 Ozone Concentration-Response Relationship

1	An important consideration in characterizing the O ₃ -morbidity and mortality association
2	is whether the concentration-response (C-R) relationship is linear across the full
3	concentration range that is encountered or if there are concentration ranges where there
4	are departures from linearity (i.e., nonlinearity). In this ISA studies have been identified
5	that attempt to characterize the shape of the O_3 C-R curve along with possible O_3
6	"thresholds" (i.e., O ₃ concentrations which must be exceeded in order to elicit an
7	observable health response). The controlled human exposure and epidemiologic studies
8	that examined the shape of the C-R curve and the potential presence of a threshold have
9	indicated a generally linear C-R function with no indication of a threshold in analyses
10	that have examined 8-h max and 24-h avg O_3 concentrations. However, there is less
11	certainty in the shape of the C-R curve at the lower end of the distribution of O_3
12	concentrations due to the low density of data in this range.
13	Controlled human exposure studies have provided strong and quantifiable C-R data on
14	the human health effects of O_3 . The magnitude of respiratory effects in these studies is
15	generally a function of O ₃ exposure, i.e., the product of concentration (C), minute
16	ventilation (\dot{V}_E), and exposure duration. Several studies provide evidence for a smooth
17	C-R curve without indication of a threshold in young healthy adults exposed during
18	moderate exercise for 6.6 hours to O ₃ concentrations between 40 and 120 ppb
19	(Figure 6-1). It is difficult to characterize the C-R relationship below 40 ppb due to
20	uncertainty associated with the sparse data at these lower concentrations.
21	Although relatively few epidemiologic studies have examined the O3-health effects C-R
22	relationship, the C-R relationship has been examined across multiple health endpoints
23	and exposure durations. Some studies of populations engaged in outdoor activity found
24	that associations between O_3 and lung function decrements persisted at lower O_3
25	concentrations with some studies showing larger negative associations in analyses limited
26	to lower O_3 concentrations (e.g., 60-80 ppb; <u>Table 6-6</u>) and shorter exposure durations
27	(i.e., in the range of 30 minutes to less than 8 hours; <u>Table 6-6</u>). A study examining the
28	C-R relationship between short-term O ₃ exposure and pediatric asthma ED visits found
29	no evidence of a threshold with a linear relationship evident down to 8-h max O_3
30	concentrations as low as 30 ppb (Figure 6-17). In an additional study, authors used a
31	smooth function while also accounting for the potential confounding effects of $PM_{2.5}$, to
32	examine whether the shape of the C-R curve for short-term exposure to O_3 and asthma
33	hospital admissions is linear. When comparing the curve to a linear fit, the authors found
34	that the linear fit is a reasonable approximation of the C-R relationship between O_3 and
35	asthma hospital admissions in the mid-range of the data though it can be seen that there is

1	greater uncertainty at the lower end of the distribution of ambient O_3 concentrations,
2	generally below 20 ppb (Figure 6-15) due to sparse data at these lower concentrations.
3	Several recent studies applied a variety of statistical approaches to examine the shape of
4	the O_3 -mortality C-R relationship and existence of a threshold (Section <u>6.6.2.4</u>). These
5	studies suggest that the shape of the O_3 -mortality C-R curve is linear across the range of
6	O_3 concentrations though uncertainty in the relationship increases at the lower end of the
0 7	distribution (<u>Figure 6-35</u>). Generally, the epidemiologic studies that examined the
8	O_3 -mortality C-R relationship do not provide evidence for the existence of a threshold
8 9	within the range of 24-h average (24-h avg) O_3 concentrations most commonly observed
10	
	in the U.S. during the O_3 season (i.e., above 20 ppb). It should be noted that the
11	evaluation of the C-R relationship for short-term exposure to O_3 and mortality is difficult
12	due to the evidence from multicity studies indicating highly heterogeneous O_3 -mortality
13	associations across regions of the U.S. In addition, there are numerous issues that may
14	influence the shape of the O_3 -mortality C-R relationship that need to be taken into
15	consideration including: multiday effects (distributed lags), and potential adaptation and
16	mortality displacement (i.e., hastening of death by a short period). Additionally, given the
17	effect modifiers identified in mortality analyses that are also expected to vary regionally
18	(e.g., temperature, air conditioning prevalence), a national or combined analysis may not
19	be appropriate to identify whether a threshold exists in the O ₃ -mortality C-R relationship.
20	In addition, the C-R relationship of long-term exposure to O_3 and birth outcomes has
21	been evaluated. Evidence from the southern California Children's Health Study identified
22	a C-R relationship of birth weight with 24-h avg O ₃ concentrations averaged over the
23	entire pregnancy that was clearest above the 30 ppb level (Figure 7-4).
24	Generally, both short- and long-term exposure studies indicate a linear, no threshold C-R
25	relationship when examining the association between O_3 exposure and multiple health
26	effects across the range of 8-h max and 24-h avg O_3 concentrations most commonly
27	observed in the U.S. during the O_3 season (i.e., greater than 20 ppb). However, evidence
28	from studies of respiratory health effects and mortality indicates less certainty in the
29	shape of the C-R curve at the lower end of the distribution of O_3 data, which corresponds
30	to 8-h max and 24-h avg O_3 concentrations generally below 20 ppb.

2.5.4.5 Regional Heterogeneity in Risk Estimates

31	Multicity epidemiologic studies that have examined the relationship between short-term
32	O ₃ exposures and mortality have provided evidence of city-to-city and regional
33	heterogeneity in O ₃ -mortality risk estimates. A possible explanation for this heterogeneity
34	may be differences in community characteristics (individual- or community-level) across

1cities that could modify the O3 effect. Another possible explanation for the observed2heterogeneity could be effect modification by concentrations of other air pollutants or3interactions with temperature or other meteorological factors that vary regionally in the4U.S.

5 An examination of community characteristics measured at the individual level that may 6 contribute to the observed heterogeneity in O₃-mortality risk estimates indicates increased 7 risk in older adults (i.e., ≥ 65 years of age), women, African American individuals, 8 individuals with pre-existing diseases/conditions (e.g., diabetes, atrial fibrillation), and 9 lower SES. Furthermore, studies have examined community characteristics measured at 10 the community level and found that higher O₃-mortality risk estimates were associated 11 with higher: percent unemployment, fraction of the population Black/African-American, 12 percent of the population that take public transportation to work; and with lower: 13 temperatures and percent of households with central air conditioning. There is also 14 evidence of greater effects in cities with lower mean O_3 concentrations. Additionally, 15 there is evidence of increased risk of O₃-related mortality as percentage unemployed 16 increases and a reduction in O_3 -related mortality as mean temperature increased (i.e., a 17 surrogate for air conditioning rate) in the U.S. The lack of a consistent reduction in 18 O₃-risk estimates in cities with a higher percentage of central air conditioning across 19 health outcomes complicates the interpretation of the potential modifying effects of air 20 conditioning use.

21 Overall, the epidemiologic studies that have examined the city-to-city and regional 22 heterogeneity observed in multicity studies have identified a variety of factors that may 23 modify the O₃-mortality or -respiratory hospital admission relationship. Some studies 24 have also examined the correlation with other air pollutants or the potential interactive 25 effects between O₃ and temperature to explain city-to-city heterogeneity in O₃-mortality 26 risk estimates. This includes evidence that O₃-mortality risk estimates in the U.S. varied 27 by mean SO₂ concentrations, the ratio between mean NO₂/PM₁₀ concentrations, and 28 temperatures. However, studies have not consistently identified specific community 29 characteristics that explain the observed heterogeneity.

2.6 Integration of Effects on Vegetation and Ecosystems

30	Chapter <u>9</u> presents the most policy-relevant information related to this review of the
31	NAAQS for the welfare effects of O_3 on vegetation and ecosystems. This section
32	integrates the key findings from the disciplines evaluated in this assessment of the O_3
33	scientific literature, which includes plant physiology, whole plant biology, ecosystems,
34	and exposure-response.

1	Overall, exposure to O_3 is causally related or likely to be causally related to effects
2	observed on vegetation and ecosystems. These effects are observed across the entire
3	continuum of biological organization; from the cellular and subcellular level to the whole
4	plant level, and up to ecosystem-level processes. Furthermore, there is evidence that the
5	effects observed across this continuum are related to one another; effects of O ₃ at lower
6	levels of organization, such as the leaf of an individual plant, can result in effects at
7	higher levels. Ozone enters leaves through stomata, and can alter stomatal conductance
8	and disrupt CO_2 fixation (Section 9.3). These effects can change rates of leaf gas
9	exchange, growth and reproduction at the individual plant level and result in changes in
10	ecosystems, such as productivity, C storage, water cycling, nutrient cycling, and
11	community composition (Section 9.4). Figure 2-3 is a simplified illustrative diagram of
12	the major pathway through which O ₃ enters leaves and the major endpoints O ₃ may affect
13	in vegetation and ecosystems.
14	The framework for causal determinations (see Preamble) has been applied to the body of
15	scientific evidence to examine effects attributed to O_3 exposure (<u>Table 2-3</u>). The
16	summary below provides brief integrated summaries of the evidence that supports the
17	causal determinations. The detailed discussion of the underlying evidence used to

causal determinations. The detailed discussion of the underlying evidence used to
formulate each causal determination can be found in Chapter <u>9</u>. This summary ends with
a short discussion of policy relevant considerations.

2.6.1 Visible Foliar Injury

20	Visible foliar injury resulting from exposure to O_3 has been well characterized and
21	documented over several decades of research on many tree, shrub, herbaceous, and crop
22	species (U.S. EPA, 2006b, 1996b, 1984, 1978a) (Section 9.4.2). Ozone-induced visible
23	foliar injury symptoms on certain bioindicator plant species are considered diagnostic as
24	they have been verified experimentally in exposure-response studies, using exposure
25	methodologies such as continuous stirred tank reactors (CSTRs), open-top chambers
26	(OTCs), and free-air fumigation. Experimental evidence has clearly established a
27	consistent association of visible injury with O3 exposure, with greater exposure often
28	resulting in greater and more prevalent injury. Since publication of the 2006 O ₃ AQCD,
29	the results of several multiple-year field surveys of O3-induced visible foliar injury at
30	National Wildlife Refuges in Maine, Michigan, New Jersey, and South Carolina have
31	been published. New sensitive species showing visible foliar injury continue to be
32	identified from field surveys and verified in controlled exposure studies.

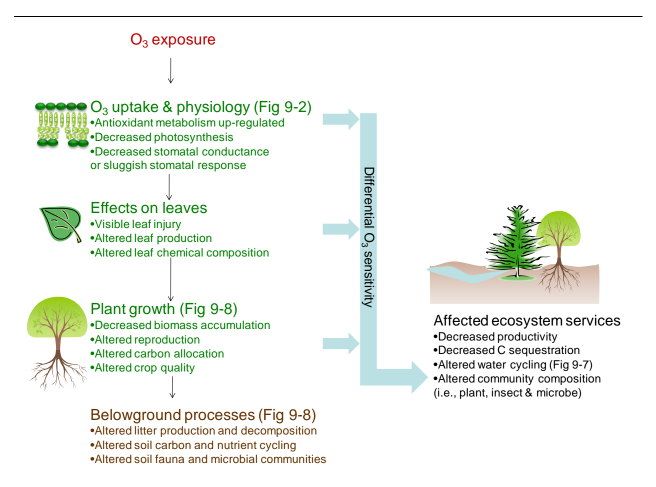


Figure 2-3 An illustrative diagram of the major pathway through which ozone enters leaves and the major endpoints that ozone may affect in plants and ecosystems.

Vegetation and Ecosystem Effects	Conclusions from 2006 O ₃ AQCD	Conclusions from 2011 2nd Draft ISA
Visible Foliar Injury Effects on Vegetation	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause impaired aesthetic quality of many native plants and trees by increasing foliar injury.	Causal Relationship
Reduced Vegetation Growth	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased growth and biomass accumulation in annual, perennial and woody plants, including agronomic crops, annuals, shrubs, grasses, and trees.	Causal Relationship
Reduced Productivity in Terrestrial Ecosystems	There is evidence that O_3 is an important stressor of ecosystems and that the effects of O_3 on individual plants and processes are scaled up through the ecosystem, affecting net primary productivity.	Causal Relationship
Reduced Carbon (C) Sequestration in Terrestrial Ecosystems	Limited studies from previous review	Likely to be a Causal Relationship
Reduced Yield and Quality of Agricultural Crops	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased yield and/or nutritive quality in a large number of agronomic and forage crops.	Causal Relationship
Alteration of Terrestrial Ecosystem Water Cycling	Ecosystem water quantity may be affected by O_3 exposure at the landscape level.	Likely to be a Causal Relationship
Alteration of Below-ground Biogeochemical Cycles	Ozone-sensitive species have well known responses to O_3 exposure, including altered C allocation to below-ground tissues, and altered rates of leaf and root production, turnover, and decomposition. These shifts can affect overall C and N loss from the ecosystem in terms of respired C, and leached aqueous dissolved organic and inorganic C and N.	Causal Relationship
Alteration of Terrestrial Community Composition	Ozone may be affecting above- and below -ground community composition through impacts on both growth and reproduction. Significant changes in plant community composition resulting directly from O_3 exposure have been demonstrated.	Likely to be a Causal Relationship

Table 2-3Summary of ozone causal determinations for vegetation and
ecosystem effects.

1 The use of biological indicators in field surveys to detect phytotoxic levels of O_3 is a 2 longstanding and effective methodology. The USDA Forest Service through the Forest 3 Health Monitoring (FHM) Program (1990-2001) and currently the Forest Inventory and 4 Analysis (FIA) Program has been collecting data regarding the incidence and severity of 5 visible foliar injury on a variety of O_3 sensitive plant species throughout the U.S. The 6 network has provided evidence that O₃ concentrations were high enough to induce visible 7 symptoms on sensitive vegetation. From repeated observations and measurements made 8 over a number of years, specific geographical patterns of visible O_3 injury symptoms can 9 be identified. In addition, a study assessed the risk of O₃-induced visible foliar injury on 10 bioindicator plants in 244 national parks in support of the National Park Service's Vital 11 Signs Monitoring Network. The results of the study demonstrated that the estimated risk 12 of visible foliar injury was high in 65 parks (27%), moderate in 46 parks (19%), and low 13 in 131 parks (54%). Some of the well-known parks with a high risk of O_3 -induced visible 14 foliar injury include Gettysburg, Valley Forge, Delaware Water Gap, Cape Cod, Fire

1Island, Antietam, Harpers Ferry, Manassas, Wolf Trap Farm Park, Mammoth Cave,2Shiloh, Sleeping Bear Dunes, Great Smoky Mountains, Joshua Tree, Sequoia and Kings3Canyon, and Yosemite. Overall, evidence is sufficient to conclude that there is a causal4relationship between ambient O3 exposure and the occurrence of O3-induced5visible foliar injury on sensitive vegetation across the U.S.

2.6.2 Growth, Productivity, Carbon Storage and Agriculture

Ambient O_3 concentrations have long been known to cause decreases in photosynthetic rates and plant growth. The O_3 -induced damages at the plant scale may translate to damages at the stand, then ecosystem scales, and cause changes in productivity and C storage. The effects of O_3 exposure on photosynthesis, growth, biomass allocation, ecosystem production, and ecosystem C sequestration were reviewed for the natural ecosystems, and crop productivity and crop quality were reviewed for the agricultural ecosystems.

2.6.2.1 Natural Ecosystems

13	The previous O ₃ AQCDs concluded that there is strong and consistent evidence that
14	ambient concentrations of O_3 decrease plant photosynthesis and growth in numerous
15	plant species across the U.S. Studies published since the last review continue to support
16	that conclusion (Section 9.4.3.1). Recent studies, based on the Aspen free-air carbon-
17	dioxide/ozone enrichment (FACE) experiment, found that O3 caused reductions in total
18	biomass relative to the control in aspen, paper birch, and sugar maple communities
19	during the first seven years of stand development. Overall, the studies at the Aspen FACE
20	experiment were consistent with the open-top chamber (OTC) studies that were the
21	foundation of previous O3 NAAQS reviews. These results strengthen the understanding
22	of O_3 effects on forests and demonstrate the relevance of the knowledge gained from
23	trees grown in OTC studies.
24	A set of meta-analyses assessed the effects of O_3 on plant photosynthesis and growth
25	across different species and fumigation methods (such as OTC and FACE). Those studies
26	reported that current O_3 concentrations in the northern hemisphere are decreasing
27	photosynthesis (~11%) across tree species, and the decreases in photosynthesis are
28	consistent with cumulative uptake of O_3 into the leaf. The current ambient O_3
29	concentrations (~40 ppb averaged across all hours of exposure) decreased annual total
30	biomass growth of forest species by an average of 7%, with potentially greater decreases
31	(11-17%) with elevated O_3 exposures (Section <u>9.4.3.1</u>). The meta-analyses further

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1	confirmed that reduction of plant photosynthesis and growth under O ₃ exposure are
2	coherent across numerous species and various experimental techniques.
2	
3	Studies during recent decades have also demonstrated O_3 alters biomass allocation and
4	plant reproduction (Section 9.4.3). Recent meta-analyses have generally indicated that O_3
5	reduced C allocated to roots. Several recent studies published since the 2006 O_3 AQCD
6	further demonstrate that O_3 altered reproductive processes, such as timing of flowering,
7	number of flowers, fruits and seeds, in herbaceous and woody plant species. However, a
8	knowledge gap still exists pertaining to the exact mechanism of the responses of
9	reproductive processes to O_3 exposure (Section <u>9.4.3.3</u>).
10	Studies at the leaf and plant scales show that O3 decreased photosynthesis and plant
11	growth, providing coherence and biological plausibility for the reported decreases in
12	ecosystem productivity. During the previous NAAQS reviews, there were very few
13	studies that investigated the effect of O_3 exposure on ecosystem productivity and
14	C sequestration. Recent studies from long-term FACE experiments and ecosystem
15	models provided evidence of the association of O ₃ exposure and reduced productivity at
16	the ecosystem scale. Elevated O_3 reduced stand biomass at Aspen FACE after 7 years of
17	O ₃ exposure, and annual volume growth at the Kranzberg Forest in Germany. Results
18	across different ecosystem models were consistent with the FACE experimental
19	evidence, which showed that O_3 reduced ecosystem productivity (Section <u>9.4.3.4</u>). In
20	addition to primary productivity, other indicators such as net ecosystem productivity
21	(NEP), net ecosystem CO ₂ exchange (NEE) and C sequestration were often assessed by
22	model studies. Model simulations consistently found that O ₃ exposure caused negative
23	impacts on these indicators (Section 9.4.3.4, Table 9-3), but the severity of these impacts
24	was influenced by multiple interactions of biological and environmental factors. The
25	suppression of ecosystem C sinks results in more CO ₂ accumulation in the atmosphere. A
26	recent study suggested that the indirect radiative forcing caused by O ₃ exposure through
27	lowering the ecosystem C sink could have an even greater impact on global warming than
28	the direct radiative forcing of O_3 .
29	Although O ₃ generally causes negative effects on ecosystem productivity, the magnitude
30	of the response varies among plant communities (Section $9.4.3.4$). For example, O ₃ had
31	little impact on white fir, but greatly reduced growth of ponderosa pine in southern
32	California. Ozone decreased net primary production (NPP) of most forest types in the
33	Mid-Atlantic region, but had small impacts on spruce-fir forest. Ozone could also affect
34	regional C budgets through interacting with multiple factors, such as N deposition,
35	elevated CO_2 and land use history. Model simulations suggested that O_3 partially offset
36	the growth stimulation caused by elevated CO_2 and N deposition in both Northeast- and
37	Mid-Atlantic-region forest ecosystems of the U.S.
51	The function region forest coolystems of the 0.5.

1Overall, evidence is sufficient to conclude that there is a causal relationship between2ambient O3 exposure and reduced native plant growth and productivity, and a likely3causal relationship between O3 exposure and reduced carbon sequestration in4terrestrial ecosystems.

2.6.2.2 Agricultural Crops

5	The detrimental effect of O_3 on crop production has been recognized since the 1960's and
6	a large body of research has subsequently stemmed from those initial findings. Previous
7	O_3 AQCDs have extensively reviewed this body of literature. Current O_3 concentrations
8	across the U.S. are high enough to cause yield loss for a variety of agricultural crops
9	including, but not limited to, soybean, wheat, potato, watermelon, beans, turnip, onion,
10	lettuce, and tomato (Section $9.4.4.1$). Continued increases in O ₃ concentration may
11	further decrease yield in these sensitive crops. Despite the well-documented yield losses
12	due to increasing O ₃ concentration, there is still a knowledge gap pertaining to the exact
13	mechanism of O ₃ -induced yield loss. Research has linked increasing O ₃ concentration to
14	decreased photosynthetic rates and accelerated senescence, which are related to yield.
15	In addition, recent research has highlighted the effects of O ₃ on crop quality. Increasing
16	O3 concentration decreases nutritive quality of grasses, decreases macro- and micro-
17	nutrient concentrations in fruits and vegetable crops, and decreases cotton fiber quality.
18	These areas of research require further investigation to determine the mechanism and
19	dose-responses (Section $9.4.4.2$).
20	During the previous NAAQS reviews, there were very few studies that estimated O_3
21	impacts on crop yields at large geographical scales (i.e., regional, national or global).
22	Recent modeling studies found that O ₃ generally reduced crop yield, but the impacts
23	varied across regions and crop species (Section 9.4.4.1). For example, the largest
24	O_3 -induced crop yield losses occurred in high-production areas exposed to high O_3
25	concentrations, such as the Midwest and the Mississippi Valley regions of the U.S.
26	Among crop species, the estimated yield loss for wheat and soybean were higher than
27	rice and maize. Satellite and ground-based O_3 measurements have been used to assess
28	yield loss caused by O3 over the continuous tri-state area of Illinois, Iowa, and
29	Wisconsin. The results showed that O ₃ concentrations reduced soybean yield, which
30	correlates well with the previous results from FACE- and OTC-type experiments
31	(Section $9.4.4.1$).
32	Evidence is sufficient to conclude that there is a causal relationship between O_3
33	exposure and reduced yield and quality of agricultural crops.

2.6.3 Water Cycling

1	Ozone can affect water use in plants and ecosystems through several mechanisms
2	including damage to stomatal functioning and loss of leaf area. Section 9.3.6 reviewed
3	possible mechanisms for O ₃ exposure effects on stomatal functioning. Regardless of the
4	mechanism, O ₃ exposure has been shown to alter stomatal performance, which may affect
5	plant and stand transpiration and therefore possibly affecting hydrological cycling.
6	Although the evidence was from a limited number of field and modeling studies, these
7	findings showed an association of O ₃ exposure and the alteration of water use and cycling
8	in vegetation and ecosystems (Section $9.4.5$). There is not a clear consensus on the nature
9	of leaf-level stomatal conductance response to O ₃ exposure. When measured at steady-
10	state high light conditions, leaf-level stomatal conductance is often found to be reduced
11	when exposed to O_3 . However, measurements of stomatal conductance under dynamic
12	light and vapor pressure deficit conditions indicate sluggish responses under elevated O_3
13	exposure which could potentially lead to increased water loss from vegetation. In
14	situations where stomata fail to close under low light or water stressed conditions water
15	loss may be greater over time. In other situations it is possible that sluggish stomata may
16	fail to completely open in response to environmental stimuli and result in decreased water
17	loss. Field studies suggested that peak hourly O3 exposure increased the rate of water loss
18	from several tree species, and led to a reduction in the late-season modeled stream flow in
19	three forested watersheds in eastern Tennessee. Sluggish stomatal responses during O ₃
20	exposure was suggested as a possible mechanism for increased water loss during peak O_3
21	exposure. Currently, the O ₃ -induced reduction in stomatal aperture is the biological
22	assumption for most process-based models. Therefore, results of those models normally
23	found that O_3 reduced water loss. For example, one study found that O_3 damage and
24	N limitation together reduced evapotranspiration and increase runoff.
25	Although the direction of the response differed among studies, the evidence is sufficient
26	to conclude that there is likely to be a causal relationship between O_3 exposure and
27	the alteration of ecosystem water cycling.

2.6.4 Below-Ground Processes

28	Below-ground processes are tightly linked with aboveground processes. The responses of
29	aboveground process to O ₃ exposure, such as reduced photosynthetic rates, increased
30	metabolic cost, and reduced root C allocation, have provided biologically plausible
31	mechanisms for the alteration of below-ground processes. Since the 2006 O ₃ AQCD,
32	more evidence has shown that although the responses are often species specific, O_3

1	altered the quality and quantity of C input to soil, microbial community composition, and
2	C and nutrient cycling.
3	Results from Aspen FACE and other experimental studies consistently found that O ₃
4	reduced litter production and altered C chemistry, such as soluble sugars, soluble
5	phenolics, condensed tannins, lignin, and macro/micro nutrient concentration in litter
6	(Section <u>9.4.6.1</u>). Under elevated O_3 , the changes in substrate quality and quantity could
7	alter microbial metabolism, and therefore soil C and nutrient cycling. Several studies
8	indicated that O_3 generally suppressed soil enzyme activities (Section <u>9.4.6.2</u>). However,
9	the impact of O ₃ on litter decomposition was inconsistent and varied among species,
10	sites, and exposure length. Similarly, O ₃ had inconsistent impacts on dynamics of micro
11	and macro nutrients (Section $9.4.6.4$).
12	Studies from the Aspen FACE experiment suggested that the response of below-ground
13	C cycle to O ₃ exposure, such as litter decomposition, soil respiration, and soil C content,
14	changed over time. For example, in the early part of the experiment (1998-2003), O ₃ had
15	no impact on soil respiration but reduced the formation rates of total soil C under
16	elevated CO ₂ . However, after 10 to 11 years of exposure, O ₃ was found to increase soil
17	respiration but have no substantial impact on soil C formation under elevated CO ₂
18	(Section $9.4.6.3$).
19	The evidence is sufficient to infer that there is a causal relationship between O_3

19The evidence is sufficient to infer that there is a causal relationship between O320exposure and the alteration of below-ground biogeochemical cycles.

2.6.5 Community Composition

21	In the 2006 O ₃ AQCD, the impact of O ₃ exposure on species competition and community
22	composition was assessed. Ozone was found to be one of the dominant factors causing a
23	decline in ponderosa and Jeffrey pine in the San Bernardino Mountains in southern
24	California. Ozone exposure also tended to shift the grass-legume mixtures in favor of
25	grass species. Since the 2006 O_3 AQCD, more evidence has shown that O_3 exposure
26	changed the competitive interactions and led to loss of O ₃ sensitive species or genotypes.
27	Studies found that the severity of O ₃ damage on growth, reproduction and foliar injury
28	varied among species (Section 9.4.3), which provided the biological plausibility for the
29	alteration of community composition. Additionally, research since the last review has
30	shown that O_3 can alter community composition and diversity of soil microbial
31	communities.
32	The decline of conifer forests under O_3 exposure was continually observed in several
33	regions. Ozone damage was believed to be an important causal factor in the dramatic

1	decline of sacred fir in the valley of Mexico, as well as cembran pine in southern France
2	and the Carpathian Mountains, although several factors, such as drought, insect outbreak
3	and forest management, may also contribute to or even be the dominant factors causing
4	the mortality of the conifer trees. Results from the Aspen FACE site indicated that O_3
5	could alter community composition of broadleaf forests as well. At the Aspen FACE site,
6	O ₃ reduced growth and increased mortality of a sensitive aspen clone, while the O ₃
7	tolerant clone emerged as the dominant clone in the pure aspen community. In the mixed
8	aspen-birch and aspen-maple communities, O ₃ reduced the competitive capacity of aspen
9	compared to birch and maple (Section $9.4.7.1$).
10	The tendency for O ₃ -exposure to shift the biomass of grass-legume mixtures in favor of
11	grass species was reported in the 2006 O ₃ AQCD and has been generally confirmed by
12	recent studies. However, in a high elevation mature/species-rich grass-legume pasture, O3
13	fumigation showed no substantial impact on community composition (Section $9.4.7.2$).
14	Ozone exposure not only altered community composition of plant species, but also
15	microorganisms. The shift in community composition of bacteria and fungi has been
16	observed in both natural and agricultural ecosystems, although no general patterns could
17	be identified (Section $9.4.7.3$).
18	The evidence is sufficient to conclude that there is likely to be a causal relationship
19	between O_3 exposure and the alteration of community composition of some
20	ecosystems.

2.6.6 Policy Relevant Considerations

2.6.6.1 Air Quality Indices

21	Exposure indices are metrics that quantify exposure as it relates to measured plant injury
22	(e.g., reduced growth). They are summary measures of monitored ambient O ₃
23	concentrations over time intended to provide a consistent metric for reviewing and
24	comparing exposure-response effects obtained from various studies. No recent
25	information is available since 2006 that alters the basic conclusions put forth in the 2006
26	and 1996 O_3 AQCDs. These AQCDs focused on the research used to develop various
27	exposure indices to help quantify effects on growth and yield in crops, perennials, and
28	trees (primarily seedlings). The performance of indices was compared through regression
29	analyses of earlier studies designed to support the estimation of predictive O3 exposure-
30	response models for growth and/or yield of crops and tree (seedling) species.

1	Another approach for improving risk assessment of vegetation response to ambient O_3 is
2	based on determining the O_3 concentration from the atmosphere that enters the leaf
3	(i.e., flux or deposition). Interest has been increasing in recent years, particularly in
4	Europe, in using mathematically tractable flux models for O_3 assessments at the regional,
5	national, and European scale. While some efforts have been made in the U.S. to calculate
6	O_3 flux into leaves and canopies, little information has been published relating these
7	fluxes to effects on vegetation. There is also concern that not all O_3 stomatal uptake
8	results in a yield reduction, which depends to some degree on the amount of internal
9	detoxification occurring with each particular species. Species having high detoxification
10	capacity may show little relationship between O_3 stomatal uptake and plant response. The
11	lack of data in the U.S. and the lack of understanding of detoxification processes have
12	made this technique less viable for vulnerability and risk assessments in the U.S.
13 14	The main conclusions from the 1996 and 2006 O_3 AQCDs regarding indices based on ambient exposure remain valid. These key conclusions can be restated as follows:
15 16 17 18 19	 O₃ effects in plants are cumulative; higher O₃ concentrations appear to be more important than lower concentrations in eliciting a response; plant sensitivity to O₃ varies with time of day and plant development stage; and
20	 quantifying exposure with indices that cumulate hourly O₃ concentrations and
21	preferentially weight the higher concentrations improves the explanatory
22	power of exposure/response models for growth and yield, over using indices
23	based on mean and peak exposure values.
24	Various weighting functions have been used, including threshold-weighted
25	(e.g., SUM06) and continuous sigmoid-weighted (e.g., W126) functions. Based on
26	statistical goodness-of-fit tests, these cumulative, concentration-weighted indices could
27	not be differentiated from one another using data from previous exposure studies.
28	Additional statistical forms for O ₃ exposure indices are summarized in Section <u>9.5</u> of this
29	ISA. The majority of studies published since the 2006 O ₃ AQCD do not change earlier
30	conclusions, including the importance of peak concentrations, and the duration and
31	occurrence of O ₃ exposures in altering plant growth and yield.
32	Given the current state of knowledge and the best available data, exposure indices that
33	cumulate and differentially weight the higher hourly average concentrations and also
34	include the mid-level values continue to offer the most defensible approach for use in
35	developing response functions and comparing studies, as well as for defining future
36	indices for vegetation protection.

2.6.6.2 Exposure-Response

1	None of the information on effects of O_3 on vegetation published since the 2006 O_3
2	AQCD has modified the assessment of quantitative exposure-response relationships that
3	was presented in that document (U.S. EPA, 2006b). This assessment updates the 2006
4	exposure-response models by computing them using the W126 metric, cumulated over
5	90 days. Almost all of the experimental research on the effects of O ₃ on growth or yield
6	of plants published since 2006 used only two levels of exposure. In addition, hourly O_3
7	concentration data that would allow calculations of exposure using the W126 metric are
8	generally unavailable. However, two long-term experiments, one with a crop species
9	(soybean), one with a tree species (aspen), have produced data that are used in
10	Section 9.6 to validate the exposure-response models presented in the 2006 O ₃ AQCD,
11	and the methodology used to derive them. EPA compared predictions from the models
12	presented in the 2006 O ₃ AQCD, updated to use the 90 day 12hr W126 metric, with more
13	recent observations for yield of soybean and biomass growth of trembling aspen. The
14	models were parameterized using data from the National Crop Loss Assessment Network
15	(NCLAN) and EPA's National Health and Environmental Effects Research Laboratory -
16	Western Ecology Division (NHEERL-WED) projects, which were conducted in OTCs.
17	The more recent observations were from experiments using FACE technology, which is
18	intended to provide conditions closer to natural environments than OTC. Observations
19	from these new experiments were exceptionally close to predictions from the models.
20	The accuracy of model predictions for two widely different plant species, grown under
21	very different conditions, provides support for the validity of the models for crops and
22	trees developed using the same methodology and data for other species. However,
23	variability observed among species in the NCLAN and NHEERL-WED projects indicates
24	that the range of sensitivity between and among species is likely quite wide.
25	Results from several meta-analyses have provided approximate values for responses of
26	yield of soybean, wheat, rice and other crops under broad categories of exposure, relative
27	to charcoal-filtered air. Additional reports have summarized yield data for six crop
28	species under various broad comparative exposure categories, and reviewed 263 studies
29	that reported effects on tree biomass. However, these analyses have proved difficult to
30	compare with exposure-response models, especially given that exposure was not
21	

31 expressed using a common metric (i.e., W126).

2.7 The Role of Tropospheric Ozone in Climate Change and UV-B Effects

Atmospheric O_3 plays an important role in the Earth's energy budget by interacting with incoming solar radiation and outgoing infrared radiation. Tropospheric O_3 makes up only a small portion of the total column of O_3 , but it has important incremental effects on the overall radiation budget. Chapter <u>10</u> assesses the specific role of tropospheric O_3 in the earth's radiation budget and how perturbations in tropospheric O_3 might affect (1) climate through its role as a greenhouse gas, and (2) health, ecology and welfare through its role in shielding the earth's surface from solar ultraviolet (UV) radiation.

2.7.1 Tropospheric Ozone as a Greenhouse Gas

Ozone is an important greenhouse gas, and increases in its abundance in the troposphere may contribute to climate change according to the 2007 climate assessment by the Intergovernmental Panel on Climate Change (IPCC). Models calculate that the global burden of tropospheric O₃ has doubled since the pre-industrial era, while observations indicate that in some regions O₃ may have increased by factors as great as 4 or 5. These increases are tied to the rise in emissions of O₃ precursors from human activity, mainly fossil fuel consumption and agricultural processes.

15 Figure 2-4 shows the main steps involved in the influence of tropospheric O_3 on climate. 16 Emissions of O_3 precursors including CO, VOCs, CH₄, and NO_x lead to production of 17 tropospheric O_3 . A change in the abundance of tropospheric O_3 perturbs the radiative 18 balance of the atmosphere, an effect quantified by the radiative forcing (RF) metric. The 19 earth-atmosphere-ocean system responds to the forcing with a climate response, typically 20 expressed as a change in surface temperature. Finally, the climate response causes 21 downstream climate-related health and ecosystem impacts, such as redistribution of 22 diseases or ecosystem characteristics due to temperature changes. Feedbacks from both 23 the climate response and downstream impacts can, in turn, affect the abundance of 24 tropospheric O₃ and O₃ precursors through multiple feedback mechanisms as indicated in 25 Figure 2-4. Direct feedbacks are discussed in Section 10.3.2.4 while downstream climate 26 impacts and their feedbacks are extremely complex and outside the scope of this 27 assessment.

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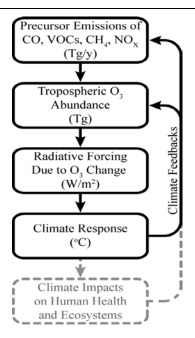
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Note: Units shown are those typical for each quantity illustrated. Feedbacks from both the climate response and climate impacts can, in turn, affect the abundance of tropospheric O_3 and O_3 precursors through multiple feedback mechanisms. Climate impacts are deemphasized in the figure since these downstream effects are extremely complex and outside the scope of this assessment.

Figure 2-4 Schematic illustrating the effects of tropospheric ozone on climate; including the relationship between precursor emissions, tropospheric ozone abundance, radiative forcing, climate response, and climate impacts.Tropospheric Ozone and UV-B related effects

1	The impact of the tropospheric O ₃ change since pre-industrial times on climate has been
-	
2	estimated to be about 25-40% of the anthropogenic CO_2 impact and about 75% of the
3	anthropogenic CH ₄ impact according to the IPCC, ranking it third in importance among
4	the greenhouse gases. There are large uncertainties in the RF estimate attributed to
5	tropospheric O ₃ , making the impact of tropospheric O ₃ on climate more uncertain than
6	the impact of the long-lived greenhouse gases. Overall, the evidence supports a causal
7	relationship between changes in tropospheric O_3 concentrations and radiative
8	forcing.
9	RF does not take into account the climate feedbacks that could amplify or dampen the
10	actual surface temperature response. Quantifying the change in surface temperature
10 11	
	actual surface temperature response. Quantifying the change in surface temperature
11	actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions
11 12	actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the
11 12 13	actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given RF is highly uncertain and can vary greatly
11 12 13 14	actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given RF is highly uncertain and can vary greatly among models and from region to region within the same model. In light of these

2.7.3 Tropospheric Ozone and UV-B Related Effects

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1	UV radiation emitted from the Sun contains sufficient energy when it reaches the Earth to
2	break (photolyze) chemical bonds in molecules, thereby leading to damaging effects on
3	living organisms and materials. Atmospheric O3 plays a crucial role in reducing exposure
4	to solar UV radiation at the Earth's surface. Ozone in the stratosphere is responsible for
5	the majority of this shielding effect, as approximately 90% of total atmospheric O_3 is
6	located there over mid-latitudes. Ozone in the troposphere provides supplemental
7	shielding of radiation in the wavelength band from 280-315 nm, referred to as UV-B
8	radiation. UV-B radiation has important effects on human health and ecosystems, and is
9	associated with materials damage.
10	Human health effects associated with solar UV-B radiation exposure include erythema,
11	skin cancer, ocular damage, and immune system suppression. A potential human health
12	benefit of increased UV-B exposure involves the UV-induced production of vitamin D
13	which may help reduce the risk of metabolic bone disease, type I diabetes, mellitus, and
14	rheumatoid arthritis, and may provide beneficial immunomodulatory effects on multiple
15	sclerosis, insulin-dependent diabetes mellitus, and rheumatoid arthritis. Ecosystem and
16	materials damage effects associated with solar UV-B radiation exposure include
17	terrestrial and aquatic ecosystem impacts, alteration of biogeochemical cycles, and
18	degradation of man-made materials.
19	There is a lack of published studies that critically examine the incremental health or
20	welfare effects (adverse or beneficial) attributable specifically to changes in UV-B
21	exposure resulting from perturbations in tropospheric O_3 concentrations. The effects are
22	expected to be small and they cannot yet be critically assessed within reasonable
23	uncertainty. Overall, the evidence is inadequate to determine if a causal relationship
24	exists between changes in tropospheric O_3 concentrations and effects on health
25	and welfare related to UV-B shielding.

2.8 Summary of Causal Determinations for Health Effects and Welfare Effects

- This chapter has provided an overview of the underlying evidence used in making the
 causal determinations for the health and welfare effects of O₃. This review builds upon
 the conclusions of the previous AQCDs for O₃.
 The evaluation of the epidemiologic, toxicological, and controlled human exposure
- 30 studies published since the completion of the 2006 O_3 AQCD have provided additional 31 evidence for O_3 -related health outcomes. Table 2-4 provides an overview of the causal

1	determinations for all of the health outcomes evaluated. Causal determinations for O_3 and
2	welfare effects are included in <u>Table 2-5</u> , while causal determinations for climate change
3	and UV-B effects are in Table 2-6. Detailed discussions of the scientific evidence and
4	rationale for these causal determinations are provided in subsequent chapters of this ISA.

Table 2-4Summary of ozone causal determinations by exposure duration
and health outcome.

Health Outcome	Conclusions from 2006 O ₃ AQCD	Conclusions from 2012 3rd Draft ISA
Short-Term Exposu	re to O ₃	
Respiratory effects	The overall evidence supports a causal relationship between acute ambient O_3 exposures and increased respiratory morbidity outcomes.	Causal Relationship
Cardiovascular effects	The limited evidence is highly suggestive that O_3 directly and/or indirectly contributes to cardiovascular-related morbidity, but much remains to be done to more fully substantiate the association.	Suggestive of a Causal Relationship
Central nervous system effects	Toxicological studies report that acute exposures to O_3 are associated with alterations in neurotransmitters, motor activity, short and long term memory, sleep patterns, and histological signs of neurodegeneration.	Suggestive of a Causal Relationship
Total Mortality	The evidence is highly suggestive that O_3 directly or indirectly contributes to non-accidental and cardiopulmonary-related mortality.	Likely to be a Causal Relationship
Long-term Exposur	e to O ₃	
Respiratory effects	The current evidence is suggestive but inconclusive for respiratory health effects from long-term O_3 exposure.	Likely to be a Causal Relationship
Cardiovascular Effects	No studies from previous review	Suggestive of a Causal Relationship
Reproductive and developmental effects	Limited evidence for a relationship between air pollution and birth-related health outcomes, including mortality, premature births, low birth weights, and birth defects, with little evidence being found for O_3 effects.	Suggestive of a Causal Relationship
Central nervous system effects	Evidence regarding chronic exposure and neurobehavioral effects was not available.	Suggestive of a Causal Relationship
Cancer	Little evidence for a relationship between chronic O_3 exposure and increased risk of lung cancer.	Inadequate to infer a Causal Relationship
Total Mortality	There is little evidence to suggest a causal relationship between chronic O_3 exposure and increased risk for mortality in humans.	Suggestive of a Causal Relationship

Vegetation and Ecosystem Effects	Conclusions from 2006 O ₃ AQCD	Conclusions from 2012 3rd Draft ISA
Visible Foliar Injury Effects on Vegetation	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause impaired aesthetic quality of many native plants and trees by increasing foliar injury.	Causal Relationship
Reduced Vegetation Growth	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased growth and biomass accumulation in annual, perennial and woody plants, including agronomic crops, annuals, shrubs, grasses, and trees.	Causal Relationship
Reduced Productivity in Terrestrial Ecosystems	There is evidence that O_3 is an important stressor of ecosystems and that the effects of O_3 on individual plants and processes are scaled up through the ecosystem, affecting net primary productivity.	Causal Relationship
Reduced Carbon (C) Sequestration in Terrestrial Ecosystems	Limited studies from previous review	Likely to be a Causal Relationship
Reduced Yield and Quality of Agricultural Crops	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased yield and/or nutritive quality in a large number of agronomic and forage crops.	Causal Relationship
Alteration of Terrestrial Ecosystem Water Cycling	Ecosystem water quantity may be affected by O_3 exposure at the landscape level.	Likely to be a Causal Relationship
Alteration of Below-ground Biogeochemical Cycles	Ozone-sensitive species have well known responses to O_3 exposure, including altered C allocation to below-ground tissues, and altered rates of leaf and root production, turnover, and decomposition. These shifts can affect overall C and N loss from the ecosystem in terms of respired C, and leached aqueous dissolved organic and inorganic C and N.	Causal Relationship
Alteration of Terrestrial Community Composition	Ozone may be affecting above- and below -ground community composition through impacts on both growth and reproduction. Significant changes in plant community composition resulting directly from O_3 exposure have been demonstrated.	Likely to be a Causal Relationship

Table 2-5 Summary of ozone causal determination for welfare effects.

Table 2-6Summary of ozone causal determination for climate and UV-B
effects.

Effects	Conclusions from 2006 O ₃ AQCD	Conclusions from 2012 3rd Draft ISA
Radiative Forcing	Climate forcing by O_3 at the regional scale may be its most important impact on climate.	Causal Relationship
Climate Change	While more certain estimates of the overall importance of global-scale forcing due to tropospheric O_3 await further advances in monitoring and chemical transport modeling, the overall body of scientific evidence suggests that high concentrations of O_3 on the regional scale could have a discernible influence on climate, leading to surface temperature and hydrological cycle changes.	Likely to be a Causal Relationship
Health and Welfare Effects Related to UV-B Shielding	UV-B has not been studied in sufficient detail to allow for a credible health benefits assessment. In conclusion, the effect of changes in surface-level O_3 concentrations on UV-induced health outcomes cannot yet be critically assessed within reasonable uncertainty.	Inadequate to Determine if a Causal Relationship Exists

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3 ATMOSPHERIC CHEMISTRY AND AMBIENT CONCENTRATIONS

3.1 Introduction

1	In the stratosphere, O_3 serves the beneficial role of absorbing the Sun's harmful
2	ultraviolet radiation and preventing the majority of this radiation from reaching the
3	Earth's surface. In the troposphere, however, O_3 and other photochemical oxidants are air
4	pollutants that can exert harmful effects on humans, animals, and vegetation. This chapter
5	discusses the atmospheric chemistry associated with tropospheric O_3 and other related
6	photochemical oxidants and provides a detailed description of their surface-level
7	concentrations. The focus of this chapter is on O_3 since it is the NAAQS indicator for all
8	photochemical oxidants. To the extent possible, other photochemical oxidants are
9	
	discussed, but limited information is currently available. Although O_3 is involved in
10	reactions in indoor air, the focus in this chapter will be on chemistry occurring in
11	outdoor, ambient air.
12	The material in this chapter is organized as follows. Section 3.2 outlines the physical and
13	chemical processes involved in O_3 formation and removal. Section <u>3.3</u> describes the latest
14	methods used to model global O_3 concentrations, and Section 3.4 describes the
15	application of these methods for estimating background concentrations of O ₃ that are
16	useful for risk and policy assessments informing decisions about the NAAQS. Section 3.5
17	includes a comprehensive description of available O ₃ monitoring techniques and
18	monitoring networks, while Section 3.6 presents information on the spatial and temporal
19	variability of O ₃ concentrations across the U.S. and their associations with other
20	pollutants using available monitoring data. Section 3.7 summarizes the main conclusions
21	from Chapter <u>3</u> . Finally, Section <u>3.8</u> provides supplemental material on atmospheric
22	model simulations of background O_3 concentrations (referenced in Section 3.4) and
23	Section 3.9 contains supplemental material on observed ambient O ₃ concentrations
24	(referenced in Section 3.6).

3.2 Physical and Chemical Processes

25	Ozone in the troposphere is a secondary pollutant formed by photochemical reactions of
26	precursor gases and is not directly emitted from specific sources. Ozone and other
27	oxidants, such as peroxyacetyl nitrate (PAN) and H_2O_2 form in polluted areas by
28	atmospheric reactions involving two main classes of precursor pollutants: VOCs and

1	NO _X . ¹ Carbon monoxide (CO) is also important for O ₃ formation in polluted areas and in
2	the remote troposphere. The formation of O ₃ , other oxidants and oxidation products from
3	these precursors is a complex, nonlinear function of many factors including (1) the
4	intensity and spectral distribution of sunlight; (2) atmospheric mixing; (3) concentrations
5	of precursors in the ambient air and the rates of chemical reactions of these precursors;
6	and (4) processing on cloud and aerosol particles.
7	Ozone is present not only in polluted urban atmospheres, but throughout the troposphere,
8	even in remote areas of the globe. The same basic processes involving sunlight-driven
9	reactions of NO_X , VOCs and CO contribute to O_3 formation throughout the troposphere.
10	These processes also lead to the formation of other photochemical products, such as
11	PAN, HNO ₃ , and H_2SO_4 , and to other compounds, such as HCHO and other carbonyl
12	compounds, and to secondary components of particulate matter.
13	A schematic overview of the major photochemical cycles influencing O ₃ in the
14	troposphere and the stratosphere is given in Figure 3-1. The processes responsible for
15	producing summertime O_3 episodes are fairly well understood, and were covered in detail
16	in the 2006 O_3 AQCD (U.S. EPA, 2006b). This section focuses on topics that form the
17	basis for discussions in later chapters and for which there is substantial new information
18	since the previous O_3 review.

¹ The term VOCs refers to all organic gas-phase compounds in the atmosphere, both biogenic and anthropogenic in origin. This definition excludes CO and CO₂. NO_X, also referred to as nitrogen oxides, is equal to the sum of NO and NO₂.

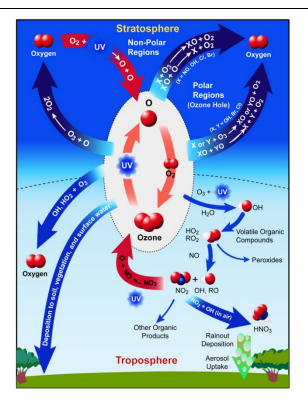


Figure 3-1 Schematic overview of photochemical processes influencing stratospheric and tropospheric ozone.

1	Major episodes of high O ₃ concentrations in the eastern U.S. and in Europe are associated
2	with slow moving high pressure systems. High pressure systems during the warmer
3	seasons are associated with the sinking of air, resulting in warm, generally cloudless
4	skies, with light winds. The sinking of air results in the development of stable conditions
5	near the surface which inhibit or reduce the vertical mixing of O_3 precursors.
6	Photochemical activity involving these precursors is enhanced because of higher
7	temperatures and the availability of sunlight during the warmer seasons. In the eastern
8	U.S., concentrations of O_3 and other secondary pollutants are determined by
9	meteorological and chemical processes extending typically over areas of several hundred
10	thousand square kilometers (Civerolo et al., 2003; Rao et al., 2003). Ozone episodes are
11	thus best regarded as regional in nature. The conditions conducive to formation of high
12	O ₃ can persist for several days. These conditions have been described in greater detail in
13	the 1996 and 2006 O ₃ AQCDs (U.S. EPA, 2006b, 1996a). The transport of pollutants
14	downwind of major urban centers is characterized by the development of urban plumes.
15	Mountain barriers limit mixing (as in Los Angeles and Mexico City) and result in a
16	higher frequency and duration of days with high O3 concentrations. However, orographic
17	lifting over the San Gabriel Mountains results in O3 transport from Los Angeles to areas

- hundreds of kilometers downwind (e.g., in Colorado and Utah) (Langford et al., 2009). Ozone concentrations in southern urban areas (such as Houston, TX and Atlanta, GA) tend to decrease with increasing wind speed. In northern U.S. cities (such as Chicago, IL; New York, NY; Boston, MA; and Portland, ME), the average O₃ concentrations over the metropolitan areas increase with wind speed, indicating that transport of O₃ and its precursors from upwind areas is important (<u>Schichtel and Husar, 2001; Husar and Renard, 1998</u>).
- 8 Aircraft observations indicate that there can be substantial differences in mixing ratios of
 9 key species between the surface and the overlying atmosphere (Berkowitz and Shaw,
 10 1997; Fehsenfeld et al., 1996). In particular, mixing ratios of O₃ can (depending on time
- 11and location) be higher in the lower free troposphere (aloft) than in the planetary12boundary layer (PBL) during multiday O3 episodes (Taubman et al., 2006; Taubman et13al., 2004). Convective processes and turbulence transport O3 and other pollutants both14upward and downward throughout the planetary boundary layer and the free troposphere.15During the day, convection is driven by heating of the earth's surface results in a deeper
- 16 PBL with vertically well mixed O_3 and precursors. As solar heating of the surface 17 decreases going into night, the daytime boundary layer collapses leaving behind O_3 and 18 its precursors in a residual layer above a shallow nighttime boundary layer. Pollutants in 19 the residual layer have now become essentially part of the free troposphere, as described 20 in Annex AX2.3.2 of the 2006 O₃ AQCD (U.S. EPA, 2006b). Winds in the free 21 troposphere tend to be stronger than those closer to the surface and so are capable of 22 transporting pollutants over long distances. Thus, O_3 and its precursors can be transported 23 vertically by convection into the upper part of the mixed layer on one day, then
 - transported overnight as a layer of elevated mixing ratios, and then entrained into a growing convective boundary layer downwind and brought back down to the surface.
- High O₃ concentrations showing large diurnal variations at the surface in southern New
 England were associated with the presence of such layers (Berkowitz et al., 1998). Winds
 several hundred meters above the ground can bring pollutants from the west, even though
 surface winds are from the southwest during periods of high O₃ in the eastern U.S.
 (Blumenthal et al., 1997). These considerations suggest that in many areas of the U.S., O₃
 and its precursors can be transported over hundreds if not thousands of kilometers.
- 32Nocturnal low level jets (LLJs) are an efficient means for transporting pollutants that33have been entrained into the residual boundary layer over hundreds of kilometers. LLJs34are most prevalent in the central U.S. extending northward from eastern Texas, and along35the Atlantic states extending southwest to northeast. LLJs have also been observed off the36coast of California. Turbulence induced by wind shear associated with LLJs brings37pollutants to the surface and results in secondary O3 maxima during the night and early

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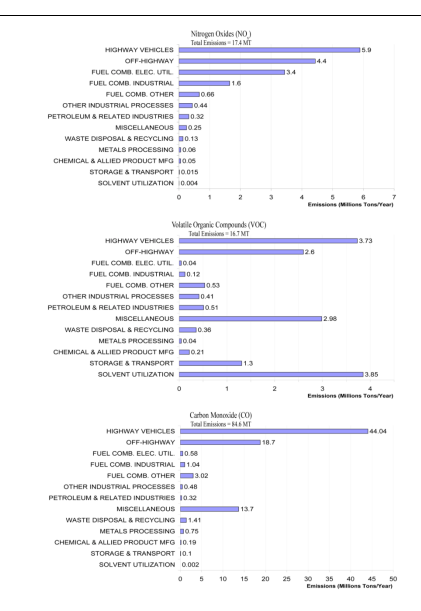
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1	morning in many locations (Corsmeier et al., 1997). Comparison of observations at
2	low-elevation surface sites with those at nearby high-elevation sites at night can be used
3	to discern the effects of LLJs. For example, <u>Fischer (2004</u>) found occasions when O_3 at
4	the base of Mt. Washington during the night was much higher than typically observed,
5	and closer to those observed at the summit of Mt. Washington. They suggested that
6	mechanically driven turbulence due to wind shear caused O ₃ from aloft to penetrate the
7	stable nocturnal inversion thus causing O ₃ to increase near the base of Mt. Washington.
8	The high wind speeds causing this mechanically driven turbulence could have resulted
9	from the development of a LLJ. Stratospheric intrusions and intercontinental transport of
10	O_3 are also important and are covered in Section <u>3.4</u> in relation to background
11	concentrations.

3.2.1 Sources of Precursors Involved in Ozone Formation

12	Emissions of O ₃ precursor compounds (NO _X , VOCs, and CO) can be divided into natural
13	and anthropogenic source categories. Natural sources can be further divided into biogenic
14	from vegetation, microbes, and animals, and abiotic from biomass combustion, lightning,
15	and geogenic sources. However, the distinction between natural and anthropogenic
16	sources is often difficult to make in practice, as human activities directly or indirectly
17	affect emissions from what would have been considered natural sources during the
18	preindustrial era. Thus, emissions from plants and animals used in agriculture have been
19	referred to as anthropogenic or biogenic in different applications. Wildfire emissions can
20	be considered natural, except that forest management practices can lead to buildup of
21	fuels on the forest floor, thereby altering the frequency and severity of forest fires.
22	Estimates of emissions for NO _X , VOCs, and CO from the 2005 National Emissions
23	Inventory (NEI) (U.S. EPA, 2008a) are shown in Figure 3-2 to provide a general
24	indication of the relative importance of the different sources in the U.S. as a whole. The
25	magnitudes of the sources are strongly location and time dependent and so should not be
26	used to apportion sources of exposure. Shown in Figure 3-2 are Tier 1 categories. The
27	miscellaneous category can be quite large compared to total emissions, especially for CO
28	and VOCs. The miscellaneous category includes agriculture and forestry, wildfires,
29	prescribed burns, and a much more modest contribution from structural fires.



Note: NO_X (top), VOCs (middle), and CO (bottom) in the U.S. in million metric tons (MT) per year. Source: U.S. EPA (2008a).

Figure 3-2 Estimated anthropogenic emissions of ozone precursors for 2005.

1	Anthropogenic NO_X emissions are associated with combustion processes. Most emissions
2	are in the form of NO, which is formed at high combustion temperatures from
3	atmospheric nitrogen (N_2) and oxygen (O_2) and from fuel nitrogen (N) . According to the
4	2005 NEI, the largest sources of NO_X are on- and off-road (such as construction
5	equipment, agricultural equipment, railroad trains, ships, and aircraft) mobile sources and
6	electric power generation plants. Emissions of NO _X therefore are highest in areas having
7	a high density of power plants and in urban regions having high traffic density. Dallmann

1	and Harley (2010) compared NO _x emissions estimates from the 2005 NEI mobile sector
2	data with an alternative method based on fuel consumption and found reasonable
3	agreement in total U.S. anthropogenic emissions between the two techniques (to within
4	about 5%). However, emissions from on-road diesel engines in the fuel based inventory
5	constituted 46% of total mobile source NO_X compared to 35% in the EPA inventory. As a
6	result, emissions from on-road diesel engines in the fuel based approach are even larger
7	than electric power generation as estimated in the 2005 NEI, and on-road diesel engines
8	might represent the largest single NO_X source category. Differences between the two
9	techniques are largely accounted for by differences in emissions from on-road gasoline
10	engines. Uncertainties in the fuel consumption inventory ranged from 3% for on-road
11	gasoline engines to 20% for marine sources, and in the EPA inventory uncertainties
12	ranged from 16% for locomotives to 30% for off-road diesel engines. It should be noted
13	that the on-road diesel engine emissions estimate by Dallmann and Harley (2010) is still
14	within the uncertainty of the EPA estimate (22%). Because of rapid changes to heavy
15	duty diesel NO _X controls, emissions are likely to also rapidly change.
16	Satellite-based techniques have been used to obtain tropospheric concentrations of O ₃
17	precursors (e.g., NO ₂ , VOCs and CO). Such satellite-based measurements provide a
18	large-scale picture of spatial and temporal distribution of NO2, VOCs and CO that can be
19	used to evaluate emissions inventories produced using the bottom-up approach and to
20	produce top-down emissions inventories of these species. Although there are
21	uncertainties associated with satellite-based measurements, several studies have shown
22	the utility of top-down constraints on the emissions of O_3 precursors (McDonald-Buller et
23	al., 2011 and references therein). Following mobile sources, power plants are considered
24	the second largest anthropogenic source of NO _X . Over the past decade, satellite
25	measurements have shown appreciable reductions in NO_X power plant emissions across
26	the U.S. as a result of emission abatement strategies (Stavrakou et al., 2008; Kim et al.,
27	<u>2006</u>). For instance, <u>Kim et al. (2006</u>) observed a 34% reduction in NO _X emission over
28	the Ohio River Valley from 1999-2006 due to such strategies. Based on these results, less
29	than 25% of anthropogenic NO_X emissions were expected to originate from power plants
30	in this region. Uncertainty in NO _X satellite measurements are impacted by several factors,
31	such as cloud and aerosol properties, surface albedo, stratospheric NO _X concentration,
32	and solar zenith angle. Boersma et al. (2004) estimated an overall uncertainty between
33	35-60% for satellite-retrieved NO_X measurements in urban, polluted regions. Although
34	trends in satellite-retrieved NO_X power plant emissions reported by <u>Kim et al. (2006</u>) are
35	uncertain to some extent, similar reductions were reported by region-wide power plant
36	measurements (e.g., Continuous Emission Monitoring System observations, CEMS).
37	Major natural sources of NO_X in the U.S. include lightning, soils, and wildfires.
38	Uncertainties in natural NO _x emissions are much larger than for anthropogenic NO _x

- 1 emissions. Fang et al. (2010) estimated lightning generated NO_X of ~0.6 MT for July 2 2004. This value is \sim 40% of the anthropogenic emissions for the same period, but the 3 authors estimated that \sim 98% is formed in the free troposphere and so contributions to the 4 surface NO_x burden are low because most of this NO_x is oxidized to nitrate containing 5 species during downward transport into the planetary boundary layer. The remaining 2% 6 is formed within the planetary boundary layer. Both nitrifying and denitrifying organisms 7 in the soil can produce NO_x, mainly in the form of NO. Emission rates depend mainly on 8 fertilization amount and soil temperature and moisture. Nationwide, about 60% of the 9 total NO_x emitted by soils is estimated to occur in the central corn belt of the U.S. Spatial 10 and temporal variability in soil NO_x emissions leads to considerable uncertainty in 11 emissions estimates. However, these emissions are relatively low, only ~0.97 MT/year, or 12 about 6% of anthropogenic NO_x emissions. However, these emissions occur mainly 13 during summer when O_3 is of most concern and occur across the entire country including 14 areas where anthropogenic emissions are low.
- 15 Hundreds of VOCs, containing mainly 2 to ~ 12 carbon (C) atoms, are emitted by 16 evaporation and combustion processes from a large number of anthropogenic sources. 17 The two largest anthropogenic source categories in the U.S. EPA's emissions inventories 18 are industrial processes and transportation. Emissions of VOCs from highway vehicles 19 account for roughly two-thirds of the transportation-related emissions. The accuracy of 20 VOC emission estimates is difficult to determine, both for stationary and mobile sources. 21 Evaporative emissions, which depend on temperature and other environmental factors, 22 compound the difficulties of assigning accurate emission factors. In assigning VOC 23 emission estimates to the mobile source category, models are used that incorporate 24 numerous input parameters (e.g., type of fuel used, type of emission controls, and age of 25 vehicle), each of which has some degree of uncertainty.
- 26 On the U.S. and global scales, emissions of VOCs from vegetation are much larger than 27 those from anthropogenic sources. Emissions of VOCs from anthropogenic sources in the 28 2005 NEI were ~17 MT/year (wildfires constitute ~1/6 of that total and were included in 29 the 2005 NEI under the anthropogenic category, but see Section 3.4 for how wildfires are 30 treated for background O₃ considerations), but were 29 MT/year from biogenic sources. 31 Uncertainties in both biogenic and anthropogenic VOC emission inventories prevent 32 determination of the relative contributions of these two categories, at least in many areas. 33 Vegetation emits substantial quantities of VOCs, such as terpenoid compounds (isoprene, 34 2-methyl-3-buten-2-ol, monoterpenes), compounds in the hexanal family, alkenes, 35 aldehydes, organic acids, alcohols, ketones, and alkanes. The major chemicals emitted by 36 plants are isoprene (40%), other terpenoid and sesqui-terpenoid compounds (25%) and 37 the remainder consists of assorted oxygenated compounds and hydrocarbons according to 38 the 2005 NEI. Most biogenic emissions occur during the summer because of their

- dependence on temperature and incident sunlight. Biogenic emissions are also higher in southern states than in northern states for these reasons and because of species variations. The uncertainty in natural emissions is about 50% for isoprene under midday summer conditions and could be as much as a factor of ten higher for some compounds (Guenther et al., 2000). In EPA's regional modeling efforts, biogenic emissions of VOCs are estimated using the Biogenic Emissions Inventory System (BEIS) model (U.S. EPA, 2010b) with data from the Biogenic Emissions Landcover Database (BELD) and annual meteorological data. However, other emissions models are used such as Model of Emissions of Gases and Aerosols from Nature (MEGAN) (Guenther et al., 2006), especially in global modeling efforts.
- 11 Satellite measurements of HCHO, produced by the oxidation of isoprene and other 12 VOCs, have also been used to estimate biogenic VOC emissions attributed to isoprene 13 (Millet et al., 2008; Millet et al., 2006). Millet et al. (2008) demonstrated that both 14 satellite-based and model techniques capture the spatial variability of biogenic isoprene emissions in the U.S. reasonably well (satellite vs. MEGAN isoprene estimates, $R^2 = 0.48$ 15 16 or 0.68 depending on vegetation data base used). However, MEGAN tends to 17 overestimate emissions compared to satellite-based measurements. The uncertainty in 18 satellite derived isoprene emissions is roughly 40%, based on combined uncertainty in 19 satellite retrieval and isoprene yield from isoprene oxidation (Millet et al., 2006), which 20 is similar to the error associated with model-based techniques (~50%) (e.g., Millet et al., 21 2006; Guenther et al., 2000).
- 22 Anthropogenic CO is emitted primarily by incomplete combustion of carbon-containing 23 fuels. In general, any increase in fuel oxygen content, burn temperature, or mixing time in 24 the combustion zone will tend to decrease production of CO relative to CO₂. However, it 25 should be noted that controls mute the response of CO formation to fuel-oxygen. CO 26 emissions from large fossil-fueled power plants are typically very low since the boilers at 27 these plants are tuned for highly efficient combustion with the lowest possible fuel 28 consumption. Additionally, the CO-to- CO_2 ratio in these emissions is shifted toward CO_2 29 by allowing time for the furnace flue gases to mix with air and be oxidized by OH to CO_2 30 in the hot gas stream before the OH concentrations drop as the flue gases cool. 31 Nationally, on-road mobile sources constituted about half of total CO emissions in the 32 2005 NEI. When emissions from non-road vehicles are included, it can be seen from 33 Figure 3-2 that all mobile sources accounted for about three-quarters of total 34 anthropogenic CO emissions in the U.S.
- Analyses by <u>Harley et al. (2005)</u> and <u>Parrish (2006)</u> are consistent with the suggestion in
 Pollack et al. (2004) that the EPA MOBILE6 vehicle emissions model (U.S. EPA, 2010d)
 overestimates vehicle CO emissions by a factor of ~2. Field measurements by <u>Bishop and</u>

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1	Stedman (2008) were in accord with the Parrish (2006) findings that the measured trends
2	of CO and NO_X concentrations from mobile sources in the U.S. indicated that modeled
3	CO emission estimates were substantially too high. Hudman et al. (2008) found that the
4	NEI overestimated anthropogenic CO emissions by 60% for the eastern U.S. during the
5	period July 1-August 15, 2004 based on comparison of aircraft observations of CO from
6	the International Consortium for Atmospheric Research on Transport and Transformation
7	(ICARTT) campaign (Fehsenfeld et al., 2006) and results from a tropospheric chemistry
8	model (GEOS-Chem). Improvements in emissions technologies not correctly represented
9	in MOBILE emission models have been suggested as one cause for this discrepancy. For
10	example, Pokharel et al. (2003, 2002) demonstrated substantial decrements in the CO
11	fraction of tailpipe exhaust in several U.S. cities and Burgard et al. (2006) documented
12	improvements in emission from heavy-duty on-road diesel engines. Some of the largest
13	errors in the MOBILE models are addressed in the successor model, MOVES (U.S. EPA,
14	<u>2011e</u>).
15	Estimates of biogenic CO emissions in the 2005 NEI are made in a manner similar to that
16	for VOCs. National biogenic emissions, excluding fires, were estimated to contribute
17	~7% and wildfires added another ~16% to the national CO emissions total.
18	Photodecomposition of organic matter in oceans, rivers, lakes, and other surface waters,
19	and from soil surfaces also releases CO (Goldstein and Galbally, 2007). However, soils
20	can act as a CO source or a sink depending on soil moisture, UV flux reaching the soil
21	surface, and soil temperature (Conrad and Seiler, 1985). Soil uptake of CO is driven by
22	anaerobic bacteria (Inman et al., 1971). Emissions of CO from soils appear to occur by
23	abiotic processes, such as thermodecomposition or photodecomposition of organic
24	matter. In general, warm and moist conditions found in most soils favor CO uptake,
25	whereas hot and dry conditions found in deserts and some savannas favor the release of
26	CO (<u>King, 1999</u>).

3.2.2 Gas Phase Reactions Leading to Ozone Formation

27	Photochemical processes involved in O ₃ formation have been extensively reviewed in a
28	number of books (Jacobson, 2002; Jacob, 1999; Seinfeld and Pandis, 1998; Finlayson-
29	Pitts and Pitts, 1986) and in the 1996 and 2006 O ₃ AQCDs (U.S. EPA, 2006b, 1996a).
30	The photochemical formation of O_3 in the troposphere proceeds through the oxidation of
31	NO to nitrogen dioxide (NO ₂) by organic-peroxy (RO ₂) or hydro-peroxy (HO ₂) radicals.
32	The peroxy radicals oxidizing NO to NO_2 are formed during the oxidation of VOCs as
33	presented in Annex AX2.2.2 of the 2006 O ₃ AQCD (U.S. EPA, 2006b). The photolysis of
34	NO_2 yields NO and a ground-state oxygen atom, $O(^{3}P)$, which then reacts with molecular
35	oxygen to form O ₃ .

1VOCs important for the photochemical formation of O3 include alkanes, alkenes,2aromatic hydrocarbons, carbonyl compounds (e.g., aldehydes and ketones), alcohols,3organic peroxides, and halogenated organic compounds (e.g., alkyl halides). This array of4compounds encompasses a wide range of chemical properties and lifetimes: isoprene has5an atmospheric lifetime of approximately an hour, whereas methane has an atmospheric6lifetime of about a decade.

7 In urban areas, compounds representing all classes of VOCs and CO are important for O₃ 8 formation. In non-urban vegetated areas, biogenic VOCs emitted from vegetation tend to 9 be the most important. In the remote troposphere, methane (CH_4) and CO are the main 10 carbon-containing precursors to O_3 formation. The oxidation of VOCs is initiated mainly by reaction with hydroxyl (OH) radicals. The primary source of OH radicals in the 11 atmosphere is the reaction of electronically excited oxygen atoms, $O(^{1}D)$, with water 12 13 vapor. $O(^{1}D)$ is produced by the photolysis of O_{3} in the Hartley bands. In polluted areas, the photolysis of aldehydes (e.g., HCHO), HONO and H₂O₂ can also be appreciable 14 15 sources of OH, or HO_2 radicals that can rapidly be converted to OH (Eisele et al., 1997). 16 Ozone can oxidize alkenes, as can NO₃ radicals. NO₃ radicals are most effective at night 17 when they are most abundant. In coastal environments and other selected environments, 18 atomic Cl and Br radicals can also initiate the oxidation of VOCs as discussed in Annex 19 AX2.2.3 of the 2006 O₃ AQCD (U.S. EPA, 2006b). It was also emphasized in Annex 20 AX2.2.9 of the 2006 O₃ AQCD (U.S. EPA, 2006b) that the reactions of oxygenated 21 VOCs are important components of O_3 formation. They may be present in ambient air not 22 only as the result of the atmospheric oxidation of hydrocarbons but also by direct 23 emissions. For example, motor vehicles (including compressed natural gas vehicles) and 24 some industrial processes emit formaldehyde (Rappenglück et al., 2009) and vegetation 25 emits methanol.

26 There are a large number of oxidized N-containing compounds in the atmosphere 27 including NO, NO₂, NO₃, HNO₂, HNO₃, N₂O₅, HNO₄, PAN and its homologues, other 28 organic nitrates, such as alkyl nitrates, isoprene nitrates, and particulate nitrate. 29 Collectively these species are referred to as NO_Y. Oxidized nitrogen compounds are 30 emitted to the atmosphere mainly as NO which rapidly interconverts with NO₂ and so NO 31 and NO_2 are often "lumped" together into their own group or family, which is referred to 32 as NO_x. All the other species mentioned above in the definition of NO_y are products of 33 NO_X reactions are referred to as NO_7 , such that $NO_Y = NO_X + NO_7$. The major reactions 34 involving interconversions of oxidized N species were covered in Annex AX2.2.4 of the 35 2006 O₃ AQCD (U.S. EPA, 2006b). Mollner et al. (2010) identified pernitrous acid 36 (HOONO), an unstable isomer of nitric acid, as a product of the major gas phase reaction 37 forming HNO₃. However, since pernitrous acid is unstable, it is not a substantial reservoir

1	for NO _x . This finding stresses the importance of identifying products in addition to
2	measuring the rate of disappearance of reactants in kinetic studies.
3	The photochemical cycles by which the oxidation of hydrocarbons leads to O ₃ production
4	are best understood by considering the oxidation of methane, structurally the simplest
5	VOC. The CH ₄ oxidation cycle serves as a model for the chemistry of the relatively clean
6	or unpolluted troposphere (although this is a simplification because vegetation releases
7	large quantities of complex VOCs, such as isoprene, into the atmosphere). In the polluted
8	atmosphere, the underlying chemical principles are the same, as discussed in Annex
9	AX2.2.5 of the 2006 O_3 AQCD (U.S. EPA, 2006b). The conversion of NO to NO_2
10	occurring with the oxidation of VOCs is accompanied by the production of O_3 and the
11	efficient regeneration of the OH radical, which in turn can react with other VOCs as
12	shown in <u>Figure 3-1</u> .
13	The oxidation of alkanes and alkenes in the atmosphere has been treated in depth in the
14	1996 O ₃ AQCD (U.S. EPA, 1996a) and was updated in Annexes AX2.2.6 and AX2.2.7
15	of the 2006 O_3 AQCD (U.S. EPA, 2006b). In contrast to simple hydrocarbons containing
16	one or two C atoms, detailed kinetic information about the gas phase oxidation pathways
17	of many anthropogenic hydrocarbons (e.g., aromatic compounds such as benzene and
18	toluene), biogenic hydrocarbons (e.g., isoprene, the monoterpenes), and their
19	intermediate oxidation products (e.g., peroxides, nitrates, carbonyls and epoxides) is
20	lacking. This information is crucial even for compounds formed in low yields, such as
21	isoprene epoxides, as they are major precursors to secondary organic aerosol formation
22	(see, e.g., Surratt et al., 2010). Reaction with OH radicals represents the major loss
23	process for alkanes. Reaction with chlorine (Cl) atoms is an additional sink for alkanes.
24	Stable products of alkane photooxidation are known to include a wide range of
25	compounds and concentrations including carbonyl compounds, alkyl nitrates, and
26	d-hydroxycarbonyls. Major uncertainties in the atmospheric chemistry of the alkanes
27	concern the chemistry of alkyl nitrate formation; these uncertainties affect the amount of
28	NO-to-NO ₂ conversion occurring and, hence, the amounts of O ₃ formed during
29	photochemical degradation of the alkanes.
30	The reaction of OH radicals with aldehydes produced during the oxidation of alkanes
31	forms acyl (R'CO) radicals, and acyl peroxy radicals (R'C(O) $-O_2$) are formed by the
32	further addition of O_2 . As an example, the oxidation of ethane (C_2H_5 –H) yields
33	acetaldehyde (CH ₃ -CHO). The reaction of CH ₃ -CHO with OH radicals yields acetyl
34	radicals (CH ₃ -CO). The acetyl radicals will then participate with O_2 in a termolecular
35	recombination reaction to form acetyl peroxy radicals, which can then react with NO to
36	form $CH_3 + CO_2$ or they can react with NO_2 to form PAN. PAN acts as a temporary

1 2	reservoir for NO ₂ . Upon the thermal decomposition of PAN, either locally or elsewhere, NO ₂ is released to participate in the O_3 formation process again.
3	Alkenes react in ambient air with OH, NO ₃ , and Cl radicals and with O ₃ . All of these
4	reactions are important atmospheric transformation processes, and all proceed by initial
5	addition to the carbon double bonds. Major products of alkene photooxidation include
6	carbonyl compounds. Hydroxynitrates and nitratocarbonyls, and decomposition products
7	from the energy-rich biradicals formed in alkene- O_3 reactions are also produced. Major
8	uncertainties in the atmospheric chemistry of the alkenes concern the products and
9	mechanisms of their reactions with O_3 , especially the yields of radicals that participate in
10	O_3 formation. Examples of oxidation mechanisms of complex alkanes and alkenes can be
11	found in comprehensive texts such as Seinfeld and Pandis (1998).
12	Although the photochemistry of isoprene is crucial for understanding O ₃ formation, there
13	are major uncertainties in its oxidation pathways that still need to be addressed. Apart
14	from the effects of the oxidation of isoprene on production of radicals and O ₃ formation,
15	isoprene nitrates (RONO ₂) appear to play an important role as NO _X reservoirs over the
16	eastern U.S. (e.g., Perring et al., 2009). Their decomposition leads to the recycling of
17	NO_X , which can participate in the O_3 formation process. Laboratory and field-based
18	approaches support yields for RONO ₂ formation from isoprene oxidation ranging from 4
19	to 12% (see summaries in, Lockwood et al., 2010; Perring et al., 2009; Horowitz et al.,
20	<u>2007</u> ; <u>von Kuhlmann et al., 2004</u>). The rate at which RONO ₂ reacts to recycle NO _X is
21	poorly understood (Archibald et al., 2010; Paulot et al., 2009) with ranges from 0 to
22	100% in global chemical transport models. This range affects the sign of the O ₃ response
23	to changes in biogenic VOC emissions as well as the sensitivity of O_3 to changes in NO_X
24	emissions (Archibald et al., 2011; Ito et al., 2009; Weaver et al., 2009; Horowitz et al.,
25	2007; Fiore et al., 2005). In models that assume zero RONO ₂ recycling (<u>Zhang et al.</u> ,
26	2011; Wu et al., 2007; Fiore et al., 2003) O ₃ production is suppressed relative to a model
27	that recycles NO_X from $RONO_2$ (Kang et al., 2003). A related issue concerns the lack of
28	regeneration of $OH + HO_2$ radicals especially in low NO_X (<~1 ppb) environments. The
29	isomerization of the isoprene peroxy radicals that are formed after initial OH attack and
30	subsequent reactions could help resolve this problem (Peeters and Müller, 2010; Peeters
31	et al., 2009) and result in increases in OH concentrations from 20 to 40% over the
32	southeastern U.S. (Archibald et al., 2011). However, the effectiveness of this pathway is
33	uncertain and depends on the fraction of isoprene-peroxy radicals reacting by
34	isomerization. Crounse et al. (2011) estimated that only 8-11% of the isoprene-peroxy
35	radicals isomerizes to reform HO ₂ radicals. <u>Hofzumahaus et al. (2009</u>) also found under
36	predictions of OH in the Pearl River Delta and they also note that the sequence of
37	reactions beginning with OH attack on VOCs introduces enormous complexity which is
38	far from being fully understood.

- 1 The oxidation of aromatic hydrocarbons constitutes an important component of the 2 chemistry of O_3 formation in urban atmospheres as discussed in Annex AX2.2.8 of the 3 2006 O₃ AQCD (U.S. EPA, 2006b). Virtually all of the important aromatic hydrocarbon 4 precursors emitted in urban atmospheres are lost through reaction with the hydroxyl 5 radical. Loss rates for these compounds vary from slow (e.g., benzene) to moderate 6 (e.g., toluene), to very rapid (e.g., xylene and trimethylbenzene isomers). However, the 7 mechanism for the oxidation of aromatic hydrocarbons following reaction with OH is 8 poorly understood, as is evident from the poor mass balance of the reaction products. The 9 mechanism for the oxidation of toluene has been studied most thoroughly, and there is 10 general agreement on the initial steps in the mechanism. However, at present there is no 11 promising approach for resolving the remaining issues concerning the later steps. The 12 oxidation of aromatic hydrocarbons also leads to particle formation that could remove 13 gas-phase constituents that participate in O₃ formation.
- 14 Adequate analytical techniques needed to identify and quantify key intermediate species 15 are not available for many compounds. In addition, methods to synthesize many of the 16 suspected intermediate compounds are not available so that laboratory studies of their 17 reaction kinetics cannot be performed. Similar considerations apply to the oxidation of 18 biogenic hydrocarbons besides isoprene. These considerations are important because 19 oxidants, other than O_3 , that are formed from the chemistry described above could exert 20 effects on human health and perhaps also on vegetation (Doyle et al., 2007; Doyle et al., 21 2004; Sexton et al., 2004). Gas phase oxidants include PAN, H₂O₂, CH₃OOH, and other 22 organic hydroperoxides.
- 23Ozone is lost through a number of gas phase reactions and deposition to surfaces. The24reaction of O_3 with NO to produce NO_2 , for example in urban centers near roads, mainly25results in the recycling of O_3 downwind via the recombination of $O({}^3P)$ with O_2 to re-26form O_3 . By itself, this reaction does not lead to a net loss of O_3 unless the NO_2 is27converted to stable end products such as HNO_3 . Ozone reacts with unsaturated28hydrocarbons and with OH and HO_2 radicals.
- 29 Perhaps the most recent field study aimed at obtaining a better understanding of 30 atmospheric chemical processes was the Second Texas Air Quality Field Study 31 (TexAQS-II) conducted in Houston in August and September 2006 (Olaguer et al., 2009). 32 The TexAQS-II Radical and Aerosol Measurement Project (TRAMP) found evidence for 33 the importance of short-lived radical sources such as HCHO and HONO in increasing O₃ 34 productivity. During TRAMP, daytime HCHO pulses as large as 32 ppb were observed 35 and attributed to industrial activities upwind in the Houston Ship Channel (HSC) and 36 HCHO peaks as large as 52 ppb were detected by in situ surface monitors in the HSC. 37 Primary HCHO produced in flares from local refineries and petrochemical facilities could

1	increase peak O_3 by ~30 ppb (Webster et al., 2007). Other findings from TexAQS-II
2	included substantial concentrations of HONO during the day, with peak concentrations
3	approaching 1 ppb at local noon. These concentrations are well in excess of current air
4	quality model predictions using gas phase mechanisms alone (Sarwar et al., 2008) and
5	multiphase processes are needed to account for these observations. Olaguer et al. (2009)
6	also noted that using measured HONO brings modeled O_3 concentrations into much
7	better agreement with observations and could result in the production of an additional
8	10 ppb O ₃ . Large nocturnal vertical gradients indicating a surface or near-surface source
9	of HONO, and large concentrations of night-time radicals (~30 ppt HO ₂) were also found
10	during TRAMP.

3.2.3 Multiphase Processes

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In addition to the gas phase, chemical reactions also occur on the surfaces of or within cloud droplets and airborne particles. Their collective surface area is huge, implying that collisions with gas phase species occur on very short time scales. In addition to hydrometeors (e.g., cloud and fog droplets and snow and ice crystals) there are also potential reactions involving atmospheric particles of varying composition (e.g., wet [deliquesced] inorganic particles, mineral dust, carbon chain agglomerates and organic carbon particles). Multiphase reactions are involved in the formation of a number of species such as particulate nitrate, and gas phase HONO that can act to both increase and reduce the rate of O_3 formation in the polluted troposphere. Data collected in Houston as part of TexAQS-II summarized by <u>Olaguer et al. (2009</u>) indicate that concentrations of HONO are much higher than can be explained by gas phase chemistry and by tailpipe emissions. Photolysis of HONO formed in multiphase reactions in addition to the other sources can help to reduce the model underestimate of simulated O_3 in Houston.

24 Multiphase processes have been associated with the release of gaseous halogen 25 compounds from marine aerosol, mainly in marine and coastal environments. However, 26 Thornton et al. (2010) found production rates of gaseous nitryl chloride near Boulder, CO 27 from reaction of N₂O₅ with particulate Cl⁻, similar to those found in coastal and marine 28 environments. ClNO₂ readily photolyzes to yield Cl. They also found that substantial 29 quantities of N₂O₅ are recycled through ClNO₂ back into NO_x instead of forming HNO₃ 30 (a stable reservoir for reactive nitrogen compounds). The oxidation of hydrocarbons by 31 Cl radicals released from the marine aerosol could lead to the rapid formation of peroxy 32 radicals and higher rates of O₃ production. It should be noted that in addition to 33 production from marine aerosol, reactive halogen species are also produced by the 34 oxidation of halogenated organic compounds (e.g., CH₃Cl, CH₃Br, and CH₃I). The 35 atmospheric chemistry of halogens is complex because Cl, Br and I containing species

1	can react among themselves and with hydrocarbons and other species and could also be
2	important for O ₃ destruction, as has been noted for the lower stratosphere (McElroy et al.,
3	<u>1986; Yung et al., 1980</u>). For example, the reactions of Br and Cl containing radicals
4	deplete O_3 in selected environments such as the Arctic during the spring (Barrie et al.,
5	<u>1988</u>), the tropical marine boundary layer (<u>Dickerson et al., 1999</u>), and inland salt flats
6	and salt lakes (Stutz et al., 2002). Mahajan et al. (2010) found that I and Br species acting
7	together resulted in O_3 depletion that was much larger than would have been expected if
8	they acted individually and did not interact with each other; see Annex AX2.2.10.3 of the
9	2006 O ₃ AQCD (<u>U.S. EPA, 2006b</u>).
10	Multiphase processes have also been associated with the uptake of reactive gas phase
11	species affecting global budgets of O ₃ and nitrogen oxides among others. The uptake of
12	N_2O_5 on aerosols or cloud droplets leads to the loss of O_3 and NO_X and the production of
13	aqueous phase nitric acid, aerosol nitrate, and gaseous halogen nitrites. In addition to loss
14	of HO ₂ , the uptake of HO ₂ radicals on aerosol surfaces potentially reduces O ₃
15	concentrations and increases formation of sulfate (if H ₂ O ₂ is formed after uptake).
16	Macintyre and Evans (2011) developed a parameterization for uptake of HO ₂ based on
17	laboratory studies, which were about a factor of seven lower than previously estimated.
18	However they note that some of the earlier studies reporting higher values might have
19	been influenced by transition metal ions (e.g., Cu(II), Fe(II)), which are highly spatially
20	variable and could be important catalysts in areas with high concentrations of these ions.
21	Although the global change in O_3 was small (~-0.3%) much larger regional changes were
22	found (e.g., up to -27% at the surface over China).
23	Uptake coefficients for these species vary widely among laboratory studies. Macintyre
24	and Evans (2010) showed that the sensitivity of key tropospheric species such as O_3
25	varies from very small to significant over the range of uptake coefficients for N_2O_5
26	obtained in laboratory studies. For example, global O ₃ loss ranges from 0 to over 10%,
27	with large regional variability over the range of N_2O_5 uptake coefficients reported. In this
28	regard, it should be stressed that knowledge of multiphase processes is still evolving and
29	there are still many questions that remain to be answered. However, it is becoming clear
30	that multiphase processes are important for O_3 chemistry.
31	Reactions of O ₃ with monoterpenes have been shown to produce oxidants in the aerosol
32	phase, principally as components of ultrafine particles. Docherty et al. (2005) found
33	evidence for the substantial production of organic hydroperoxides in secondary organic
34	aerosol (SOA) resulting from the reaction of monoterpenes with O3. Analysis of the SOA
35	formed in their environmental chamber indicated that the SOA consisted mainly of
36	organic hydroperoxides. In particular, they obtained yields of 47% and 85% of organic
37	peroxides from the oxidation of α - and β -pinene. The hydroperoxides then react with

1	aldehydes in particles to form peroxyhemiacetals, which can either rearrange to form
2	other compounds such as alcohols and acids or revert back to the hydroperoxides. The
3	aldehydes are also produced in large measure during the ozonolysis of the monoterpenes.
4	Monoterpenes also react with OH radicals resulting in the production of more
5	lower-molecular-weight products than in the reaction with monoterpenes and O ₃ . Bonn et
6	al. (2004) estimated that hydroperoxides lead to 63% of global SOA formation from the
7	oxidation of terpenes. The oxidation of anthropogenic aromatic hydrocarbons by OH
8	radicals could also produce organic hydroperoxides in SOA (Johnson et al., 2004).
9	Recent measurements show that the abundance of oxidized SOA exceeds that of more
10	reduced hydrocarbon like organic aerosol in Pittsburgh (Zhang et al., 2005) and in about
11	30 other cities across the Northern Hemisphere (Zhang et al., 2007b). Based on aircraft
12	and ship-based sampling of organic aerosols over coastal waters downwind of
13	northeastern U.S. cities, de Gouw et al. (2008) reported that 40-70% of measured organic
14	mass was water soluble and estimated that approximately 37% of SOA is attributable to
15	aromatic precursors, using PM yields estimated for NO _X -limited conditions. Uncertainties
16	still exist as to the pathways by which the oxidation of isoprene leads to the formation of
17	SOA. Nozière et al. (2011) found that a substantial fraction of 2-methyltetrols are
18	primary in origin, although these species have been widely viewed solely as products of
19	the atmospheric oxidation of isoprene. This finding points to lingering uncertainty in
20	reaction pathways in the oxidation of isoprene and in estimates of the yield of SOA from
21	isoprene oxidation.
22	Departience of Q on the surfaces of norticles in norticular these with humis estimite

22	Reactions of O ₃ on the surfaces of particles, in particular those with humic acid like
23	composition, are instrumental in the processing of SOA and the release of
24	low-molecular-weight products such as HCHO (D'Anna et al., 2009). However, direct
25	reactions of O_3 and atmospheric particles appear to be too slow to represent a major O_3
26	sink in the troposphere (<u>D'Anna et al., 2009</u>).

3.2.3.1 Indoor Air

27	Except when activities such as photocopying or welding are occurring, the major source
28	of O_3 to indoor air is through infiltration of outdoor air. Reactions involving ambient O_3
29	with NO either from exhaled breath or from gas-fired appliances, surfaces of furnishings
30	and terpenoid compounds from cleaning products, air fresheners and wood products also
31	occur in indoor air as was discussed in the 2006 O_3 AQCD (U.S. EPA, 2006b). The
32	previous O3 review also noted that the ozonolysis of terpenoid compounds could be a
33	substantial source of secondary organic aerosol in the ultrafine size fraction. Chen et al.
34	(2011) examined the formation of secondary organic aerosol from the reaction of O_3 that
35	has infiltrated indoors with terpenoid components of commonly used air fresheners. They

1focused on the formation and decay of particle bound reactive oxygen species (ROS) and2on their chemical properties. They found that the ROS content of samples can be3decomposed into fractions that differ in terms of reactivity and volatility; however, the4overall ROS content of samples decays and over 90% is lost within a day at room5temperature. This result also suggests loss of ROS during sampling periods longer than a6couple of hours.

3.2.4 Temperature and Chemical Precursor Relationships

7 As might be expected based on the temperature dependence of many reactions involved 8 in the production and destruction of O_3 and the temperature dependence of emissions 9 processes such as evaporation of hydrocarbon precursors and the emissions of 10 biogenically important precursors such as isoprene, ambient concentrations of O_3 also 11 show temperature dependence. Bloomer et al. (2009) determined the sensitivity of O_3 to 12 temperature at rural sites in the eastern U.S. They found that O₃ increased on average at 13 rural Clean Air Status and Trends Network (CASTNET) sites by ~3.2 ppbv/°C before 14 2002, and after 2002 by $\sim 2.2 \text{ ppby/}^{\circ}\text{C}$. This change in sensitivity was largely the result of 15 reductions in NO_x emissions from power plants. These results are in accord with model 16 predictions by Wu et al. (2008b) showing that the sensitivity of O_3 to temperature 17 decreases with decreases in precursor emissions. Rasmussen et al. (2012) recently 18 extended the work of Bloomer et al. (2009) to quantify seasonal changes in the sensitivity 19 of O₃ to temperature as well as regional variability (3-6 ppb/°C over the Northeast and 20 mid-Atlantic; 3-4 ppb/°C over the Great Lakes region) and to evaluate the capability of a 21 chemistry-climate model to capture O₃ sensitivity to temperature. However, the 22 associations of O_3 with temperature are not as clear in the West as they might be in the 23 East. For example, sites downwind of Phoenix, AZ showed basically no sensitivity of O_3 to temperature ($R^2 = 0.02$) (U.S. EPA, 2006b). However, Wise and Comrie (2005) did 24 25 find that meteorological parameters (mixing height and temperature) typically accounts 26 for 40 to 70% of the variability in O_3 in the five southwestern cities (including Phoenix) 27 they examined. It is likely that differences in the nature of sites chosen (urban vs. rural) 28 accounted for this difference and are at least partially responsible for the difference in 29 results. Jaffe et al. (2008) regressed O_3 on temperature at Yellowstone and Rocky Mountain NP and found weak associations ($R^2 = 0.09$ and 0.16). They found that 30 31 associations with area burned by wildfires are much stronger. Other sources as discussed 32 in Section 3.4 might also be more important in the West than in the East. 33 The warmer months of the year are generally regarded as being the most conducive to 34 higher O_3 concentrations. However, Schnell et al. (2009) reported observations of high O_3

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concentrations (maximum 1-h avg of 140 ppb; maximum 8-h avg of 120 ppb) in the

1	Jonah-Pinedale gas fields in Wyoming during winter at temperatures of -17°C. Potential
2	factors contributing to these anomalously high concentrations include a highly reflective
3	snow surface, emissions of NO _X , hydrocarbons and short-lived radical reservoirs
4	(e.g., HONO and HCHO) and a very shallow, stable boundary layer trapping these
5	emissions close to the surface (Schnell et al., 2009). Multiphase processes might also be
6	involved in the production of these short-lived reservoirs. At a temperature of -17°C, the
7	production of hydroxyl radicals (by the photolysis of O_3 yielding O^1D followed by the
8	reaction, $O(^{1}D) + H_{2}O$, needed to initiate hydrocarbon oxidation) is severely limited,
9	suggesting that another source of radicals is needed. Radicals can be produced by the
10	photolysis of molecules such as HONO and HCHO which photolyze in optically thin
11	regions of the solar spectrum. A similar issue, in part due to the under-prediction of
12	radicals, has arisen in the Houston airshed where chemistry-transport models (CTMs;
13	discussed further in Section $\underline{3.3}$) under-predict O ₃ (<u>Olaguer et al., 2009</u>). <u>Carter and</u>
14	Seinfeld (2012) modeled several of the events using the SAPRC-07 chemical mechanism
15	and found that the release of HONO from the snow surface aids in the formation of O_3 .
16	The chemical mechanism they used-including the temperature dependence of rate
17	coefficients-was developed for application at higher temperatures. They also note that
18	temperature changes will also affect the distribution of products and radicals formed
19	when individual VOCs react, but the current version of the mechanism represents these
20	by lumped overall processes in which the product and radical distributions are treated as
21	if they are temperature independent. It is not clear how this treatment of radical
22	production might affect their results.

23 Rather than varying directly with emissions of its precursors, O_3 changes in a nonlinear 24 fashion with the concentrations of its precursors. At the low NO_X concentrations found in 25 remote continental areas to rural and suburban areas downwind of urban centers (low-26 NO_X regime), the net production of O_3 typically increases with increasing NO_X . In the 27 low-NO_X regime, the overall effect of the oxidation of VOCs is to generate (or at least 28 not consume) free radicals, and O₃ production varies directly with NO_X. In the high-NO_X 29 regime, NO₂ scavenges OH radicals which would otherwise oxidize VOCs to produce 30 peroxy radicals, which in turn would oxidize NO to NO_2 . In this regime, O_3 production is 31 limited by the availability of free radicals and O_3 shows only a weak dependence on NO_x 32 concentrations. The production of free radicals is in turn limited by the availability of 33 solar UV radiation capable of photolyzing O_3 (in the Hartley bands) or aldehydes and/or 34 by the abundance of VOCs whose oxidation produce more radicals than they consume. 35 At even higher NO_X concentrations, as found in downtown metropolitan areas, especially 36 near busy streets and roads, and in power plant plumes, there is scavenging (titration) of 37 O₃ by reaction with NO.

1	There are a number of ways to refer to the chemistry in these chemical regimes.
2	Sometimes the terms VOC-limited and NO _X -limited are used. However, there are
3	difficulties with this usage because (1) VOC measurements are not as abundant as they
4	are for nitrogen oxides; (2) rate coefficients for reaction of individual VOCs with radicals
5	(e.g., OH, Cl) vary over an extremely wide range; and (3) consideration is not given to
6	CO nor to reactions that can produce radicals without involving VOCs. The terms NO _X -
7	limited and NO_X -saturated (<u>Jaegle et al., 2001</u>) will be used wherever possible to more
8	adequately describe these two regimes. However, the terminology used in original
9	articles will also be used here. In addition to these two regimes, there is also a "very low
10	NO_X regime" in the remote marine troposphere in which NO_X concentrations are ~20 ppt
11	or less. Under these very low NO_X conditions, which are not likely to be found in the
12	continental U.S, HO ₂ and CH ₃ O ₂ radicals react with each other and HO ₂ radicals undergo
13	self-reaction (to form H_2O_2), and OH and HO_2 react with O_3 , leading to net destruction of
14	O_3 and inefficient OH radical regeneration by comparison with much higher NO_X
15	concentrations found in polluted areas. In polluted areas, HO ₂ and CH ₃ O ₂ radicals react
16	with NO to convert NO to NO ₂ , regenerate the OH radical, and, through the photolysis of
17	NO ₂ , produce O ₃ as noted in Annex AX2.2.5 of the 2006 O ₃ AQCD (U.S. EPA, 2006b).
18	There are no sharp transitions between these regimes. For example, in the "low NO_X
19	regime" there still may be appreciable peroxy-peroxy radical reactions depending on the
20	local NO_X concentration. In any case, in all of these NO_X regimes, O_3 production is also
21	limited by the abundance of HO_X radicals.
22	The chemistry of OH radicals, which are responsible for initiating the oxidation of

- 23 hydrocarbons, shows behavior similar to that for O_3 with respect to NO_X concentrations 24 (Poppe et al., 1993; Zimmermann and Poppe, 1993; Hameed et al., 1979). These 25 considerations introduce a high degree of uncertainty into attempts to relate changes in 26 O_3 concentrations to emissions of precursors. There are no definitive rules governing the 27 concentrations of NO_x at which the transition from NO_x-limited to NO_x-saturated 28 conditions occurs. The transition between these two regimes is highly spatially and 29 temporally dependent and depends also on the nature and abundance of the hydrocarbons 30 that are present.
- 31 Trainer et al. (1993) and Olszyna et al. (1994) have shown that O_3 and NO_Y are highly 32 correlated in rural areas in the eastern U.S. Trainer et al. (1993) also showed that O₃ 33 concentrations correlate even better with NO_Z than with NO_Y, as may be expected 34 because NO_Z represents the amount of NO_X that has been oxidized, forming O₃ in the 35 process. NO_Z is equal to the difference between measured total reactive nitrogen (NO_Y) 36 and NO_X and represents the summed products of the oxidation of NO_X . NO_Z is composed 37 mainly of HNO₃, PAN and other organic nitrates, particulate nitrate, and HNO₄. Trainer 38 et al. (1993) also suggested that the slope of the regression line between O_3 and NO_2 can

1	be used to estimate the rate of O_3 production per NO _x oxidized (also known as the O_3
2	production efficiency [OPE]). Ryerson et al. (2001); Ryerson et al. (1998) used measured
3	correlations between O_3 and NO_Z to identify different rates of O_3 production in plumes
4	from large point sources. A number of studies in the planetary boundary layer over the
5	continental U.S. have found that the OPE ranges typically from 1 to nearly 10. However,
6	it may be higher in the upper troposphere and in certain areas, such as the Houston-
0 7	
	Galveston area in Texas. Observations indicate that the OPE depends mainly on the
8	abundance of NO_X and also on availability of solar UV radiation, VOCs and O_3 itself.
9	Various techniques have been proposed to use ambient $\ensuremath{\text{NO}_X}$ and $\ensuremath{\text{VOC}}$ measurements to
10	derive information about the dependence of O ₃ production on their concentrations. For
11	example, it has been suggested that O_3 formation in individual urban areas could be
12	understood in terms of measurements of ambient NO_X and VOC concentrations during
13	the early morning (<u>NRC, 1991</u>). In this approach, the ratio of summed (unweighted) VOC
14	to NO_X is used to determine whether conditions were NO_X -limited or VOC-limited. This
15	procedure is inadequate because it omits many factors that are important for O_3
16	production such as the impact of biogenic VOCs (which are typically not present in urban
17	centers during early morning); important differences in the ability of individual VOCs to
18	generate radicals (rather than just total VOC) and other differences in O ₃ forming
19	potential for individual VOCs (<u>Carter, 1995</u>); and changes in the VOC to NO _x ratio due
20	to photochemical reactions and deposition as air moves downwind from urban areas
21	(Milford et al., 1994).
22	
22	Photochemical production of O_3 generally occurs simultaneously with the production of
23	various other species such as HNO ₃ , organic nitrates, and other oxidants such as
24	hydrogen peroxide. The relative rate of production of O_3 and other species varies
25	depending on photochemical conditions, and can be used to provide information about
26	O_3 -precursor sensitivity. <u>Sillman (1995</u>) and <u>Sillman and He (2002</u>) identified several
27	secondary reaction products that show different correlation patterns for NO _X -limited and
28	NO_X -saturated conditions. The most important correlations are for O_3 versus NO_Y , O_3
29	versus NO_Z , O_3 versus HNO_3 , and H_2O_2 versus HNO_3 . The correlations between O_3 and
30	NO_Y , and O_3 and NO_Z are especially important because measurements of NO_Y and NO_X
31	are more widely available than for VOCs. Measured O_3 versus NO_Z (Figure 3-3) shows
32	distinctly different patterns in different locations. In rural areas and in urban areas such as

Los Angeles.

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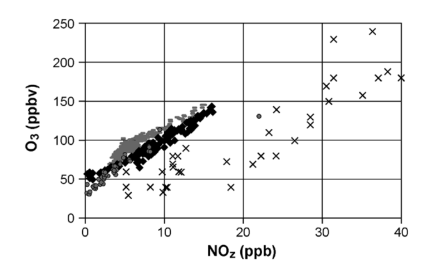
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37 38 Nashville, TN, O₃ is highly correlated with NO_Z. By contrast, in Los Angeles, CA, O₃ is

not as highly correlated with NO_Z, and the rate of increase of O₃ with NO_Z is lower and

the O₃ concentrations for a given NO_Z value are generally lower. The different O₃ versus

 NO_Z relations in Nashville, TN and Los Angeles, CA reflects the difference between NO_X -limited conditions in Nashville versus an approach to NO_X -saturated conditions in



Note: (NO_Y–NO_X) during the afternoon at rural sites in the eastern U.S. (grey circles) and in urban areas and urban plumes associated with Nashville, TN (gray dashes); Paris, France (black diamonds); and Los Angeles, CA (Xs). Source: Adapted with permission of American Geophysical Union, (<u>Sillman and He, 2002</u>; <u>Sillman et al., 1998</u>; <u>Trainer et al., 1993</u>).

Figure 3-3 Measured concentrations of ozone and NO_z.

1	The difference between NO _X -limited and NO _X -saturated regimes is also reflected in
2	measurements of H ₂ O ₂ . H ₂ O ₂ production is highly sensitive to the abundance of radicals
3	and is thus favored in the NO _X -limited regime. Measurements in the rural eastern U.S.
4	(Jacob et al., 1995), Nashville, TN (Sillman et al., 1998), and Los Angeles, CA
5	(Sakugawa and Kaplan, 1989), show large differences in H_2O_2 concentrations between
6	likely NO _x -limited and NO _x -saturated locations.
7	The applications of indicator species mentioned above are limited to individual urban
8	areas either because they are based on point measurements or by the range of the aircraft
9	carrying the measurement instruments. Satellites provide a platform for greatly extending
10	the range of applicability of the indicator technique and also have the resolution
11	necessary to examine urban to rural differences. Duncan et al. (2010) used satellite data
12	from Ozone Monitoring Instrument (OMI) for HCHO to NO2 column ratios to diagnose
13	NO _X -limited and radical-limited (NO _X -saturated) regimes. HCHO can be used as an
14	indicator of VOCs as it is a common, short-lived, oxidation product of many VOCs that
15	is a source of HO_X (Sillman, 1995). In adopting the satellite approach, CTMs are used to
16	estimate the fractional abundance of the indicator species in the planetary boundary layer.
17	Duncan et al. (2010) found that O_3 formation over most of the U.S. became more
18	sensitive to NO_X over most of the U.S. from 2005 to 2007 largely because of decreases in
19	NO_{X} emissions. They also found that surface temperature is correlated with the ratio of

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HCHO to NO_2 especially in cities in the Southeast where emissions of isoprene (a major source of HCHO) are high due to high temperatures in summer.

3.3 Atmospheric Modeling

CTMs have been widely used to compute the interactions among atmospheric pollutants and their transformation products, and the transport and deposition of pollutants. They have also been widely used to improve basic understanding of atmospheric chemical processes and to develop control strategies. The spatial scales over which pollutant fields are calculated range from intra-urban to regional to global. Generally, these models are applied to problems on different spatial scales but efforts are underway to link across spatial scales for dealing with global scale environmental issues that affect population health within cities. Many features are common to all of these models and hence they share many of the same problems. On the other hand, there are appreciable differences in approaches to parameterizing physical and chemical processes that must be addressed in applying these models across spatial scales.

14 CTMs solve a set of coupled, non-linear partial differential equations, or continuity 15 equations, for relevant chemical species. Jacobson (2005) described the governing partial 16 differential equations, and the methods that are used to solve them. Because of limitations 17 imposed by the complexity and spatial-temporal scales of relevant physical and chemical 18 processes, the CTMs must include parameterizations of these processes, which include 19 atmospheric transport; the transfer of solar radiation through the atmosphere; chemical 20 reactions; and removal to the surface by turbulent motions and precipitation. 21 Development of parameterizations for use in CTMs requires data for three dimensional 22 wind fields, temperatures, humidity, cloudiness, and solar radiation; emissions data for 23 primary (i.e., directly emitted from sources) species such as NO_X, SO₂, NH₃, VOCs, and 24 primary PM; and chemical reactions.

25 The domains of CTMs extend from a few hundred kilometers on a side to the entire 26 globe. Most major regional (i.e., sub-continental) scale air-related modeling efforts at 27 EPA rely on the Community Multi-scale Air Quality (CMAQ) modeling system (Byun 28 and Schere, 2006; Byun and Ching, 1999). CMAQ's horizontal domain typically extends 29 over North America with efforts underway to extend it over the entire Northern 30 Hemisphere. Note that CTMs can be 'nested' within each other as shown in Figure 3-4 31 which shows domains for CMAQ (Version 4.6.1); additional details on the model 32 configuration and application are found elsewhere (U.S. EPA, 2009e). The figure shows 33 the outer domain (36 km horizontal grid spacing) and two 12 km spatial resolution (east 34 and west) sub-domains. The upper boundary for CMAQ is typically set at about 100 hPa, or at about 16 km altitude on average, although in some recent applications the upper
 boundary has been set at 50 hPa. These domains and grid spacings are quite common and
 can also be found in a number of other models.



Note: Figure depicts a 36 km grid-spacing outer parent domain in black; 12 km western U.S. domain in red; 12 km eastern U.S. domain in blue.

Figure 3-4 Sample Community Multi-scale Air Quality (CMAQ) modeling domains.

4	The main components of a CTM such as EPA's CMAQ are summarized in Figure 3-5.
5	The capabilities of a number of CTMs designed to study local- and regional-scale air
6	pollution problems were summarized by <u>Russell and Dennis (2000</u>) and in the 2006 O_3
7	AQCD (U.S. EPA, 2006b). Historically, CMAQ has been driven most often by the MM5
8	mesoscale meteorological model (Seaman, 2000), though it could be driven by other
9	meteorological models including the Weather Research and Forecasting (WRF) model
10	and the Regional Atmospheric Modeling System (RAMS) (ATMET, 2011).

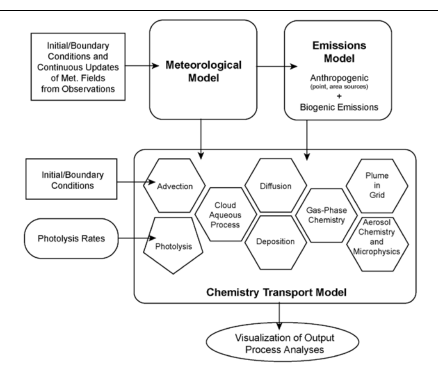


Figure 3-5 Main components of a comprehensive atmospheric chemistry modeling system, such as the U.S. EPA's Community Multi-scale Air Quality (CMAQ) modeling system.

1	Simulations of pollution episodes over regional domains have been performed with a
2	horizontal resolution down to 1 km; see the application and general survey results
3	reported in Ching et al. (2006). However, simulations at such high resolution require
4	better parameterizations of meteorological processes such as boundary layer fluxes, deep
5	convection, and clouds (Seaman, 2000). Finer spatial resolution is necessary to resolve
6	features such as urban heat island circulation; sea, bay, and land breezes; mountain and
7	valley breezes; and the nocturnal low-level jet; all of which can affect pollutant
8	concentrations. Other major air quality systems used for regional scale applications
9	include the Comprehensive Air Quality Model with extensions (CAMx) (ENVIRON,
10	2005) and the Weather Research and Forecast model with Chemistry (WRF/Chem)
11	(<u>NOAA, 2010</u>).
12	CMAQ and other grid-based or Eulerian air quality models subdivide the modeling
13	domain into a three-dimensional array of grid cells. The most common approach to
14	setting up the horizontal domain is to nest a finer grid within a larger domain of coarser
15	resolution. The use of finer horizontal resolution in CTMs will necessitate finer-scale
16	inventories of land use and better knowledge of the exact paths of roads, locations of
17	factories, and, in general, better methods for locating sources and estimating their

- 1 emissions. The vertical resolution of CTMs is variable and usually configured to have 2 more layers in the PBL and fewer in the free troposphere. 3 The meteorological fields are produced either by other numerical prediction models such 4 as those used for weather forecasting (e.g., MM5, WRF), and/or by assimilation of 5 satellite data. The flow of information shown in Figure 3-5 has most often been 6 unidirectional in the sense that information flows into the CTM (large box) from outside; 7 feedbacks on the meteorological fields and on boundary conditions (i.e., out of the box) 8 have not been included. However, CTMs now have the capability to consider these 9 feedbacks as well; see, for example, Binkowski et al. (2007) and WRF/Chem (NOAA, 10 2010). 11 Because of the large number of chemical species and reactions that are involved in the 12 oxidation of realistic mixtures of anthropogenic and biogenic hydrocarbons, condensed 13 mechanisms must be used in atmospheric models. These mechanisms can be tested by 14 comparison with smog chamber data. However, the existing chemical mechanisms often 15 neglect many important processes such as the formation and subsequent reactions of 16 long-lived carbonyl compounds, the incorporation of the most recent information about 17 intermediate compounds, and heterogeneous reactions involving cloud droplets and 18 aerosol particles. To the extent that information is available, models like CMAO and 19 CAMx do include state-of-the-science parameterization for some of these processes such 20 as heterogeneous N₂O₅ chemistry. 21 The initial conditions, or starting concentration fields of all species computed by a model, 22 and the boundary conditions, or concentrations of species along the horizontal and upper 23 boundaries of the model domain throughout the simulation, must be specified at the 24 beginning of the simulation. Both initial and boundary conditions can be estimated from 25 models or data or, more generally, model plus data hybrids. Because data for vertical 26 profiles of most species of interest are very sparse, results of model simulations over 27 larger, usually global, domains are often used. 28 Chemical kinetics mechanisms representing the important reactions occurring in the 29 atmosphere are used in CTMs to estimate the rates of chemical formation and destruction 30 of each pollutant simulated as a function of time. The Master Chemical Mechanism 31 (MCM) (Univ of Leeds, 2010) is a comprehensive reaction database providing as near an 32 explicit treatment of chemical reactions in the troposphere as is possible. The MCM 33 currently includes over 12,600 reactions and 4,500 species. However, mechanisms that 34 are this comprehensive are still computationally too demanding to be incorporated into 35 CTMs for regulatory use. Simpler treatments of tropospheric chemistry have been 36 assembled by combining chemical species into mechanisms that group together
 - compounds with similar chemistry. It should be noted that because of different

1	approaches to the lumping of organic compounds into surrogate groups for computational
2	efficiency, chemical mechanisms can produce different results under similar conditions.
3	Jimenez et al. (2003) briefly described the features of the seven main chemical
4	mechanisms in use and compared concentrations of several key species predicted by
5	these mechanisms in a box-model simulation over 24 hours. Several of these mechanisms
6	have been incorporated into CMAQ including extensions of the Carbon Bond (CB)
7	mechanism (Luecken et al., 2008), SAPRC (Luecken et al., 2008), and the Regional
8	Atmospheric Chemistry Mechanism, version 2 (RACM2) (Fuentes et al., 2007). The CB
9	mechanism is currently undergoing extension (CB06) to include, among other things,
10	longer lived species to better simulate chemistry in the remote and upper troposphere.
11	These mechanisms were developed primarily for homogeneous gas phase reactions and
12	treat multiphase chemical reactions in a very cursory manner, if at all. As a consequence
13	of neglecting multiphase chemical reactions, models such as CMAQ could have
14	difficulties capturing the regional nature of O ₃ episodes, in part because of uncertainty in
15	the chemical pathways converting NO_X to HNO_3 and recycling of NO_X (Godowitch et al.,
16	2008; Hains et al., 2008). Much of this uncertainty also involves multiphase processes as
17	described in Section $3.2.3$.
18	CMAQ and other CTMs incorporate processes and interactions of aerosol-phase
18 19	CMAQ and other CTMs incorporate processes and interactions of aerosol-phase chemistry (<u>Zhang and Wexler, 2008; Gaydos et al., 2007; Binkowski and Roselle, 2003</u>).
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19 20	chemistry (<u>Zhang and Wexler, 2008</u> ; <u>Gaydos et al., 2007</u> ; <u>Binkowski and Roselle, 2003</u>). There have also been several attempts to study the feedbacks of chemistry on
19 20 21	chemistry (<u>Zhang and Wexler, 2008; Gaydos et al., 2007; Binkowski and Roselle, 2003</u>). There have also been several attempts to study the feedbacks of chemistry on atmospheric dynamics using meteorological models like MM5 and WRF (<u>Liu et al.</u> ,
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19 20 21 22 23 24 25 26 27 28 29	 chemistry (Zhang and Wexler, 2008; Gaydos et al., 2007; Binkowski and Roselle, 2003). There have also been several attempts to study the feedbacks of chemistry on atmospheric dynamics using meteorological models like MM5 and WRF (Liu et al., 2001; Park et al., 2001; Grell et al., 2000; Lu et al., 1997). This coupling is necessary to accurately simulate feedbacks from PM (Park et al., 2001; Lu et al., 1997) over areas such as Los Angeles or the Mid-Atlantic region. Photolysis rates in CMAQ can now be calculated interactively with model produced O₃, NO₂, and aerosol fields (Binkowski et al., 2007). Spatial and temporal characterizations of anthropogenic and biogenic precursor emissions can be specified as inputs to a CTM or these emissions can be calculated in-line in CMAQ. Emissions inventories have been compiled on grids of varying resolution for
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19 20 21 22 23 24 25 26 27 28 29 30 31	 chemistry (Zhang and Wexler, 2008; Gaydos et al., 2007; Binkowski and Roselle, 2003). There have also been several attempts to study the feedbacks of chemistry on atmospheric dynamics using meteorological models like MM5 and WRF (Liu et al., 2001; Park et al., 2001; Grell et al., 2000; Lu et al., 1997). This coupling is necessary to accurately simulate feedbacks from PM (Park et al., 2001; Lu et al., 1997) over areas such as Los Angeles or the Mid-Atlantic region. Photolysis rates in CMAQ can now be calculated interactively with model produced O₃, NO₂, and aerosol fields (Binkowski et al., 2007). Spatial and temporal characterizations of anthropogenic and biogenic precursor emissions can be specified as inputs to a CTM or these emissions can be calculated in-line in CMAQ. Emissions inventories have been compiled on grids of varying resolution for many hydrocarbons, aldehydes, ketones, CO, NH₃, and NO_X. Preprocessing of emissions data for CMAQ is done by the Spare-Matrix Operator Kernel Emissions (SMOKE)

characteristic of the time of day and season. Appreciable errors in emissions can occur if inappropriate time dependence is applied.

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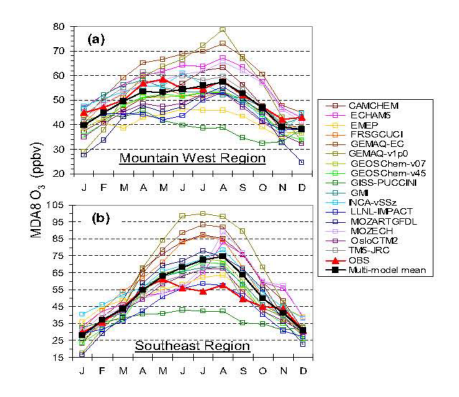
- 1 Each of the model components described above has associated uncertainties; and the 2 relative importance of these uncertainties varies with the modeling application. Large 3 errors in photochemical modeling arise from the meteorological, chemical and emissions 4 inputs to the model (Russell and Dennis, 2000). While the effects of poorly specified 5 boundary conditions propagate through the model's domain, the effects of these errors 6 remain undetermined. Because many meteorological processes occur on spatial scales 7 smaller than the model's vertical or horizontal grid spacing and thus are not calculated 8 explicitly, parameterizations of these processes must be used. These parameterizations 9 introduce additional uncertainty.
- 10 The performance of CTMs must be evaluated by comparison with field data as part of a 11 cycle of model evaluations and subsequent improvements (NRC, 2007). However, they 12 are too computationally demanding to have the full range of their sensitivities examined 13 using Monte Carlo techniques (NRC, 2007). Models of this complexity are evaluated by 14 comparison with field observations for O_3 and other species. Evaluations of the 15 performance of CMAQ are given in Arnold et al. (2003), Eder and Yu (2005), Appel et 16 al. (2005), and Fuentes and Raftery (2005). Discrepancies between model predictions and 17 observations can be used to point out gaps in current understanding of atmospheric chemistry and to spur improvements in parameterizations of atmospheric chemical and 18 19 physical processes. Model evaluation does not merely involve a straightforward 20 comparison between model predictions and the concentration field of the pollutant of 21 interest. Such comparisons may not be meaningful because it is difficult to determine if 22 agreement between model predictions and observations truly represents an accurate 23 treatment of physical and chemical processes in the CTM or the effects of compensating 24 errors in complex model routines (in other words, it is important to know if the right 25 answer is obtained for the right reasons). Each of the model components (emissions 26 inventories, chemical mechanism, and meteorological driver) should be evaluated 27 individually as has been done to large extent in some major field studies such as TexAQS 28 I and II and CalNex. Comparisons of correlations between measured and modeled VOCs 29 and NO_x are useful for evaluating results from CTMs and can provide information about 30 the chemical state of the atmosphere. A CTM that accurately computes both VOC and 31 NO_x along with the spatial and temporal relations among the critical secondary species 32 associated with O₃ has a higher probability of representing O₃-precursor relations 33 correctly than one that does not.
- The above evaluation techniques are sometimes referred to as "static" in the sense that individual model variables are compared to observations. It is also crucial to understand the dynamic response to changes in inputs and to compare the model responses to those that are observed. These tests might involve changes in some natural forcing or in emissions from an anthropogenic source. As an example, techniques such as the direct

decoupled method (DDM) (<u>Dunker et al., 2002; Dunker, 1981</u>) could be used. However,
 the observational basis for comparing a model's response is largely unavailable for many
 problems of interest, in large part because meteorological conditions are also changing
 while the emissions are changing. As a result, methods such as DDM are used mainly to
 address the effectiveness of emissions controls.

3.3.1 Global Scale CTMs

- 6 With recognition of the global nature of many air pollution problems, global scale CTMs 7 have been applied to regional scale pollution problems (NRC, 2009). Global-scale CTMs 8 are used to address issues associated with global change, to characterize long-range 9 transport of air pollutants, and to provide boundary conditions for the regional-scale 10 models. The upper boundaries of global scale CTMs extend anywhere from the 11 tropopause (~ 8 km at the poles to ~ 16 km in the tropics) to the mesopause at ~ 80 km, in 12 order to obtain more realistic boundary conditions for problems involving stratospheric 13 dynamics and chemistry. The global-scale CTMs consider the same processes shown in 14 Figure 3-5 for the regional scale models. In addition, many of the same issues that have 15 arisen for the regional models have also arisen for the global scale models (Emmerson 16 and Evans, 2009). For example, after adjusting lightning NO_x to better match observed 17 constraints in the MOZART-4 model, simulated HNO₃ was too low and PAN too high in 18 the mid-troposphere, though observations were captured in the upper troposphere, over 19 the U.S. during summer 2004 in the MOZART-4 model (Fang et al., 2010). In contrast, 20 summer 2004 simulations with improved lightning NO_x in GEOS-Chem indicate that 21 PAN is too low but HNO₃ is overestimated throughout the mid- and upper troposphere 22 (Hudman et al., 2007). Predictions of HNO_3 were too high and PAN too low over the 23 U.S. during summer in the MOZART model (Fang et al., 2010). Similar findings were 24 obtained in a box model of upper tropospheric chemistry (Henderson et al., 2011), 25 indicating a need for improved constraints on processes controlling NO_Y distributions in 26 the free troposphere.
- 27 The GEOS-Chem model is a community-owned, global scale CTM that has been widely 28 used to study issues associated with the hemispheric transport of pollution and global 29 change (Harvard University, 2010a). Comparisons of the capabilities of GEOS-Chem and 30 several other models to simulate intra-hemispheric transport of pollutants are given in a 31 number of articles (Fiore et al., 2009; Reidmiller et al., 2009). Reidmiller et al. (2009) 32 compared 18 global models and their ensemble average to spatially and monthly 33 averaged observations of O_3 at CASTNET sites in the U.S. (see Figure 3-6). These results 34 show that the multi-model ensemble agrees much better with observations than do most 35 of the individual models. The GEOS-Chem model was run for two grid spacings

1 $(4^{\circ} \times 4.5^{\circ} \text{ and } 2^{\circ} \times 2.5^{\circ})$ over the U.S. with very similar results that lie close to the2ensemble average. In general, the model ensemble mean and the two GEOS-Chem3simulations are much closer to observations in the Intermountain West than in the4Southeast during summer, when most major O_3 episodes occur (Note, though, that more5current versions of GEOS-Chem are now in use.) However, there are also sizable over-6predictions by many models in both regions during summer.



Source: Reprinted with permission of Copernicus Publications, (Reidmiller et al., 2009).

Figure 3-6 Comparison of global chemical-transport model (CTM) predictions of maximum daily 8-h avg ozone concentrations and multi-model mean with monthly averaged CASTNET observations in the Intermountain West and Southeast regions of the U.S.

7	In their review, McDonald-Buller et al. (2011) noted that global scale chemical transport
8	models exhibit biases in monthly mean daily maximum 8-h avg (MDA8) O_3 in some
9	regions of the U.S., including the Gulf Coast, regions affected by fires, and regions with
10	complex topography, which have implications for model estimates of background O_3 ; and
11	they also have difficulty representing the fine structures of O_3 events at sub-grid scales at

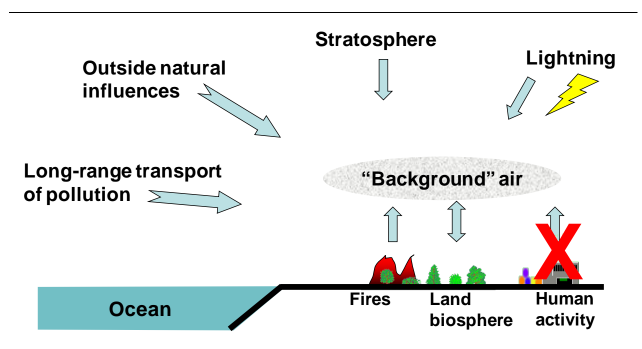
relatively remote monitoring sites that include contributions to O₃ from background sources.

- 3 Global models are not alone in overestimating O₃ in the Southeast. Godowitch et al. 4 (2008), Gilliland et al. (2008) and Nolte et al. (2008) found positive O_3 biases in regional 5 models over the eastern U.S., as well, which they largely attributed to uncertainties in 6 temperature, relative humidity and planetary boundary layer height. Agreement between 7 monthly average values is expected to be better than with daily values because of a 8 number of factors including the increasing uncertainty of emissions at finer time 9 resolution. Kasibhatla and Chameides (2000) found that the accuracy of simulations 10 improved as the averaging time of both the simulation and the observations increased.
- 11Simulations of the effects of long-range transport at particular locations must be able to12link multiple horizontal resolutions from the global to the local scale. Because of
- 13 computational limitations, global simulations are not made at the same horizontal 14 resolutions found in the regional scale models, i.e., down to 1-4 km² horizontal 15 resolution. They are typically conducted with a horizontal grid spacing of 1°-2° of latitude 16 and longitude (or roughly 100-200 km at mid-latitudes). Some models such as GEOS-17 Chem have the capability to include nested models at a resolution of $0.5^{\circ} \times 0.667^{\circ}$ (Wang 18 et al., 2009a) and efforts are underway to achieve even higher spatial resolution. Another 19 approach is to nest regional models within GEOS-Chem. Caution must be exercised with 20 nesting different models because of differences in chemical mechanisms and numerical 21 schemes, and in boundary conditions between the outer and inner models. As an example 22 of these issues, surface O_3 concentrations that are too high have been observed in models 23 in which CMAQ was nested inside of GEOS-Chem. The high O₃ results in large measure 24 from stratospheric O₃ intruding into the CMAQ domain (see (Lam and Fu, 2010) for one 25 way to address this issue). Large vertical O_3 gradients in the upper troposphere must be 26 preserved to accurately represent downward transport of stratospheric O₃. This 27 complicates efforts to link global and regional models with different vertical grid spacing. 28 Efforts are also underway to extend the domain of CMAQ over the entire Northern 29 Hemisphere. In this approach, the same numerical schemes are used for transporting 30 species and the same chemistry is used throughout all spatial scales. Finer resolution in 31 models of any scale can only improve scientific understanding to the extent that the 32 governing processes are accurately described. Consequently, there is a crucial need for 33 observations at the appropriate scales to evaluate the scientific understanding represented
- 34 by the models.

1

3.4 Background Ozone Concentrations

1	Background concentrations of O ₃ have been given various definitions in the literature
2	over time. An understanding of the sources and contributions of background O_3 to
3	O_3 concentrations in the U.S. is potentially useful in reviewing the O_3 NAAQS,
4	especially related to days at the upper end of the distribution of O_3 concentrations. In the
5	context of a review of the NAAQS, it is useful to define background O_3 concentrations in
6	a way that distinguishes between concentrations that result from precursor emissions that
7	are relatively less controllable from those that are relatively more controllable through
8	U.S. policies. In previous NAAQS reviews, a specific definition of background
9	concentrations was used and referred to as policy relevant background (PRB). In those
10	previous reviews, PRB concentrations were defined by EPA as those concentrations that
10	would occur in the U.S. in the absence of anthropogenic emissions in continental North
12	America (CNA), defined here as the U.S., Canada, and Mexico. There is no chemical
12	difference between background O_3 and O_3 attributable to CNA anthropogenic sources.
13	However, to inform policy considerations regarding the current or potential alternative
15	standards, it is useful to understand how total O_3 concentrations can be attributed to
15	different sources.
10	different sources.
17	For this document, EPA has considered background O_3 concentrations more broadly by
18	considering three different definitions of background. The first is natural background
19	which includes contributions resulting from emissions from natural sources
20	(e.g., stratospheric intrusion, wildfires, biogenic methane and more short-lived VOC
21	emissions) throughout the globe simulated in the absence of all anthropogenic emissions.
22	The second is North American background (NA background) which includes
23	contributions from natural background throughout the globe and emissions of
24	anthropogenic pollutants contributing to global concentrations of O3 (e.g., anthropogenic
25	methane) from countries outside North America. The third is United States background
26	(U.S. background) which includes contributions from natural background throughout the
27	globe and emissions from anthropogenic pollutants contributing to global concentrations
28	of O_3 from countries outside the U.S. U.S. background differs from NA background in
29	that it includes anthropogenic emissions from neighboring Canada and Mexico. These
30	three definitions have been explored in recent literature and are discussed further below.
31	Sources included in the definitions of NA background and U.S. background O ₃ are shown
32	schematically in Figure 3-7. Definitions of background and approaches to derive
33	background concentrations were reviewed in the 2006 O_3 AQCD (U.S. EPA, 2006b) and
34	in <u>Reid et al. (2008</u>). Further detail about the processes involved in these sources is given
35	in Section <u>3.4.1</u> and Section <u>3.4.2</u> and application to models calculating background
36	concentrations is presented in Section $3.4.3$.



Note: Background concentrations are ozone concentrations that would exist in the absence of anthropogenic emissions from the U.S., Canada, and Mexico. United States background is similarly defined, but includes transport from Canada and Mexico in addition to intercontinental transport.

Figure 3-7 Schematic overview of contributions to North American background concentrations of ozone.

3.4.1 Contributions from Natural Sources

1	Natural sources contributing to background O ₃ include the stratospheric-tropospheric
2	exchange (STE) of O_3 and photochemical reactions involving natural O_3 precursor
3	emissions of VOCs, NO _X , and CO. Natural sources of O ₃ precursors include biogenic
4	emissions, wildfires, and lightning. Biogenic emissions from agricultural activities in
5	CNA (or the U.S.) are not considered in the formation of NA (or U.S.) background O ₃ .
6	Contributions from natural sources are an important component of background
7	concentrations and are discussed in greater detail below.

3.4.1.1 Contributions from the Stratosphere

8	The basic atmospheric dynamics and thermodynamics of STE were outlined in the 2006
9	O_3 AQCD (U.S. EPA, 2006b); as noted there, stratospheric air rich in O_3 is transported
10	into the troposphere. Ozone is produced naturally by photochemical reactions in the
11	stratosphere as shown in Figure 3-1. Some of this O_3 is transported downward into the
12	troposphere throughout the year, with maximum contributions at mid-latitudes during late

- 1 winter and early spring mainly coming from a process known as tropopause folding. 2 These folds occur behind most cold fronts, bringing stratospheric air with them. The 3 tropopause should not be interpreted as a material surface through which there is no 4 exchange. Rather these folds should be thought of as regions in which mixing of 5 tropospheric and stratospheric air is occurring (Shapiro, 1980). This imported 6 stratospheric air contributes to the natural background of O_3 in the troposphere, especially 7 in the free troposphere during winter and spring. STE also occurs during other seasons 8 including summer.
- 9 Methods for estimating the contribution of stratospheric intrusions rely on the use of 10 tracers of stratospheric origin that can be either dynamical or chemical. Thompson et al. 11 (2007) found that roughly 20-25% of tropospheric O₃ over northeastern North America 12 during July-August 2004 was of stratospheric origin based on an analysis of ozonesonde 13 data. This O_3 can be mixed into the PBL where it can either be destroyed or transported 14 to the surface. They relied on the combined use of low relative humidity and high 15 (isentropic) potential vorticity (PV) (>2 PV units) to identify stratospheric contributions. 16 PV has been a widely used tracer for stratospheric air; see the 2006 O_3 AQCD (U.S. 17 EPA, 2006b). Lefohn et al. (2011) used these and additional criteria to assess 18 stratospheric influence on sites in the Intermountain West and in the Northern Tier. 19 Additional criteria include consideration of trajectories originating at altitudes above the 20 380 K potential temperature surface with a residence time requirement at these heights. 21 Based on these criteria, they identified likely stratospheric influence at the surface sites 22 on a number of days during spring of 2006 to 2008. However, they noted that their 23 analysis of stratospheric intrusions captures only the frequency and vertical penetration of 24 the intrusions but does not provide information about the contribution of the intrusions to 25 the measured O_3 concentration. These results are all generally consistent with what was 26 noted in the 2006 O₃ AQCD (U.S. EPA, 2006b). Fischer (2004) analyzed the O₃ record 27 during summer at Mount Washington and identified a stratospheric contribution to 5% of 28 events during the summers of 1998 -2003 when O_3 was >65 ppb; the air was dry and 29 trajectories originated from altitudes where PV was elevated (PV >1 PV unit). However, 30 this analysis did not quantify the relative contributions of anthropogenic and stratospheric 31 O_3 sources, because as they note identifying stratospheric influences is complicated by 32 transport over industrialized/urban source regions. Stratospheric O₃ was hypothesized to 33 influence the summit during conditions also potentially conducive to photochemical O_3 34 production, which make any relative contribution calculations difficult without additional 35 measurements of anthropogenic and stratospheric tracers.
- Although most research has been conducted on tropopause folding as a source of
 stratosphere to troposphere exchange, this is not the only mechanisms by which
 stratospheric O₃ can be brought to lower altitudes. <u>Tang et al. (2011)</u> estimated that deep

1	convection capable of penetrating the tropopause can increase the overall downward flux
2	of O_3 by ~20%. This mechanism operates mainly during summer in contrast with
3	tropopause folding which is at a maximum from late winter through spring and at lower
4	latitudes. <u>Yang et al. (2010</u>) estimated that roughly 20% of free tropospheric O_3 above
5	coastal California in 2005 and 2006 was stratospheric in origin. Some of this O_3 could
6	also contribute to O_3 at the surface.
7	It should be noted that there is considerable uncertainty in the magnitude and distribution
8	of this potentially important source of tropospheric O ₃ . Stratospheric intrusions that reach
9	the surface are much less frequent than intrusions which penetrate only to the middle and
10	upper troposphere. However, O ₃ transported to the upper and middle troposphere can still
11	affect surface concentrations through various exchange mechanisms that mix air from the
12	free troposphere with air in the PBL.
13	Several instances of STE producing high concentrations of O ₃ around Denver and
14	Boulder, CO were analyzed by Langford et al. (2009) and several likely instances of
15	STE, including one of the cases analyzed by Langford et al. (2009) were also cited in
16	Annex AX2-3 of the 2006 O ₃ AQCD (U.S. EPA, 2006b). Clear examples of STE have
17	also been observed in southern Quebec province by Hocking et al. (2007), in accord with
18	previous estimates by Wernli and Bourqui (2002) and James et al. (2003). As also noted
19	in the 2006 O ₃ AQCD (U.S. EPA, 2006b), the identification of stratospheric O ₃ and the
20	calculation of its contributions to ambient air requires data for other tracers of
21	stratospheric origin. In some cases, stratospheric ozone intrusions can be identified based
22	on measurements of low relative humidity, high potential vorticity and low ratios of
23	O ₃ /PM. Strong stratospheric ozone intrusion events that typically occur during winter or
24	spring have been readily identified using these types of data (Langford et al., 2009).
25	However, it remains challenging to accurately estimate the contributions from smaller
26	direct or indirect (i.e., resulting from shallow intrusions into the mid and upper
27	troposphere that are then mixed downward into the planetary boundary layer)
28	contributions of stratospheric ozone to ambient air.

3.4.1.2 Contributions from Other Natural Sources

Biomass burning consists of wildfires and the intentional burning of vegetation to clear
new land for agriculture and for population resettlement; to control the growth of
unwanted plants on pasture land; to manage forest resources with prescribed burning; to
dispose of agricultural and domestic waste; and as fuel for cooking, heating, and water
sterilization. Biomass burning also exhibits strong seasonality and interannual variability
(van der Werf et al., 2006), with most biomass burned during the local dry season. This is

- 1 true for both prescribed burns and wildfires. Globally, most wildfires may be ignited 2 directly as the result of human activities, leaving only 10-30% initiated by lightning 3 (Andreae, 1991). However, because fire management practices suppress natural wildfires, 4 the buildup of fire fuels increases the susceptibility of forests to more severe but less 5 frequent fires. Thus there is considerable uncertainty in attributing the fraction of wildfire 6 emissions to human activities because the emissions from naturally occurring fires that 7 would have been present in the absence of fire suppression practices are not known. 8 Contributions to NO_x, CO and VOCs from wildfires and prescribed fires are considered 9 as precursors to background O₃ formation in this assessment.
- 10 Estimating contributions from wildfires is subject to considerable uncertainty.
- 11 <u>McDonald-Buller et al. (2011)</u> note that "Models generally find little O_3 production in 12 wildfire plumes for short aging times (days) because NO_X emissions are low and 13 conversion to peroxyacetylnitrate (PAN) is rapid. In contrast, observations show large O_3 14 production from at least some regional wildfires that may significantly elevate O_3 at low 15 altitude sites on a monthly basis, and persist over long distances from the burned region." 16 They also note that fire plumes transported on intercontinental scales can contain very 17 high O_3 concentrations. However Singh et al. (2010b) found appreciable increases of O_3 18 in California fire plumes only when they are mixed with urban pollution. Jaffe and 19 Wigder (2012) note that this result could have also been due to suppression of O_3 20 production near the source. Factors such as the stage of combustion (smoldering to 21 flaming), fuel nitrogen content, ambient meteorological conditions, and the availability of 22 solar ultraviolet radiation need to be considered when evaluating the potential of fires for 23 producing O₃.
- 24 Jaffe et al. (2008) examined the effects of wildfires on O_3 in the western U.S. They found a strong relation ($R^2 = 0.60$) between summer mean O₃ measured at various national park 25 and CASTNET sites and area burned in the western U.S. They also found generally 26 27 higher concentrations within surrounding $5^{\circ} \times 5^{\circ}$ and $10^{\circ} \times 10^{\circ}$ of burned areas. Smaller 28 correlations were found within the surrounding $1^{\circ} \times 1^{\circ}$ areas, reflecting near source 29 consumption of O_3 and the time necessary for photochemical processing of emissions to 30 form O_3 . Jaffe et al. (2008) estimate that burning 1 million acres in the western U.S. 31 during summer results in an increase in O_3 of 2 ppb across the region; this translates to an 32 average O_3 increase across the entire western U.S. of 3.5 and 8.8 ppb during mean and 33 maximum fire years. The unusually warm and dry weather in central Alaska and western 34 Yukon in the summer of 2004 contributed to the burning of 11 million acres there. 35 Subsequent modeling by Pfister et al. (2005) showed that the CO contribution from these 36 fires in July 2004 was 33.1 (\pm 5.5) MT that summer, roughly comparable to total U.S. 37 anthropogenic CO emissions during the same period.

1	These results underscore the importance of wildfines as a source of important O
	These results underscore the importance of wildfires as a source of important O_3
2	precursors. In addition to emissions from forest fires in the U.S., emissions from forest
3	fires in other countries can be transported to the U.S., for example from boreal forest fires
4	in Canada (Mathur, 2008), Siberia (Generoso et al., 2007) and tropical forest fires in the
5	Yucatan Peninsula and Central America (Wang et al., 2006). These fires have all resulted
6	in notable increases in O_3 concentrations in the U.S.
7	Estimates of biogenic VOC, NO and CO emissions can be made using the BEIS model
8	with data from the BELD and annual meteorological data or MEGAN. VOC emissions
9	from vegetation were described in Section 3.2 .
10	As discussed in Section 3.2.1, NO _X is produced by lightning. Kaynak et al. (2008) found
11	lightning contributes 2 to 3 ppb to surface-level background O3 centered mainly over the
12	southeastern U.S. during summer. Although total column estimates of lightning produced
13	NO_X are large compared to anthropogenic NO_X during summer, lightning produced NO_X
14	does not contribute substantially to the NO_X burden in the continental boundary layer.
15	For example, (Fang et al., 2010) estimated that only 2% of NO _X production by lightning
16	occurs within the boundary layer and most occurs in the free troposphere. In addition,
17	much of the NO_X produced in the free troposphere is converted to more oxidized
18	N species during downward transport. Note that contributions of natural sources to North
19	American background arise from everywhere in the world.

3.4.2 Contributions from Anthropogenic Emissions

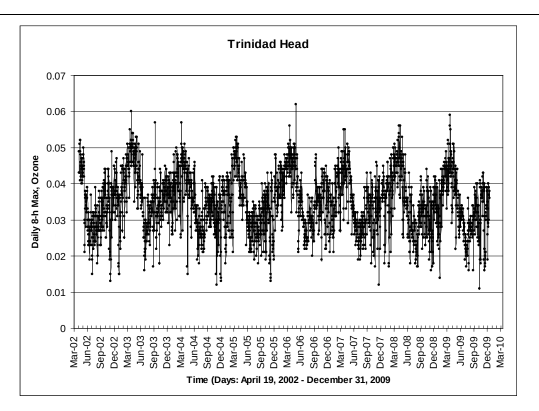
20	In addition to emissions from North America, anthropogenic emissions from Eurasia
21	have contributed to the global burden of O_3 in the atmosphere and to the U.S. (<u>NRC</u> ,
22	<u>2009, and references therein</u>). Because the mean tropospheric lifetime of O_3 is on the
23	order of a few weeks (<u>Hsu and Prather, 2009</u>), O_3 can be transported from continent to
24	continent and around the globe in the Northern Hemisphere. Ozone produced by U.S.
25	emissions can, therefore, be recirculated around northern mid-latitudes back to the U.S.
26	High elevation sites are most susceptible to the intercontinental transport of pollution
27	especially during spring. For example, a number of occurrences of O ₃ >60 ppb from mid-
28	April to mid-May of 2006 were observed at Mt. Bachelor Observatory, OR (elevation
29	2,700 m) with a maximum O_3 concentration of ~85 ppb observed on April 22, 2006.
30	Calculations using GEOS-Chem, a global-scale CTM, indicate that Asia contributed
31	9 ± 3 ppb to a modeled mean concentration of 53 ± 9 ppb O_3 at Mt. Bachelor during the
32	same period compared to measured concentrations of 54 ± 10 ppb (Zhang et al., 2008).
33	<u>Zhang et al. (2008</u>) also calculated a contribution of 5 to 7 ppb to surface O_3 over the
34	western U.S. during that period from Asian anthropogenic emissions. They also estimated

1	an increase in NO_X emissions of ~44% from Asia from 2001 to 2006 resulting in an
2	increase of 1-2 ppb in O ₃ over North America.

3 Cooper et al. (2010) analyzed all available O_3 measurements in the free troposphere 4 above western North America at altitudes of 3-8 km (above sea level) during April and 5 May of 1995 to 2008 (i.e., times when intercontinental transport is most prominent). 6 They derived a trend of $+0.63 \pm 0.34$ ppb/year in median O₃ concentrations with 7 indication of a similar rate of increase since 1984. Back trajectories that were likely to 8 have been strongly and recently influenced by North American emissions were filtered 9 out, resulting in a trend of $+0.71 \pm 0.45$ ppb/year. Considering only trajectories with an 10 Asian origin resulted in a trend of $+0.80 \pm 0.34$ ppb/year. These results suggest that local 11 North American emissions were not responsible for the measured O_3 increases. This O_3 12 could have been produced from natural and anthropogenic precursors in Asia and Europe 13 with some contribution from North American emissions that have circled the globe. 14 Cooper et al. (2010) also found that it is unlikely that the trends in tropospheric O_3 are 15 associated with trends in stratospheric intrusions. Note, however, that these results relate 16 to O_3 trends above ground level and not to surface O_3 . Model results (Zhang et al., 2008) 17 show that surface O₃ contributions from Asia are much smaller than those derived in the 18 free troposphere because of dilution and chemical destruction during downward transport 19 to the surface. These processes tend to reduce the strength of associations between free 20 tropospheric and surface O_3 especially if air from other sources is sampled by the surface 21 monitoring sites.

22 Trinidad Head, CA is one sampling location at which measurements might be expected to 23 reflect in large measure NA background O_3 contributions, at times during the spring 24 (Oltmans et al., 2008; Goldstein et al., 2004). The monitoring station at Trinidad Head is 25 on an elevated peninsula extending out from the mainland of northern California, and so 26 might be expected at times to intercept air flowing in from the Pacific Ocean with little or 27 no influence from sources on the mainland. Figure 3-8 shows the time series of MDA8 28 O₃ concentrations measured at Trinidad Head from April 18, 2002 through December 31, 29 2009. The data show pronounced seasonal variability with spring maxima and summer 30 minima. Springtime concentrations typically range from 40 to 50 ppb with a number of 31 occurrences >50 ppb. The two highest daily maxima were 60 and 62 ppb. The data also 32 show much lower concentrations during summer, with concentrations typically ranging 33 between 20 and 30 ppb. Oltmans et al. (2008) examined the time series of O_3 and back 34 trajectories reaching Trinidad Head. They found that springtime maxima (April-May) 35 were largely associated with back trajectories passing over the Pacific Ocean and most 36 likely entraining emissions from Asia, with minimal interference from local sources. 37 However, Parrish et al. (2009) noted that only considering trajectories coming from a given direction is not sufficient for ruling out local continental influences, as sea breeze 38

1	circulations are complex phenomena involving vertical mixing and entrainment of long-
2	shore components. They found that using a wind speed threshold in addition to a criterion
3	for wind direction allowed determination of background trajectories not subject to local
4	influence. This was confirmed by measurements of chemical tracers of local influence
5	such as CO ₂ , MTBE and radon. By applying the two criteria for wind speed and
6	direction, they found that Trinidad Head met these criteria only 30% of the time during
7	spring. Goldstein et al. (2004) used CO_2 as an indicator of exchange with the local
8	continental environment and found that O3 concentrations were higher by about 2-3 ppb
9	when filtered against local influence indicating higher O_3 in air arriving from over the
10	Pacific Ocean. At other times of the year, Trinidad Head is less strongly affected by air
11	passing over Asia and the northern Pacific Ocean; and many trajectories have long
12	residence times over the semi-tropical and tropical Pacific Ocean where O_3
13	concentrations are much lower than they are at mid-latitudes. The use of the Trinidad
14	Head data to derive contributions from background sources requires the use of screening
15	procedures adopted by Parrish et al. (2009) and the application of photochemical models
16	to determine the extent either of titration of O_3 by fresh NO_X emissions and the extent of
17	local production of O_3 from these emissions. As noted above, anthropogenic emissions
18	from North America also contribute to hemispheric background and must be filtered out
19	from observations even when it is thought that air sampled came directly from over the
20	Pacific Ocean and was not influenced by local pollutant emissions.



Source: Reprinted with permission of Elsevier Ltd., (Oltmans et al., 2008); and NOAA Climate Monitoring Diagnostics Laboratory for data from 2008-2009.

Figure 3-8 Time series of daily maximum 8-h avg (MDA8) ozone concentrations (ppm) measured at Trinidad Head, CA, from April 18, 2002 through December 31, 2009.

1	Parrish et al. (2009) also examined data obtained at other marine boundary layer sites on
2	the Pacific Coast. These include Olympic NP, Redwood NP, Point Arena, and Point
3	Reyes. Using data from these sites, they derived trends in O_3 of +0.46 ppb/year (with a
4	95% confidence interval of 0.13 ppb/year) during spring and +0.34 ppb/year
5	(0.09 ppb/year) for the annual mean O_3 increase in air arriving from over the Pacific
6	during the past two decades. Although O_3 data are available from the Channel Islands,
7	Parrish et al. (2009) noted that these data are not suitable for determining background
8	influence because of the likelihood of circulating polluted air from the South Coast Basin.
9	The 2010 Intercontinental Chemical Transport Experiment Ozone Network Study (IONS-
10	2010) and Research at the Nexus of Air Quality and Climate Change (CalNex) study
11	conducted in May through June of 2010 had discerning the contributions of Asian
12	emissions to air quality in California as a major focus. Cooper et al. (2011) examined O_3
13	profiles measured above four coastal sites in California, including Trinidad Head. Based
14	on trajectory analyses coupled with comparison with the O ₃ profiles, they suggested that

- 1 Asian pollution, stratospheric intrusions and international shipping made substantial 2 contributions to lower tropospheric O_3 (typically 0 to 3 km above sea level, meant as a 3 rough approximation of planetary boundary layer height) measured at inland California 4 sites. These contributions tended to increase on a relative basis in going from south to 5 north. In particular, no contribution from local pollution was needed to explain lower 6 tropospheric O_3 in the northern Central Valley; and the contribution of local pollution to 7 lower tropospheric O_3 in the LA basin ranged from 32 to 63% (depending on layer depth; 8 either 0 to 1.5 km or 0 to 3 km). It should be noted that the extent of photochemical 9 production and loss occurring in the descending air masses between the coastal and 10 inland sites remains to be determined. Cooper et al. (2011) also note that very little of the 11 O_3 observed above California reaches the eastern U.S. However, this does not necessarily 12 mean that the pathways by which Asian O_3 could reach the eastern U.S. were fully 13 captured in this analysis.
- 14 Lin et al. (2012) used the AM3 model (\sim 50 × 50 km resolution globally) and satellite data 15 to characterize the influence of Asian emissions and stratospheric intrusions on O₃ 16 concentrations in southern California and Arizona during CalNex (May-June 2010). The 17 model simulates sharp O_3 gradients in the upper troposphere and the interweaving and 18 mixing of stratospheric air and Asian plumes. Similar phenomena were also found during 19 field campaigns conducted in the North Atlantic as noted in Annex AX2.3.1 of the 2006 20 O_3 AQCD (U.S. EPA, 2006b) and introduces uncertainty into attempts to attribute O_3 to 21 these sources, based solely on observations, because this mixing will affect relationships 22 between CO (mainly a marker for polluted air that is commonly used to separate air 23 influenced by anthropogenic pollution from stratospheric air) and O_3 (a pollutant and a 24 stratospheric component). Lin et al. (2012) found that Asian emissions contributed from 25 20-30% to O_3 in the mid troposphere over the California coast and remnants of 26 stratospheric intrusions contributed from 50 to 60% to O_3 in discrete layers in the same 27 altitude range. This O_3 then has the potential to mix downward into the planetary 28 boundary layer. Lin et al. (2012) also found evidence of Asian contributions of up to 8 -29 15 ppb in surface air during strong transport events in southern California. These 30 contributions are in addition to contributions from dominant local pollution sources. 31 Their results suggest that the influence of background sources on high surface O_3 32 concentrations is not always confined to high elevation sites. However, it is not clear to 33 what extent the contributions inferred by Cooper et al. (2011) and Lin et al. (2012) are 34 likely to be found in other years or how long they would extend into summer.

3.4.3 Estimating Background Concentrations

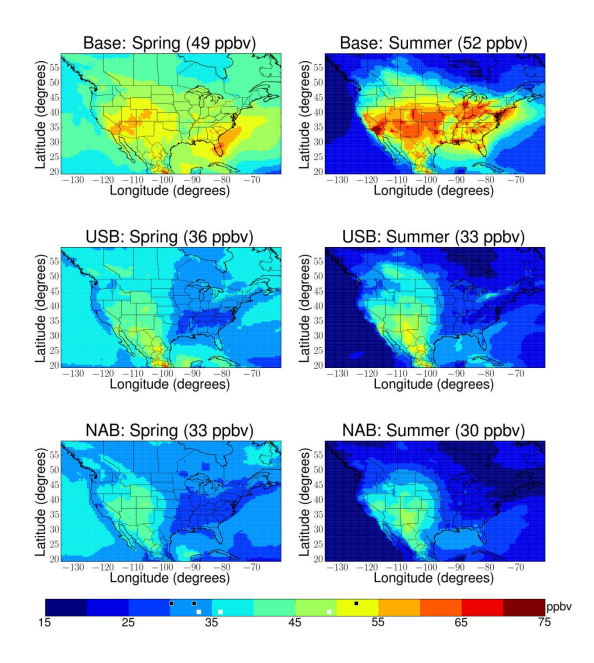
1	Historically, two approaches to estimating NA background concentrations (previously
2	referred to as PRB) have been considered in past O ₃ assessments. In the 1996 and earlier
3	O_3 AQCDs, measurements from remote monitoring sites were used. In the 2006 O_3
4	AQCD, the use of CTMs was adopted, because as noted in Section 3.9 of the 2006 O_3
5	AQCD (U.S. EPA, 2006b), estimates of background concentrations cannot be obtained
6	directly by examining measurements of O ₃ obtained at relatively remote monitoring sites
7	in the U.S. because of the long-range transport from anthropogenic source regions within
8	North America. The 2006 O ₃ AQCD (<u>U.S. EPA, 2006b</u>) also noted that it is impossible to
9	determine sources of O ₃ without ancillary data that could be used as tracers of sources or
10	to calculate photochemical production and loss rates. As further noted by Reid et al.
11	(2008), the use of monitoring data for estimating background concentrations is essentially
12	limited to the edges of the domain of interest. The current definition of NA background
13	implies that only CTMs (see Section 3.3 for description and associated uncertainties) can
14	be used to estimate the range of background concentrations. An advantage to using
15	models is that the entire range of O_3 concentrations measured in different environments
16	can be used to evaluate model performance. In this regard, data from the relatively small
17	number of monitoring sites at which large background contributions are expected are best
18	used to evaluate model predictions.
19	Estimates of NA background concentrations in the 2006 O ₃ AQCD (U.S. EPA, 2006b)
20	were based on output from the GEOS-Chem (v4.3.3) model (Fiore et al., 2003) with
21	$2^{\circ} \times 2.5^{\circ}$ horizontal resolution. The GEOS-Chem model estimates indicated that NA
22	background O_3 concentrations in eastern U.S. surface air were 25 ± 10 ppb (or generally
23	15-35 ppb) from June through August, based on conditions for 2001. Values reported by
24	Fiore et al. (2003) represent averages from 1 p.m. to 5 p.m.; all subsequent values given
25	for background concentrations refer to MDA8 O ₃ concentrations. Background
26	concentrations decline from spring to summer. Background O_3 concentrations may be
27	higher, especially at high altitude sites during the spring, due to enhanced contributions
28	from (1) pollution sources outside North America; and (2) stratospheric O_3 exchange. At
29	the time, only the GEOS-Chem model (<u>Harvard University, 2010b</u>) was documented in
30	the literature for calculating background O_3 concentrations (Fiore et al., 2003). The
31	simulated monthly mean concentrations in different quadrants of the U.S. were typically
32	within 5 ppbv of observations at CASTNET sites, with no descernible bias, except in the
33	Southeast in summer when the model was 8-12 ppbv too high. This bias was attributed to
34	excessive background O_3 transported in from the Gulf of Mexico and the tropical Atlantic
35	Ocean in the model (Fiore et al., 2003).
55	Seean in the model (<u>11010 et al., 2003</u>).

1	Although many of the features of the day-to-day variability in O_3 at relatively remote
2	monitoring sites in the U.S. were simulated reasonably well by GEOS-Chem (Fiore et al.,
3	2003), uncertainties in the calculation of the temporal variability of O ₃ originating from
4	different sources on shorter time scales must be recognized. The uncertainties stem in
5	part from an underestimate in the seasonal variability in the STE of O ₃ (Fusco and Logan,
6	2003), the geographical variability of this exchange, and the variability in the exchange
7	between the free troposphere and the PBL in the model. In addition, the relatively coarse
8	spatial resolution in that version of GEOS-Chem ($2^{\circ} \times 2.5^{\circ}$) limited the ability to provide
9	separate estimates for cities located close to each other, and so only regional estimates
10	were provided for the 2006 O ₃ AQCD (U.S. EPA, 2006b) based on the results of Fiore et
11	<u>al. (2003</u>).
12	Wang et al. (2009a) recomputed NA background concentrations for 2001 using GEOS-
13	Chem (v7-01-01) at higher spatial resolution $(1^{\circ} \times 1^{\circ})$ over North America and not only
14	for afternoon hours but for the daily maximum 8-h O ₃ concentration. The resulting
15	background concentrations, 26.3 ± 8.3 ppb for summer, are consistent with those of
16	26 ± 7 ppb for summer reported by Fiore et al. (2003), suggesting horizontal resolution
17	was not a substantial factor limiting the accuracy of the earlier results. In addition to
18	computing NA background concentrations, Wang et al. (2009a) also computed U.S.
19	background concentrations of 29.6 ± 8.3 ppb with higher concentrations in the Northeast

(up to 15 ppb higher) and the Southwest (up to 13 ppb higher) for summer means.

3.4.3.1 Updated GEOS-Chem Model Estimates of Background Concentrations

21	Zhang et al. (2011) computed NA background, U.S. background and natural background
22	O ₃ concentrations using GEOS-Chem (v8-02-03) at an even finer grid spacing of
23	$0.5^{\circ} \times 0.667^{\circ}$ over North America for 2006 through 2008. For March through August
24	2006, mean NA background O_3 concentrations of 29 ± 8 ppb at low elevation (<1,500 m)
25	and 40 \pm 8 ppb at high elevation (>1,500 m) were predicted. Spring and summer mean O ₃
26	concentrations calculated for the base case (i.e., including all natural and anthropogenic
27	sources worldwide), U.S. background, and NA background in surface air for spring and
28	summer 2006 calculated by Zhang et al. (2011) are shown in the upper, middle and lower
29	panels of <u>Figure 3-9</u> .



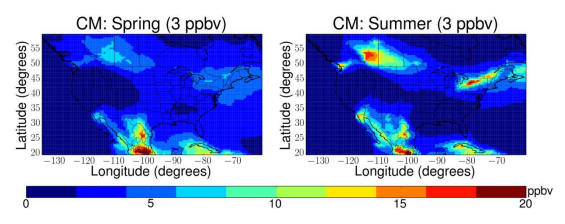
Note: Values in parentheses refer to continental U.S. means and are shown as black squares in the color bar for summer and white squares for spring.

Source: Adapted from Zhang et al. (2011).

Figure 3-9 Mean MDA8 ozone concentrations in surface air for spring and summer 2006 calculated by GEOS-Chem for the base case (Base), U.S. background (USB), and NA background (NAB).

1	As noted above Zhang et al. (2011) found increases in Asian emissions only accounted
2	for an average increase of between 1 to 2 ppb in background O ₃ across the U.S. even
3	though Asian emissions have increased by about 44% from 2001 to 2006. As can be seen
4	from Figure 3-9, U.S. background and NA background concentrations are very similar
5	throughout most of the U.S. Zhang et al. (2011) also found that NA background
6	concentrations are ~4 ppb higher, on average, in the $0.5^{\circ} \times 0.667^{\circ}$ version than in the
7	coarser $2^{\circ} \times 2.5^{\circ}$ version. This difference was not entirely due to higher resolution, but to
8	the combination of changes in lightning and Asian emission estimates as well as higher
9	model resolution.
10	As can be seen from the middle and lower panels in Figure 3-9, U.S. background and NA
11	background concentrations tend to be higher in the West, particularly in the
12	Intermountain West and in the Southwest compared to the East in both spring and
13	summer. U.S. background and NA background concentrations tend to be highest in the
14	Southwest during summer in the GEOS-Chem model, driven by lightning NO_X .
15	Intercontinental transport and stratospheric intrusions are major contributors to the high
16	elevation, Intermountain West during spring with wildfires becoming more important
17	sources during summer. The base case O_3 concentrations (upper panels) show two broad
18	maxima with highest concentrations extending throughout the Southwest, Intermountain
19	West and the East in both spring and summer. These maxima extend over many
20	thousands of kilometers demonstrating that O_3 is a regional pollutant. Low-level outflow
21	from the Northeast out over the Atlantic Ocean and from the Southeast out over the Gulf
22	of Mexico is also apparent.
23	Lower bounds to NA background concentrations tend to be higher by several parts per
24	billion at high elevations than at low elevations, reflecting the increasing importance of
25	background sources such as stratospheric intrusions and intercontinental transport with
26	altitude. In addition, background concentrations tend to increase with increasing base
27	model (and measured) concentrations at higher elevation sites, particularly during spring.
28	Although results of Zhang et al. (2011) are broadly consistent with results from earlier
29	coarser resolution versions of GEOS-Chem used by Fiore et al. (2003) and Wang et al.
30	(2009a), there are some apparent differences. Concentrations of O_3 for both the base case
31	and the NA background case in Zhang et al. (2011) are higher in the Intermountain West
32	than in earlier versions. In addition, background concentrations in many eastern areas
33	tend to be higher on days when predicted total O_3 is >60 ppb or at least do not decrease
34	with increasing total $O_3 $ <u>Zhang et al. (2011</u>).
35	Figure 3-10 shows seasonal mean estimates of contributions to O_3 from Canadian and
36	Mexican emissions calculated by Zhang et al. (2011) as the difference between U.S.

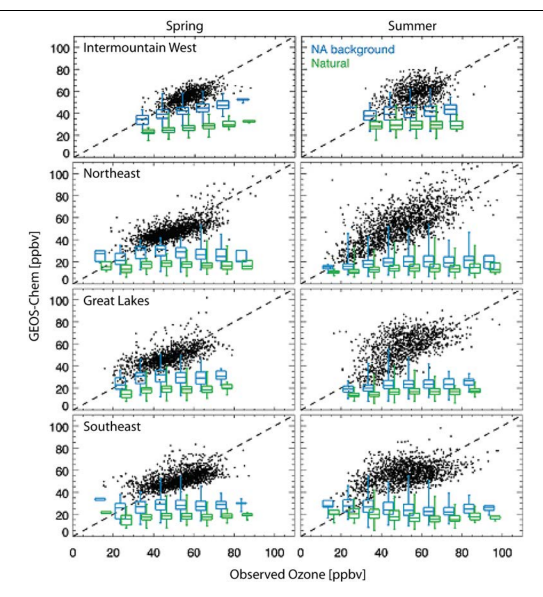
1	background and NA background values and then averaged over spring and summer
2	following the procedure in Wang et al. (2009a). U.S. background concentrations are on
3	average 3 ppb higher than NA background concentrations during spring and summer
4	across the United States. Highest values in Figure 3-10 (in the U.S.) are found over the
5	Northern Tier of New York State (19.1 ppb higher than NA background) in summer.
6	High values are also found in other areas bordering Canada and Mexico. Although the
7	contributions from Canada and Mexico were obtained by differencing, it should be
8	remembered that relations between O3 and precursors are subject to non-linear effects
9	that are strongest near concentrated sources of precursors, as noted in Section $3.2.4$.
10	Therefore, the values shown in the figure are only estimates of contributions to total O_3
11	coming from Canada and Mexico.



Note: Values in parentheses show mean difference (ppb) across the U.S. Source: Adapted from <u>Zhang et al. (2011</u>).

Figure 3-10 Spring and summer mean Canadian and Mexican (CM) contributions to MDA8 ozone determined as the difference between the U.S. background and NA background.

12	Figure 3-11 shows MDA8 O_3 concentrations for spring (March-May) and summer (June-
13	August) 2006 simulated by GEOS-Chem vs. measured by the ensemble of CASTNET
14	sites in the Intermountain West, Northeast, Great Lakes, and Southeast (Zhang et al.,
15	2011). Shown is the 1:1 line and NA background (blue) and natural background (green)
16	model statistics as box plots (minimum, 25th, 50th, 75th percentile, and maximum) for
17	10-ppbv bins of observed ozone concentrations. These plots show that NA background
18	constitute a larger fraction of modeled base case O_3 at the upper end of the concentration
19	distribution for the Intermountain West than for other regions of the country.



Note: Shown is the 1:1 line and North American (NA) background (blue) and natural background (green) model statistics as box plots (minimum, 25th, 50th, 75th percentile, and maximum) for 10-ppbv bins of observed ozone concentrations. Source: Adapted from <u>Zhang et al. (2011</u>).

Figure 3-11 MDA8 ozone concentrations for spring (March-May) and summer (June-August) 2006 simulated by GEOS-Chem vs. measured by the ensemble of CASTNET sites in the Intermountain West, Northeast, Great Lakes, and Southeast.

1	Comparisons between GEOS-Chem and measurements of the mean MDA8 O ₃ between
2	March and August at individual CASTNET sites across the country are shown as
3	supplemental material in Section <u>3.8</u> , <u>Figure 3-58</u> through <u>Figure 3-64</u> . In general, the
4	GEOS-Chem predictions tend to show better agreement with observations at the high-

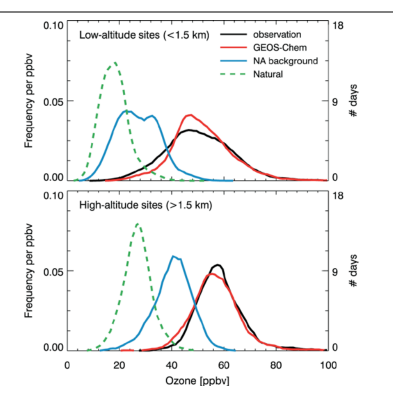
1	altitude sites than at the low-altitude sites. Overall agreement between model results for
2	the base case and measurements is within a few parts per billion for spring-summer
3	means in the Northeast (see Figure 3-58 in Section 3.8) and the Southeast (see
4	<u>Figure 3-59</u> in Section <u>3.8</u>), except in and around Florida where the base case
5	overpredicts O_3 by 10 ppb on average. In the Upper Midwest (Figure 3-60 in
6	Section <u>3.8</u>), the Intermountain West (Figure 3-61 and Figure 3-62 in Section <u>3.8</u>), and
7	the West (Figure 3-63 in Section 3.8) including most sites in California (Figure 3-64 in
8	Section 3.8), the model predictions are within 5 ppb of measurements. The model
9	underpredicts O_3 by 10 ppb at the Yosemite site (Figure 3-64 in Section 3.8). These
10	results suggest that the model is capable of calculating March to August mean MDA8 O_3
11	to within ~5 ppb at most (26 out of 28) sites chosen.

Comparison between results in Wang et al. (2009a) for 2001 with data obtained in the 12 13 Virgin Islands indicate that GEOS-Chem over-predicts summer mean MDA8 O₃ 14 concentrations there by 10 ppb (28 vs. 18 ppb). Ozone concentrations at the Virgin 15 Islands NP site appear not to have been affected by U.S. emissions, based on the close 16 agreement between the base case and the NA background case. Wind roses calculated for 17 the Virgin Islands site indicate that wind patterns affecting this site are predominantly 18 easterly/southeasterly in spring and summer. The over-predictions at the Virgin Islands 19 site imply that modeled O₃ over the tropical Atlantic Ocean is too high. As a result, 20 inflow of O₃ over Florida and into the Gulf of Mexico is also likely to be too high as 21 winds are predominantly easterly at these low latitudes. Similar considerations apply to 22 the results of Zhang et al. (2011). Possible explanations include deficits in model 23 chemistry (for example, reactions involving halogens are not included) and/or subsidence 24 that is too strong over tropical oceans in the model. No clear explanation can be provided 25 on why the model under-predicts mean O_3 at Yosemite (elevation 1,680 m) by ~10 ppb 26 (see Figure 3-64 in Section 3.8). However, March to August mean MDA8 O₃ 27 concentrations are simulated to within a few parts per billion at an even higher elevation 28 site in California (Converse Station, elevation 1,837 m) and at the low elevation sites.

29 Figure 3-65 in Section 3.8 shows a comparison of GEOS-Chem output with 30 measurements at Mt. Bachelor, OR and Trinidad Head, CA from March-August, 2006 31 from Zhang et al. (2011). For the Mt. Bachelor model runs, model estimates are given for 32 both a coarse $(2^{\circ} \times 2.5^{\circ})$ and fine $(0.5^{\circ} \times 0.667^{\circ})$ resolution model. In general, mean 33 concentrations are simulated reasonably well at both coarse and finer grid resolution 34 versions of the model with mean values 2 ppb higher in the finer resolution model. 35 Although the finer resolution version provides some additional day to day variability and 36 can capture the timing of peaks, it still does not adequately resolve peak concentrations as 37 can be seen for an event in the second half of April.

1	
1	Figure 3-66 in Section 3.8 shows a comparison of vertical profiles (mean $\pm 1\sigma$) calculated
2	by GEOS-Chem with ozonesondes launched at Trinidad Head, CA and Boulder, CO. As
3	can be seen from the figure, variability in both model and measurements increases with
4	altitude, but variability in the model results is much smaller at high altitudes than seen in
5	the observations. This may be due in part to the inability of grid-point models to capture
6	the fine-scale, layered structure often seen in O_3 in the mid and upper troposphere
7	(Rastigejev et al., 2010; Newell et al., 1999) and to inadequacies in parameterizations of
8	relevant chemistry and dynamics. Figure 3-67 and Figure 3-68 in Section 3.8 show a
9	comparison of vertical profiles simulated by AM3 at 50×50 km global resolution (Lin et
10	al., 2012) with ozonesondes launched at several locations in California during May-June
11	2010. Note that in contrast to comparing measured mean monthly O_3 profiles to monthly
12	mean profiles calculated by GEOS-CHEM (see, for example, <u>Figure 3-66</u> in Section <u>3.8</u>),
13	AM3 is sampled for comparison to individual measurements of O ₃ profiles. This model
14	has likely had the most success in simulating vertical O_3 gradients in the upper
15	troposphere and in capturing layered structures in the mid and upper troposphere.
16	The natural background for O_3 averages 18 ± 6 ppbv at the low-elevation sites and
17	27 ± 6 ppbv at the high-elevation sites in the GEOS-Chem model <u>Zhang et al. (2011</u>). In
18	regions where non-linear effects are small, far from concentrated sources of O_3
19	precursors, the difference between NA background and natural background O_3
20	concentrations provides an estimate of contributions from intercontinental pollution
21	including anthropogenic methane (given by the difference between values in 2006 and
22	the pre-industrial era, or 1,760 ppb and 700 ppb). The difference between the two
23	backgrounds averages 9 ppbv at the low-elevation sites and 13 ppbv at sites in the
24	Intermountain West. Based on the Zhang et al. (2011) model runs, anthropogenic
25	methane emissions are estimated to contribute \sim 4-5 ppb to global annual mean O ₃ surface
26	concentrations. North American emissions of methane are uncertain, but are a small
27	fraction of total anthropogenic input. This suggests that slightly less than half of the
28	difference between North American background and natural background is due to the
29	increase of methane since the beginning of the industrial era and the other half is due to
30	anthropogenic emissions of shorter lived VOCs and NO_x . However, the relative
31	importance of methane for O_3 production is expected to increase in the future. Indeed,
32	variations in methane concentrations account for approximately 75% of the wide spread
33	(~5 ppb) in tropospheric O_3 projections between Representative Concentration Pathway
34	(RCP) scenarios for the next century (<u>Wild et al., 2012</u>); see Section <u>10.3.6.1</u> in
35	Chapter $\underline{10}$ for more on the RCP scenarios.
36	Figure 3-12 shows frequency distributions for observations at low-altitude and high-
37	altitude CASTNET sites along with GEOS-Chem frequency distributions for the base
38	case, NA background and natural background. Most notable is the shift to higher

concentrations and the narrowing of the concentration distributions for all three
 simulations and the observations in going from low to high altitudes. However, maximum
 concentrations show little if any dependence on altitude, except for the natural
 background which tends to be slightly higher at high altitude sites.



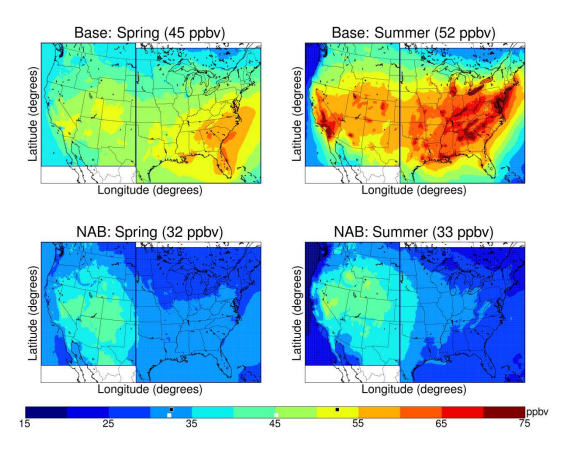
Note: Observations (black) as well as GEOS-Chem estimates for the base case (red), NA background (blue), and natural background (green dashed). Source: Zhang et al. (2011).

Figure 3-12 Frequency distributions of MDA8 ozone concentrations in March-August 2006 for the ensemble of low-altitude (<1,500 meters) and high-altitude CASTNET sites (>1,500 meters) in the U.S.

3.4.3.2 Using Other Models to Estimate Background Concentrations

5	Another approach to modeling background concentrations involves using a regional CTM
6	such as CMAQ or CAMx with boundary conditions taken from a global scale CTM such
7	as GEOS-Chem (see Section 3.3 for discussion of this approach). Mueller and Mallard
8	(2011a), while not calculating NA background values exactly as defined here, calculated
9	contributions from natural sources and inflow from the boundaries to O_3 for 2002 using

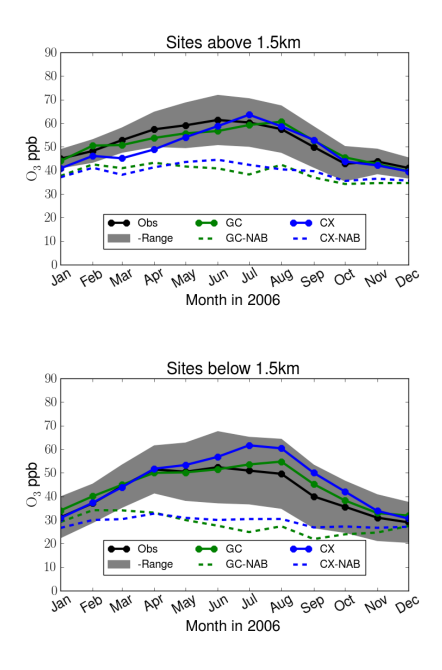
1	MM5 and CMAQ for the outermost domain (36 km resolution) shown in Figure 3-4 with
2	boundary conditions from GEOS-Chem. The overall bias based on comparison with AQS
3	monitors for the base case is about 3 ppb; the annual mean fractional bias and mean
4	fractional error were 7% and 21% for the O_3 season across the U.S. Note that Figure 2 in
5	their paper is mislabeled, as it should refer to the case with total emissions - not to natural
6	emissions in North America only (<u>Mueller and Mallard, 2011b</u>). However, boundary
7	conditions are fixed according to monthly averages based on an earlier version of GEOS-
8	Chem and do not reflect shorter term variability or trends in Northern Hemispheric
9	emissions of pollution. In addition, fluxes of O_3 from the stratosphere are not included
10	explicitly. Note that their natural background includes North American natural
11	background emissions only and influence from boundary conditions and thus is not a
12	global natural background. Calculated values including natural emissions from North
13	America and from fluxes through the boundaries are somewhat larger than given in
14	Zhang et al. (2011), in large measure because of much larger contributions from wildfires
15	and lightning. Wildfire contributions reach values of ~140 ppb in Redwoods National
16	Park, CA and higher elsewhere in the U.S. and in Quebec in the simulations by <u>Mueller</u>
17	and Mallard (2011a). Lightning contributions (ranging up to ~30 ppb) are substantially
18	larger than estimated by <u>Kaynak et al. (2008</u>) (see Section <u>$3.4.1.2$</u>). The reasons for much
19	larger contributions from wildfires and lightning found by <u>Mueller and Mallard (2011a</u>)
20	are not clear and need to be investigated further.
21	Emery et al. (2012) used CAMx in conjunction with boundary conditions from a coarse
22	resolution version of GEOS-Chem ($2^{\circ} \times 2.5^{\circ}$ or ~200 km resolution) to derive NA
23	background concentrations of O ₃ . The nested CAMx simulations were run at a horizontal
24	resolution of 12 km separately for the eastern and western U.S. The following paragraphs
25	compare results from the Emery et al. (2012) nested GEOS-Chem/CAMx simulations
26	(hereafter referred to as CAMx) at 12 km resolution with those obtained by Zhang et al.
27	(2011) using GEOS-Chem simulations at $0.5^{\circ} \times 0.667^{\circ}$ (~50 km) resolution. This is in
28	contrast to the comparison reported in Emery et al. (2012) using a $2^{\circ} \times 2.5^{\circ}$ (~200 km)
29	resolution GEOS-Chem model. Figure 3-13 shows seasonal mean MDA8 O ₃
30	concentrations calculated by Emery et al. (2012) using CAMx for 2006 for the base case
31	and for NA background. Figure 3-14 shows a comparison of monthly average O_3
32	concentrations calculated by GEOS-Chem (Zhang et al., 2011) with those calculated by
33	CAMx (Emery et al., 2012). Comparison of the base case for GEOS-Chem with that for
34	CAMx in Figure 3-14 indicates broad agreement in spatial patterns.



Note: Values in parentheses refer to continental U.S. means and are shown as black squares in the color bar for summer and white squares for spring.

Source: Adapted from Emery et al. (2012).

Figure 3-13 Mean MDA8 ozone concentrations in surface air during spring and summer 2006 (top) calculated by GEOS-Chem/CAMx for the base case (Base, top) and NA background (NAB, bottom).



Note: Shaded area shows 1 SD range about the mean of observations. Source: Adapted [with permission of Elsevier, <u>Emery et al. (2012)</u>] and <u>Zhang et al. (2011)</u>.

Figure 3-14 Monthly average MDA8 ozone concentrations observed (Obs) and predicted for the base case and NA background (NAB) by GEOS-Chem (GC) and GEOS-Chem/CAMx (CX) at CASTNET sites above 1,500 meters elevation (upper panel) and CASTNET sites below 1,500 meters elevation (lower panel).

1	Second second of the second the second
1 2	Supplemental figures (Figure 3-69 through Figure 3-74) in Section 3.8 show box plots comparing MDA8 O ₃ concentrations calculated by GEOS-Chem at $0.5^{\circ} \times 0.667^{\circ}$
2 3	
3 4	resolution and CAMx for March-August 2006 at the combined set of CASTNET sites
4 5	used by both groups for model evaluation. Note that the individual model results and the
5 6	observations are un-paired in time. At CASTNET sites in the Northern Rockies, both models tend to underpredict maximum O_3 concentrations, but they are generally higher in
0 7	
8	CAMx than in GEOS-Chem (typically by 5-10 ppb). The distribution of MDA8 values
8 9	from GEOS-Chem is consistent with measured distributions (i.e., cannot be rejected using Mann Whitney contrast, n yolug (0.01) at 18 of 20 sites in apring and 21 of
9 10	using Mann-Whitney rank sum test, p-value <0.01) at 18 of 39 sites in spring and 21 of 20 sites in symmer. The distribution of MDA8 values from CAMy is consistent with
	39 sites in summer. The distribution of MDA8 values from CAMx is consistent with
11	measured distributions at 13 of 39 sites in spring and 18 of 39 sites in summer. When
12	spring and summer are pooled, both simulations are consistent with measured
13	distributions at 16 out of 39 sites (but not the same 16 sites). There are examples in which
14	either model over- or under- simulates maximum concentrations. However, over-
15	predictions are made more often by CAMx. At high elevations in the Intermountain West
16	(see <u>Figure 3-72</u> in Section <u>3.8</u>), both models tend to under-predict maxima, but their
17	interquartile range agrees much better with observations. As <u>McDonald-Buller et al.</u>
18	(2011) noted, complex topography in some regions of the U.S. could influence surface O_3
19	through fine-scale, orographically induced flow regimes. In addition, numerical diffusion
20	broadly affects the ability of models to capture observed maxima, particularly at
21	mountain sites. <u>McDonald-Buller et al. (2011</u>) also note there are regions in the U.S.
22	where global models show consistent biases. For example, models are generally unable to
23	simulate the low O ₃ concentrations observed at Gulf Coast sites in summer during
24	onshore flow from the Gulf of Mexico, which could reflect marine boundary layer
25	chemistry and/or stratification that is not properly represented in the model. Both models
26	overpredict O ₃ at two sites in Florida (Sumatra and Indian River Lagoon). However,
27	further inland, CAMx tended to overpredict O3 at the Coffeeville, MS; Sand Mountain,
28	AL; and Georgia Station, GA sites whereas GEOS-Chem did not. The same is true for
29	higher elevation sites (Great Smoky Mountain, NC-TN; Shenandoah, VA). In the
30	Northeast, there is a general tendency for both models to overpredict the measured
31	distributions with somewhat higher maximum concentrations in CAMx compared to
32	GEOS-Chem and observations (see <u>Figure 3-69</u> to <u>Figure 3-74</u> in Section <u>3.8</u>).
33	The most readily discernible differences in model formulation are in the model grid
34	spacing and the treatment of wildfires. The finer resolution in CAMx allows for
35	topography to be better-resolved producing higher maximum O ₃ concentrations in the
36	Intermountain West. For wildfires, treatment differences include emission composition,
37	emission time averaging, and associated chemistry. Wildfires produce more O_3 in CAMx
38	simulations than in GEOS-Chem simulations, and Emery et al. (2012) attribute these
39	enhancements to shorter emission time averaging. The CAMx emissions average fire

1	emissions at hourly resolution based on the SmartFire algorithm, whereas GEOS-Chem
2	uses monthly averages from GFED2. Each model representation also uses different
3	emission compositions. The emissions used by Emery et al. (2012) include a larger
4	number of VOCs and additional categories of VOCs than used by Zhang et al. (2011).
5	Following emission, Emery et al. (2012) note that photochemical aging of wildfire
6	emissions depends on the chemical mechanism. Neither chemical mechanism was
7	designed specifically for these type of events. GEOS-Chem has traditionally focused on
8	the chemistry of the non-urban troposphere and does not represent secondary products of
9	fast reacting VOCs as does CB05. A lack of reactivity of secondary products would cause
10	a dampening of fire contributions to O ₃ . CB05 has traditionally focused on urban
11	chemistry and does not explicitly includes ketones (Henderson et al., 2011), which are
12	among the top ten VOCs emitted from fires (Andreae and Merlet, 2001). The O_3
13	increases seen in Emery et al. (2012) and Mueller and Mallard (2011a), however, are
14	subject to uncertainties in the representation of physics in the wildfire plumes. The
15	improvements in characterizing emissions would lead to smoke plumes that attenuate
16	light, thereby reducing photolysis and photoreactivity (e.g., Real et al., 2007). The
17	wildfires would also alter temperature and convective activity that influences plume rise
18	and the height of the planetary boundary layer. Emery et al. (2012) note the need for
19	more research to improve simulation of O ₃ from fires. Using a sensitivity analysis of
20	CAMx, the authors showed that removing wildfires in the West (California, Oregon, and
21	Idaho) resulted in reductions of NA background O ₃ of 10 to 50 ppb, with smaller
22	reductions elsewhere. Further, Emery et al. (2012) note that their calculated O ₃ increases
23	in the vicinity of wildfires is consistent with that of Mueller and Mallard (2011a).
24	Emery et al. (2012) captured the timing of a possible stratospheric intrusion at Gothic,
25	CO on April 19-20, 2006 and predicted an MDA8 value of ~73 ppb using CAMx on
26	April 20 compared to a measured value of 87 ppb. GEOS-Chem (at $0.5^{\circ} \times 0.667^{\circ}$)
27	predicted ~65 ppb for this event. The higher spatial resolution in CAMx likely
28	contributed to the improvement in model performance, but this may not be the only
29	factor. AM3, another global scale CTM (Lin et al., 2012) at $\sim 2^{\circ} \times 2.5^{\circ}$ resolution
30	predicted ~75 ppb for that event suggesting that differences in dynamical cores between
31	WRF and AM3, different treatments of the stratospheric O ₃ source, and perhaps the
32	spatial extent of the intrusion's effect on surface O ₃ should be considered in addition to
33	model resolution. Note that all three models (CAMx, GEOS-Chem, and AM3) under-
34	predicted the magnitude of this event. These results indicate a need for process-oriented
35	evaluation and targeted measurements that yield insight into both chemical and
36	dynamical processes. The R^2 for comparison of AM3 with observations of MDA8 O_3

37

resolution version of GEOS-Chem, 55.0 ppb for CAMx and 58.4 ppb for AM3 compared to 56.1 ppb for measurements (see Figure 3-75 in Section 3.8).

- 3 The results from either model have also been compared to more urban oriented sites in 4 the AQS network. As noted earlier, comparisons between model results and observations 5 become problematic near concentrated sources of O₃ precursors (NO_X and VOCs) in 6 urban cores. Emery et al. (2012) note that in coarse resolution models rural biogenic and 7 urban precursor emissions are mixed immediately leading to higher production efficiency 8 for O₃. Finer resolution models are better able to separate these two source categories and 9 to resolve features of urban chemistry such as titration of O₃ by NO_X emitted by traffic 10 and subsequent processing of NO_X emissions during transport downwind. CAMx at 11 12×12 km resolution is better able to capture these features than GEOS-Chem at 12 50×50 km resolution. Both models tend to over-predict O₃ at the low O₃ concentrations 13 in areas where O_3 scavenging by NO_x is evident. In these situations, NA background O_3 14 concentrations are often higher than in the respective models for the base case. At high 15 O_3 concentrations downwind of source areas, both models predict NA background O_3 16 concentrations that are much lower than observed or base case O_3 . The latter results are in 17 accord with results shown in Figure 3-11 for rural CASTNET sites at low elevations, 18 which show lower ratios between NA background O₃ and either observations or base case 19 O_3 at high O_3 than at low O_3 concentrations.
- 20 Figure 3-15 shows the annual 4th highest MDA8 O₃ predicted by GEOS-Chem (at 21 $0.5^{\circ} \times 0.667^{\circ}$ resolution) for the base case (upper panel), and corresponding U.S. 22 background (middle panel) and NA background (lower panel) MDA8 O₃ on the same 23 days for 2006. Figure 3-16 shows corresponding values predicted by CAMx for the base 24 case (upper panel) and NA background (lower panel) MDA8 O₃ on the same days for 25 2006. As can be seen from Figure 3-15 and Figure 3-16, on those days when models predicted their annual 4th highest MDA8 O3, the corresponding NA background 26 27 concentrations are 36 ± 9 ppb in the eastern U.S. Base case concentrations are much 28 higher indicating that regional pollution is mainly responsible for the models 4th highest 29 concentrations. In the western U.S. on the other hand, NA background concentrations are 30 generally higher and make up a larger fraction of the calculated 4th highest MDA8 O_3 in 31 both models, but for different reasons. GEOS-Chem predicts highest values in the 32 Southern Rockies because of over-production of NO_x by lightning. CAMx predicts 33 highest values in ID, OR and WA from wildfires. The CAMx run includes day specific 34 values for area burned, but GEOS-Chem uses monthly averages. (A more recent version 35 of GEOS-Chem also incorporates day specific estimates for area burned.) Remaining 36 areas of relatively high background levels (>60 ppb) are due mainly to some combination 37 of stratospheric intrusions and Eurasian emissions. There are a few examples that can be 38 used to give a rough idea of the magnitudes of episodically high background

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1	contributions. A comparison of the annual 4th highest MDA8 O ₃ concentration simulated
2	by CAMx including wildfires and omitting them indicates that wildfires contributed ~ 30
3	to 40 ppb in Idaho, Montana, and Washington with a potentially larger contribution in the
4	upper northwestern corner of California. Estimated contributions from strong
5	stratospheric intrusions to surface O_3 in AM3 could range up to ~ 70 ppb in the western
6	U.S.

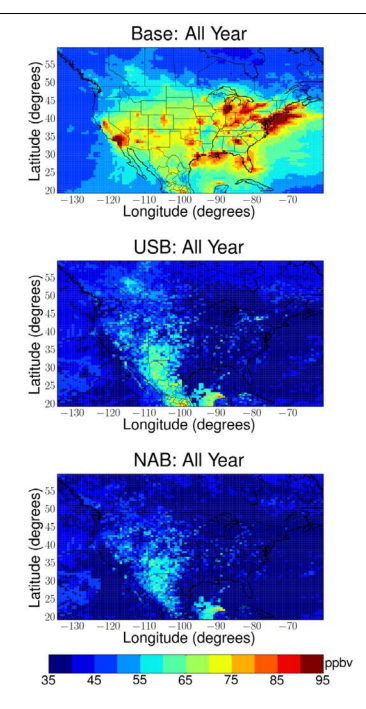
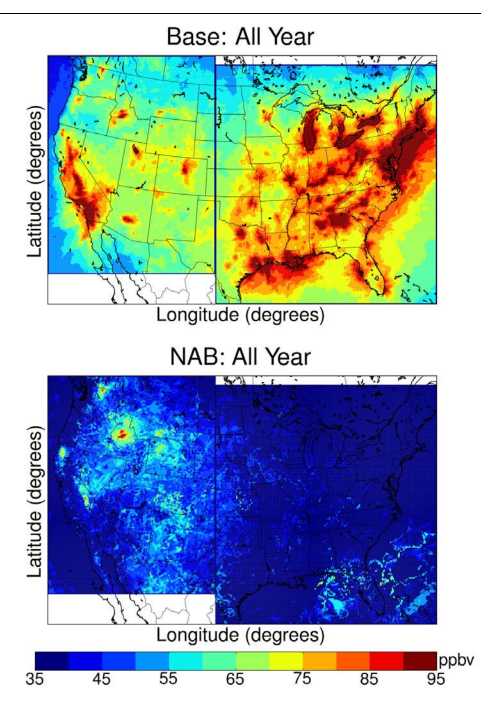




Figure 3-15 Annual 4th highest MDA8 ozone predicted by GEOS-Chem (0.5° × 0.667°) for the base case (Base) with corresponding U.S. background (USB) and NA background (NAB) MDA8 ozone for the same days in 2006.



Source: Adapted from Emery et al. (2012).

Figure 3-16 Annual 4th highest MDA8 ozone predicted by CAMx for the base case (Base) and corresponding NA background (NAB) MDA8 ozone for the same days in 2006.

1	All models undergo continuous updating of inputs, parameterizations of physical and
2	chemical processes, and improvements in model resolution. Inputs that might be
3	considered most relevant include emissions inventories, chemical reactions, and
4	meteorological fields. This leads to uncertainty in model predictions in part because there
5	is typically a lag between updated information for the above inputs—as outlined in
6	Section 3.2 for chemical processes and emissions and in Section 3.3 for model
7	construction—and their implementation in CTMs including GEOS-Chem or the other
8	models described above. Quantitative estimates of uncertainties from meteorological and
9	emission inputs and chemical mechanisms are problematic because simulations designed
10	to quantify uncertainties from these sources have not been performed for these model
11	runs. At best, these uncertainties can be estimated by comparison with observations while
12	recognizing that compensating errors likely exist.
13	Since NA background is a construct that cannot be directly measured, the range of
14	background O_3 concentrations must be estimated using CTMs. Results from the Zhang et
15	al. (2011) GEOS-Chem and Emery et al. (2012) GEOS-Chem/CAMx (hereafter referred
16	to as CAMx) model estimates were chosen for further analysis because these models
17	have produced the latest estimates for background O3 concentrations documented in the
18	open literature. The main results from these two modeling efforts can be described as
19	follows:
20	 Both models show background concentrations vary spatially and temporally;
21	 Simulated mean background concentrations are highest in the Intermountain
22	West (i.e., at high altitude) in spring and lowest in the Northeast during
23	summer;
24	 Background concentrations tend to increase with total (i.e., base case) O₃
25	concentrations at high elevation, but that tendency is not as clear at low
26	elevations.
27	The most pronounced differences between the Zhang et al. (2011) GEOS-Chem and the
28	Emery et al. (2012) CAMx models—when compared with observations—are at the upper
29	end of the concentration distribution. At high elevations, differences are likely to be the
30	result of underpredictions of background contributions which are driven mainly by
31	episodic events such as stratospheric intrusions and wildfires. In general, CAMx predicts
32	higher values at the upper end of the concentration distribution than does GEOS-Chem.
33	At low elevations (<1,500 meters)—located mainly in the East—the reasons for
34	underpredictions at the upper end of the concentration distribution are more complex and
35	likely involve extensive interactions between anthropogenic and natural sources.
36	
50	Table 3-1 summarizes modeling results for seasonal mean MDA8 O ₃ by region simulated
30 37	<u>Table 3-1</u> summarizes modeling results for seasonal mean MDA8 O_3 by region simulated by the two models. The regions in <u>Table 3-1</u> are shown in <u>Figure 3-50</u> . As can be seen

1	from the table, seasonal means predicted by GEOS-Chem are within a few parts per
2	billion of measurements in both spring and summer for all regions shown except for
3	California in the spring. Although CASTNET sites are meant to represent regional
4	background air, they can be heavily influenced by polluted air masses, particularly in
5	California where the underpredictions are largest. Seasonal means are simulated by
6	CAMx to within 2-5 ppb except in California in the spring where they are underpredicted
7	by 8 ppb and at sites in the Northeast and Southeast where they are overpredicted by 8-
8	9 ppb in summer. When compared to observations, the mean R^2 within each region—
9	except for California in the spring—is higher for CAMx than for GEOS-Chem suggesting
10	better ability to track day-to-day variability by CAMx. It is clear from these results that
11	model resolution (at least for the model resolutions considered here) is not the dominant
12	factor determining agreement of the means between simulations or between simulations
13	and measurements. Differences in model chemistry and physics must also be considered.

Table 3-1 Comparison of seasonal mean MDA8 ozone concentrations simulated by the GEOS-Chem and CAMx base case models for 2006, with measurements at CASTNET sites.

Region	CASTNET		GEOS-Chem		CAMx	
	Spring	Summer	Spring	Summer	Spring	Summer
California (5) ^a	58 ± 12 ^b	69 ± 14	52 ± 11; 0.52 ^c	66 ± 18; 0.22	50 ± 10; 0.50	66 ± 13; 0.30
			38 ± 7^{d}	37 ± 9	39 ± 6	42 ± 6
West (14)	54 ± 9	55 ± 11	53 ± 7; 0.30	55 ± 11; 0.12	49 ± 8; 0.39	57 ± 10; 0.33
			42 ± 6	40 ± 9	40 ± 7	41 ± 8
North Central (6)	47 ± 10	50 ± 12	47 ± 8; 0.52	51 ± 14; 0.44	45 ± 11; 0.63	54 ± 13; 0.48
			33 ± 6	27 ± 7	30 ± 6	31 ± 5
Northeast (5)	48 ± 10	45 ± 14	45 ± 7; 0.44	45 ± 13; 0.47	46 ± 11; 0.53	53 ± 14; 0.54
			33 ± 7	24 ± 7	30 ± 5	27 ± 6
Southeast (9)	52 ± 11	52 ± 16	51 ± 7; 0.42	54 ± 9; 0.21	54 ± 9; 0.56	61 ± 12; 0.45
			32 ± 7	29 ± 10	33 ± 6	30 ± 6

^aValues in parentheses after each region name refer to the number of sites.

^bShown are seasonal (spring, summer) mean MDA8 O₃ concentrations (ppb ± standard deviation);

^cShown are mean R² of all model-measurement pairs at individual CASTNET sites.

^dNorth American (NA) background seasonal mean MDA8 O₃ concentrations (ppb ± standard deviation) are shown beneath the base case seasonal means.

Source: Data from Zhang et al. (2011) for GEOS-Chem and Emery et al. (2012) for CAMx.

14 Table 3-2 summarizes modeling results for the annual 4th highest (99th –percentile)

- MDA8 O_3 for the same seasons and regions used in <u>Table 3-1</u>. As can be seen, the
- 16 GEOS-Chem and the CAMx models both underestimate mean day specific 4th highest
- 17
 - values in California by ~20 ppb. In general, CAMx simulates MDA8 O₃ concentrations
- 18 that are higher and in better agreement with measurements. Shown alongside the model
- 19 estimates is the number of days the modeled MDA8 O₃ concentrations are within 5 ppb

1	of observed. The lower portions of the entries for the models in <u>Table 3-2</u> show model
2	predicted 4th highest MDA8 O ₃ concentrations that are not calculated on the same day as
3	the 4th highest values measured at CASTNET sites. It can be seen that simulated regional
4	means of the 4th highest MDA8 O ₃ are in better agreement with measurements when
5	results are un-paired by date. In other words, the models do not predict their annual
6	4th highest MDA8 O ₃ concentrations on the same day as they are observed.
7	These results underscore the uncertainties inherent in any model's attempts to simulate
8	day specific 4th highest O ₃ concentrations. As noted earlier, uncertainties in calculating
9	day specific O_3 concentrations are especially challenging because of the lack of day
10	specific data for emissions of many species. While progress is being made in obtaining
11	day specific data for lightning strikes and area burned in wildfires, the emission factors
12	for precursors from these episodic sources such as lightning and wildfires are still
13	uncertain. In addition to uncertainty in emissions, uncertainties in models' treatments of
14	transport and chemical mechanisms must also be considered.
15	Comparison of GEOS-Chem results for natural and NA background indicate that
16	methane is also a major contributor to NA background O ₃ , accounting for slightly less
17	than half of the increase in background since the preindustrial era and whose relative
18	contribution is projected to grow in the future. U.S. background concentrations are on
19	average 2.6 ppb higher than NA background concentrations during spring and 2.7 ppb
20	during summer across the United States. Highest values for U.S. background (in the U.S.)
21	are found over the Northern Tier of New York State (19.1 ppb higher than local NA
22	background concentrations) in summer. High values are also found in other areas
23	bordering Canada and Mexico.

Table 3-2Comparison of annual 4th-highest MDA8 ozone concentrations
measured at CASTNET sites in 2006 with MDA8 ozone
concentrations simulated by the GEOS-Chem and CAMx base case
models.

Region	CASTNET	GEOS-Chem		CAMx	
California (5) ^a	90 ± 13 ^b	71 ± 15 [°] 85 ± 19 [°]	0 ^d	71 ± 9 85 ± 13	0
West (14)	70 ± 4	62 ± 8 68 ± 7	4	63 ± 8 71 ± 7	6
North Central (6)	71 ± 5	58 ± 10 69 ± 10	1	63 ± 7 73 ± 8	1
Northeast (5)	71 ± 4	61 ± 6 68 ± 5	0	72 ± 7 75 ± 3	3
Southeast (9)	76 ± 8	61 ± 6 71 ± 5	2	71 ± 11 79 ± 9	5

^aValues in parentheses after each region name refer to the number of sites.

^bShown are annual 4th highest (99th-percentile) MDA8 O₃ concentration regional means (ppb ± standard deviation).

^cShown are calculated MDA8 O_3 concentrations on days when the 4th highest MDA8 O_3 concentrations was measured. ^dShown are the number of days the model predicted MDA8 O_3 concentrations were within 5 ppb of observed 4th-highest concentrations.

 $^{e}\mbox{Shown}$ are model predicted annual 4th highest MDA8 O_{3} concentrations.

Source: Data from Zhang et al. (2011) for GEOS-Chem and Emery et al. (2012) for CAMx.

1	Analyses of results from GEOS-Chem and CAMx presented here and shown in <u>Table 3-1</u>
2	and Table 3-2 are in accord with results from Kasibhatla and Chameides (2000) who
3	found that the accuracy of simulations improved as the averaging time of both the
4	simulation and the observations increased (see Section 3.3). Note that any CTM—not just
5	the ones considered here-will have difficulty in predicting day specific quantities. When
6	analyzing results over long time periods (e.g., months), special care should be taken to
7	examine temporal trends in bias because this will improve understanding of the modeling
8	results.
9	Overall, these results suggest that GEOS-Chem is capable of simulating seasonal or
10	monthly mean MDA8 O_3 to within a few parts per billion on a regional basis throughout
11	the U.S., except in California. These results suggest that CAMx is capable of simulating
12	seasonal or monthly mean MDA8 O_3 to within a few ppb, though, CAMx also shows
13	relatively large disagreements in California and, in addition, shows relatively large
14	positive bias in seasonal mean MDA8 O_3 in the eastern U.S. However, differences
15	between the models in the East are likely to narrow with updates to chemistry. Neither
16	model is capable of simulating 4th highest MDA8 O_3 to within suitable bounds on a
17	day-specific basis at all sites, or even most sites. However, agreement between simulated

1	vs. observed 4th highest MDA8 O ₃ is improved for either model when the models and the
2	measurements are sampled on different days.
3	Note that the calculations of background concentrations presented in this section were
4	formulated to answer the question, "what would O3 concentrations be if there were no
5	anthropogenic sources". This is different from asking, "how much of the O ₃ measured or
6	simulated in a given area is due to background contributions". Because of potentially
7	strong non-linearities (i.e., the fate, or lifetime, of the background O ₃ transported into the
8	urban area will depend on the concentration of the background O ₃ in addition to
9	interactions of background O ₃ with the local chemical regime) in many urban areas, these
10	estimates by themselves should not be used to answer the second question posed above.
11	The extent of these non-linearities will generally depend on location and time, the
12	strength of concentrated sources and the nature of the chemical regime. Further work is
13	needed on how these estimates of regional background concentrations can be used to help
14	determine the contributions of background sources of O_3 to urban concentrations.

3.5 Monitoring

3.5.1 Routine Monitoring Techniques

15	The federal reference method (FRM) for O_3 measurement is called the
16	Chemiluminescence Method (CLM) and is based on the detection of chemiluminescence
17	resulting from the reaction of O_3 with ethylene gas. The UV absorption photometric
18	analyzers were approved as federal equivalent methods (FEMs) in 1977 and gained rapid
19	acceptance for NAAQS compliance purposes due to ease of operation, relatively low
20	cost, and reliability. The UV absorption method is based on the principle that O_3
21	molecules absorb UV radiation at a wavelength of 254 nm from a mercury lamp. The
22	concentration of O_3 is computed from Beer's law using the radiation absorbed across a
23	fixed path length, the absorption coefficient, and the measured pressure and temperature
24	in the detection cell. UV absorption photometry is the predominant method for assessing
25	compliance with the NAAQS for O ₃ . Almost all of the state and local air monitoring
26	stations (SLAMS) that reported data to EPA AQS from 2005 to 2009 used UV absorption
27	photometer FEMs. No CLM monitors, approved as FRMs or FEMs, reported O_3 data to
28	AQS from 2005 to 2009 and only one monitor reported data using a long-path or open
29	path Differential Optical Absorption Spectrometer (DOAS) FEM during this period.
30	The rationale history and calibration of Ω measurements were summarized in the 1006
	The rationale, history, and calibration of O_3 measurements were summarized in the 1996
31	and 2006 O_3 AQCDs (U.S. EPA, 2006b, 1996a) and focused on the state of ambient O_3

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measurements at that time as well as evaluation of interferences and new developments. This discussion will continue with the current state of O_3 measurements, interferences, and new developments for the period 2005 to 2010.

UV O_3 monitors use mercury lamps as the source of UV radiation and employ an O_3 scrubber (typically manganese dioxide) to generate an ozone-free air flow to serve as a reference channel for O_3 measurements. There are known interferences with UV O_3 monitors. The 2006 O_3 AQCD (U.S. EPA, 2006b) reported on the investigation of the effects of water vapor, aromatic compounds, ambient particles, mercury vapor and alternative materials in the instrument's O_3 scrubber. The overall conclusions from the review of the scientific literature covered in the 2006 O_3 AQCD (U.S. EPA, 2006b) are briefly summarized below.

12 Kleindienst et al. (1993) found water vapor to have no measurable impact and aromatic 13 compounds to have a minor impact (as much as 3% higher than the FRM extrapolated to 14 ambient conditions) on UV absorption measurements. UV O₃ monitor response evaluated 15 by chamber testing using cigarette smoke, reported an elimination of the O₃ monitor 16 response to the smoke when a particle filter was used that filtered out particles less than 17 0.2 µm in diameter (Arshinov et al., 2002). One study (Leston et al., 2005) in 18 Mexico City compared a UV O3 FEM to a CLM FRM. The UV FEM reported 19 consistently higher O_3 than the CLM FRM. They suggested that O_3 measured in ambient 20 air could be too high by 20 to 40 ppb under specific conditions due to positive 21 interference by a number of organic compounds, mainly those produced during the 22 oxidation of aromatic hydrocarbons and some primary compounds such as styrene and 23 naphthalene. However, the concentrations of these compounds were many times higher in 24 both of these environments than are typically found at ambient air monitoring sites in the 25 U.S. Although Hg is also potentially a strong interfering agent, because the Hg resonance 26 line is used in this technique, its concentration would also have to be many times higher 27 than is typically found in ambient air, e.g., as might be found in power plant plumes. 28 Thus, it seems unlikely that such interferences would amount to more than one or two 29 ppb (within the design specifications of the FEM), except under conditions conducive to 30 producing high concentrations of the substances they identified as causing interference. 31 Leston et al. (2005) also presented smog chamber data which demonstrated that heated 32 metal and heated silver wool scrubbers perform better in the presence of aromatic 33 hydrocarbon irradiations than manganese dioxide scrubbers when compared to the FRM. 34 They also suggested the use of humidified calibration gas and alternative scrubber 35 materials to improve UV O₃ measurements. Some O₃ monitor manufacturers now offer 36 heated silver wool scrubbers as an alternative to manganese dioxide. Another possible 37 solution to the O_3 scrubber problem may be the use of a gas phase scrubber such as NO. 38 A commercial version of this has recently been introduced by 2B Technologies as an

option on their model 202 FEM; however, it has not been field tested or approved for use as an FEM.

Review of the recent literature is summarized below. Study of UV monitors by <u>Williams</u> et al. (2006) concluded that well maintained monitors showed no substantial interferences when operated in locations with high concentrations of potentially interfering VOCs including Nashville, Houston, and the Gulf of Maine. Monitors were tested in urban and suburban environments, as well as on board a ship in both polluted and clean marine air. Comparisons of UV measurements to a non-FRM/FEM NO based CLM demonstrated agreement to within 1%. At the Houston location, they did observe a brief period on one day for about 30 minutes where the UV measurements exceeded the CLM by about 8 ppb (max). This was attributed to probable instrument malfunction.

- 12 Wilson and Birks (2006) investigated water vapor interference in O_3 measurements by 13 four different UV monitors. In extreme cases where a rapid step change in relative 14 humidity between 0 and 90% was presented, large transitory responses (tens to hundreds 15 of ppb) were found for all monitors tested. Rapid changes in relative humidity such as 16 this would not be expected during typical ambient O_3 measurements and could only be 17 expected during measurement of vertical profiles from balloon or aircraft. The magnitude 18 of the interference and the direction (positive or negative) was dependent on the 19 manufacturer and model. Wilson and Birks (2006) also hypothesized that water vapor 20 interference is caused by physical interactions of water vapor on the detection cell. The 21 O_3 scrubber was also thought to act as a reservoir for water vapor and either added or 22 removed water vapor from the air stream, subsequently affecting the detector signal and 23 producing either a positive or negative response. They demonstrated that the use of a 24 Nation permeation membrane just before the O_3 detection cell to remove water vapor 25 eliminated this interference.
- 26 Dunlea et al. (2006) evaluated multiple UV O₃ monitors with two different O₃ scrubber 27 types (manganese dioxide and heated metal wool) in Mexico City. Large spikes in O_3 28 concentrations were observed while measuring diesel exhaust where large increases in 29 particle number density were observed. The interference due to small particles passing 30 through the Teflon filter and scattering/absorbing light in the detection cell were 31 estimated to cause at most a 3% increase in measurements in typical ambient air 32 environments. This estimate pertains to measurements in the immediate vicinity of fresh 33 diesel emissions and most monitor siting guidelines would not place the monitor close to 34 such sources, so actual interferences are expected to be much less than 3%. Dunlea et al. 35 (2006) also observed no evidence for either a positive or negative interference or 36 dependence due to variations in aromatics during their field study.

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1	Li et al. (2006c) verified early reports of gas phase mercury interference with the UV O_3
2	measurement. They found that 300 ng/m ³ of mercury produced an instrument response of
3	about 35 ppb O_3 . Background concentrations of mercury are around 1-2 ng/m ³ and
4	expected to produce an O_3 response that would be <1 ppb.
5	Spicer et al. (2010) examined potential UV O_3 monitor interferences by water vapor,
6	mercury, aromatic compounds, and reaction products from smog chamber simulations.
7	Laboratory tests showed little effect of changing humidity on conventional FEM UV O ₃
8	monitors with manganese dioxide or heated metal wool scrubbers in the absence of other
9	interferences. Mercury vapor testing produced an O3 response by the UV monitors that
10	was <1 ppb O_3 per 1 ppt (about 8 ng/m ³) mercury vapor. Interference by aromatic
11	compounds at low (3% RH) and high (80% RH) humidity showed some positive
12	responses that varied by UV monitor and ranged from 0 to 2.2 ppb apparent O ₃ response,
13	per ppb of aromatic compound tested. The authors acknowledged that the aromatic
14	compounds most likely to interfere are rarely measured in the atmosphere and therefore,
15	make it difficult to assess the impact of these compounds during ambient air monitoring.
16	Comparison of UV and CLM responses to photochemical reaction products in smog
17	chamber simulations at 74 to 85% RH showed varied responses under low
18	(0.125 ppmv/0.06 ppmv) to high (0.50 ppmv/0.19 ppmv) hydrocarbon/NO _X conditions.
19	The conventional UV monitors were as much as 2 ppb higher than the CLM under low
20	hydrocarbon/NO _X conditions and 6 ppb higher under the high hydrocarbon/NO _X
21	conditions. Two FEM UV monitors were also co-located at six sites in Houston from
22	May to October, 2007 with one UV monitor equipped with Nafion permeation
23	membrane. The average difference between 8-h daily max O ₃ concentrations using the
24	UV and the UV with Nafion permeation membrane ranged from -4.0 to 4.1 ppb.

3.5.2 Precision and Bias

25	In order to provide decision makers with an assessment of data quality, EPA's Quality
26	Assurance (QA) group derives estimates of both precision and bias for O_3 and the other
27	gaseous criteria pollutants from the biweekly single point quality control (QC) checks
28	using calibration gas, performed at each site by the monitoring agency. The single-point
29	QC checks are typically performed at concentrations around 90 ppb. Annual summary
30	reports of precision and bias can be obtained for each monitoring site at
31	http://www.epa.gov/ttn/amtic/qareport.html. The assessment of precision and bias are
32	based on the percent-difference values, calculated from single-point QC checks. The
33	percent difference is based on the difference between the pollutant concentration
34	indicated by monitoring equipment and the known (actual) concentration of the standard
35	used during the QC check. The monitor precision is estimated from the 90% upper

1	confidence limit of the coefficient of variation (CV) of relative percent difference (RPD)
2	values. The bias is estimated from the 95% upper confidence limit on the mean of the
3	absolute values of percent differences. The data quality goal for O_3 precision and bias at
4	the 90 and 95% upper confidence limits is 7% (40 CFR Part 58, Appendix A). Table 3-3
5	presents a summary of the number of monitors that meet the precision and bias goal of
6	7% for 2005 to 2009. Greater than 96% of O_3 monitors met the precision and bias goal
7	between 2005 and 2009. Another way to look at the precision (CV) and bias (percent
8	difference) information using the single-point QC check data from the monitoring
9	network is to present box plots of the monitors' individual precision and percent-
10	difference data; Figure 3-17 and Figure 3-18 include this information for O_3 monitors
11	operating from 2005 to 2009.

Table 3-3Summary of ozone monitors meeting 40 CFR Part 58, Appendix A
Precision and Bias Goals

Year	Number of Monitors	Monitors with Acceptable Precision (%)	Monitors with Acceptable Bias (%)
2005	879	96.5	96.7
2006	881	98.1	97.6
2007	935	98.1	98.1
2008	955	97.1	96.7
2009	958	97.4	97.5

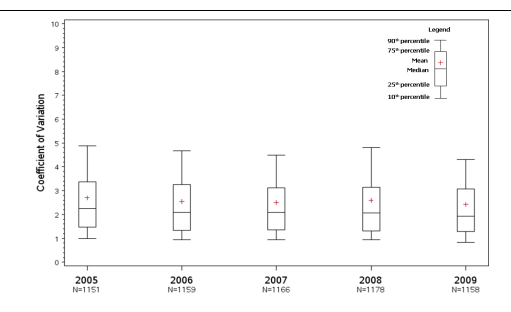


Figure 3-17 Box plots of precision data by year (2005-2009) for all ozone monitors reporting single-point QC check data to AQS.

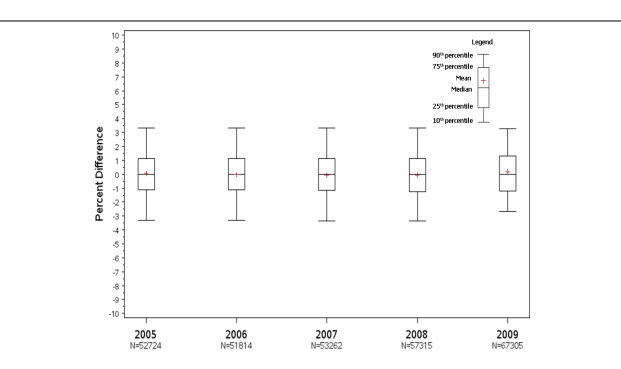


Figure 3-18 Box plots of percent-difference data by year (2005-2009) for all ozone monitors reporting single-point QC check data to AQS.

3.5.2.1 Precision from Co-located UV Ozone Monitors in Missouri

1	The Missouri Department of Natural Resources (MODNR) maintains a network of co-
2	located UV O ₃ analyzers. The MODNR provided co-located data from four monitors:
3	two co-located at the same monitoring site in Kansas City (AQS ID 290370003) and two
4	co-located at the same monitoring site in St. Louis (AQS ID 291831002). Hourly
5	observations for the co-located measurements at these two sites between April and
6	October, 2006-2009 were used to evaluate precision from co-located UV monitors. These
7	data were then compared with the precision obtained by the biweekly single point QC
8	checks for all sites reporting single-point QC check data to AQS between 2005 and 2009;
9	the method normally used for assessing precision. Box plots of the RPD between the
10	primary and co-located hourly O ₃ measurements in Missouri are shown in Figure 3-19
11	and box plots of the RPD between the actual and indicated QC check for all U.S. sites are
12	shown in Figure 3-20. As mentioned above, the average concentration of the single-point
13	QC check is 90 ppb, whereas the average ambient O_3 concentration measured at the two
14	sites in Missouri was 34 ppb. The mean RPD for the co-located monitors in Missouri and
15	the single-point QC check data from all sites were less than 1 percent.

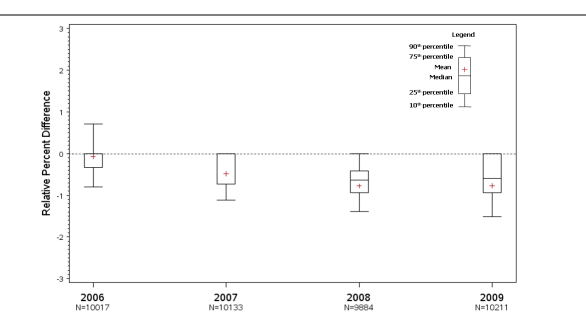


Figure 3-19 Box plots of RPD data by year for the co-located ozone monitors at two sites in Missouri from 2006-2009.

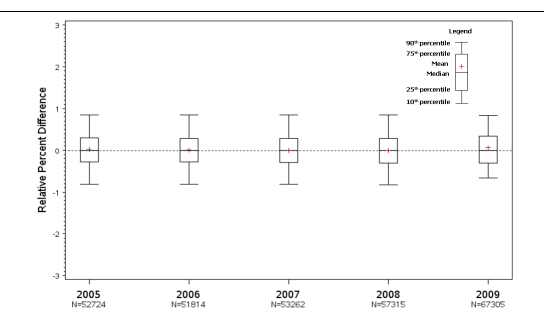


Figure 3-20 Box plots of RPD data by year for all U.S. ozone sites reporting single-point QC check data to AQS from 2005-2009.

3.5.3 Performance Specifications

1	The performance specifications for evaluating and approving new FEMs in accordance
2	with 40 CFR Part 53 are provided in <u>Table 3-4</u> . These specifications were developed and
3	originally published in the Federal Register in 1975. Modern, commercially-available
4	instruments can now perform much better than the requirements specified below. For
5	example, the lower detectable limit (LDL) performance specification is 10 ppb and the
6	typical vendor-stated performance for the LDL is now less than 0.60 ppb. The amount of
7	allowable interference equivalent for total interference substances is 60 ppb, and the
8	current NAAQS for O_3 is 75 ppb, with an averaging time of 8 hours. Improvements in
9	new measurement technology have occurred since these performance specifications were
10	originally developed. These specifications should be revised to more accurately reflect
11	the necessary performance requirements for O3 monitors used to support the current
12	NAAQS.

Parameter	Specification	
Range	0 – 0.5 ppm (500 ppb)	
Noise	0.005 ppm (5 ppb)	
LDL – defined as two times the noise	0.01 ppm (10 ppb)	
Interference equivalent		
Each interfering substance	± 0.02 ppm (20 ppb)	
Total interfering substances	0.06 ppm (60 ppb)	
Zero drift		
12 h	± 0.02 ppm (20 ppb)	
24 h	± 0.02 ppm (20 ppb)	
Span Drift, 24 h		
20% of upper range limit	± 20.0%	
80% of upper range limit	±5.0%	
Lag time	20 min	
Rise time	15 min	
Fall time	15 min	
Precision		
20% of upper range limit	0.01 ppm (10 ppb)	
80% of upper range limit	0.01 ppm (10 ppb)	

Table 3-4Performance specifications for ozone based in 40 CFR Part 53

3.5.4 Monitor Calibration

1	The calibration of O_3 monitors was summarized in detail in the 1996 O_3 AQCD (U.S.
2	EPA, 1996a). The calibration of O_3 monitors is done using an O_3 generator and UV
3	photometers. UV photometry is the prescribed procedure for the calibration of reference
4	methods to measure O ₃ in the atmosphere. Because O ₃ is unstable and cannot be stored,
5	the O_3 calibration procedure specifically allows the use of transfer standards for
6	calibrating ambient O_3 monitors. A transfer standard is calibrated against a standard of
7	high authority and traceability and then moved to another location for calibration of O_3
8	monitors. The EPA and the National Institute of Standards and Technology (NIST) have
9	established a network of standard reference photometers (SRPs) that are used to verify
10	transfer standards. The International Bureau of Weights and Measures (BIPM) maintain
11	one NIST SRP (SRP27) as the World's O_3 reference standard. NIST maintains two SRPs
12	(SRP0 and SRP2) that are used for comparability to ten other SRPs maintained by the
13	EPA's Regional QA staff.
14	SRPs have been compared to other reference standards. Tanimoto et al. (2006) compared
15	NIST SRP35, owned by the National Institute for Environmental Studies in Japan, to gas
-	

1	phase titration (GPT). The SRP was found to be 2% lower than GPT. GPT is no longer
2	used as a primary or transfer standard in the U.S. Viallon et al. (2006) compared SRP27
3	built at BIPM to four other NIST SRPs maintained by BIPM (SRP28, SRP31, SRP32,
4	and SRP33). A minimum bias of +0.5% was found for all SRP measurement results, due
5	to use of the direct cell length measurement for the optical path length; this bias was
6	accounted for by applying the appropriate correction factor. Study of the bias-corrected
7	SRPs showed systematic biases and measurement uncertainties for the BIPM SRPs. A
8	bias of -0.4% in the instrument O_3 mole fraction measurement was identified and
9	attributed to non-uniformity of the gas temperature in the instrument gas cells, which was
10	compensated by a bias of $+0.5\%$ due to an under-evaluation of the UV light path length
11	in the gas cells. The relative uncertainty of the O_3 absorption cross section was 2.1% at
12	253.65 nm and this was proposed as an internationally accepted consensus value until
13	sufficient experimental data is available to assign a new value.
14	In November, 2010, the EPA revised the Technical Assistance Document for Transfer
15	Standards for Calibration of Air Monitoring Analyzers for Ozone (U.S. EPA, 2010f) that
16	was first finalized in 1979 (U.S. EPA, 1979b). The revision removed methods no longer
17	in use and updated definitions and procedures where appropriate. In the revised
18	document, the discussion of transfer standards for O ₃ applies to the family of standards
19	that are used beyond SRPs or Level 1 standards. To reduce confusion, EPA reduced the
20	number of common terms that were used in the past such as: primary standard, local

number of common terms that were used in the past such as: primary standard, local primary standard, transfer standard, and working standard. Beyond the SRPs, all other standards are considered transfer standards.

3.5.5 Other Monitoring Techniques

3.5.5.1 Portable UV Ozone Monitors

23	Small, lightweight, and portable UV O ₃ monitors with low power consumption are
24	commercially available. These monitors are based on the same principle of UV
25	absorption by O_3 at 254 nm. Monitors of this type are typically used for vertical profiling
26	using balloons, kites, or light aircraft where space and weight are limited. They have also
27	been used for monitoring at remote locations such as National Parks. Burley and Ray
28	(2007) compared portable O_3 monitor measurements to those from a conventional UV
29	monitor in Yosemite National Park. Calibrations of the portable O_3 monitors against a
30	transfer standard resulted in an overall precision of ± 4 ppb and accuracy of $\pm 6\%$. Field
31	measurement comparisons between the portable and conventional monitor at Turtleback
32	Dome showed the portable monitor to be 3.4 ppb lower on average, with daytime

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1	deviation typically on the order of 0-3 ppb. Agreement between the portable and
2	conventional monitor during daylight hours (9:00 a.m. to 5:00 p.m. PST) resulted in an
3	R^2 of 0.95, slope of 0.95, and intercept of 0.36 ppb. Substantial deviations were observed
4	in the predawn hours where the portable monitor was consistently low. These deviations
5	were attributed to the difference in sampling inlet location. The portable monitor was
6	located at 1.3 meters above ground and the conventional monitor was located at
7	10 meters above ground. Agreement between the portable and conventional monitors for
8	all hours sampled resulted in an R^2 of 0.88, slope of 1.06, and intercept of -6.8 ppb.
9	(Greenberg et al., 2009) also compared a portable UV O ₃ monitor to a conventional UV
10	monitor in Mexico City and obtained good agreement for a 14 day period with an R^2 of
11	0.97, slope of 0.97, and intercept of 6 ppb. One portable O_3 monitor was recently
12	approved as an FEM (EQOA-0410-190) on April 27, 2010 (75 FR 22126).

3.5.5.2 NO-based Chemiluminescence Monitors

13	One commercially available NO-based chemiluminescence monitor has been approved as
14	an FEM (EQOA-0611-199) on October 7, 2011 (75 FR 62402). It may also be designated
15	as a second or replacement FRM since the ethene based FRMs are no longer
16	manufactured. Although this is a relatively new monitor, other NO-based CLM
17	instruments have been custom built for various field studies since the early 1970s. A
18	commercial version that measured both O_3 and NO_X was offered in the early 1970s but
19	failed to gain commercial acceptance. Initial testing with SO ₂ , NO ₂ , Cl ₂ , C ₂ H ₂ , C ₂ H ₄ and
20	C_3H_6 (Stedman et al., 1972) failed to identify any interferences. In the intervening years,
21	custom built versions have not been found to have any interference; however, they do
22	experience a slight decrease in response with increasing relative humidity (due to
23	quenching of the excited species by the water molecules). The new NO-based CLM
24	solves this problem with the use of a Nafion membrane dryer. A custom built NO-based
25	CLM similar to the FEM was used by Williams et al. (2006) in Houston, TX; Nashville,
26	TN; and aboard ship along the New England coast. It was found to be in good agreement
27	with a standard UV based FEM and with a custom built DOAS.

3.5.5.3 Passive Air Sampling Devices and Sensors

28	A passive O_3 sampling device depends on the diffusion of O_3 in air to a collecting or
29	indicating medium. In general, passive samplers are not adequate for compliance
30	monitoring because of the limitations in averaging time (typically one week or more),
31	particularly for O ₃ . However, these devices are valuable for personal human exposure
32	estimates and for obtaining long-term data in rural areas where conventional UV

1	monitors are not practical or feasible to deploy. The 1996 O ₃ AQCD (U.S. EPA, 1996a)
2	provided a detailed discussion of passive samplers, along with the limitations and
3	uncertainties of the samplers evaluated and published in the literature from 1989 to 1995.
4	The 2006 O ₃ AQCD (U.S. EPA, 2006b) provided a brief update on available passive
5	samplers developed for use in direct measurements of personal exposure published
6	through 2004. The 2006 O ₃ AQCD (U.S. EPA, 2006b) also noted the sensitivity of these
7	samplers to wind velocity, badge placement, and interference by other co-pollutants that
8	may result in measurement error.
9	Subsequent evaluations of passive diffusion samplers in Europe showed good correlation
10	when compared to conventional UV O ₃ monitors, but a tendency for the diffusion
11	samplers to overestimate the O ₃ concentration (Gottardini et al., 2010; Vardoulakis et al.,
12	<u>2009</u> ; <u>Buzica et al., 2008</u>). The bias of O_3 diffusion tubes were also found to vary with
13	concentration, season, and exposure duration (Vardoulakis et al., 2009). Development of
14	simple, inexpensive, passive O_3 measurement devices that rely on O_3 detection papers
15	and a variety of sensors with increased time resolution (sampling for hours instead of
16	weeks) and improved sensitivity have been reported (Maruo et al., 2010; Ebeling et al.,
17	2009; Miwa et al., 2009; Ohira et al., 2009; Maruo, 2007; Utembe et al., 2006).
18	Limitations for some of these sensors and detection papers include air flow dependence
19	and relative humidity interference.

3.5.5.4 Differential Optical Absorption Spectrometry

20	Optical remote sensing methods can provide direct, sensitive, and specific measurements
21	of O_3 over a broad area or open path in contrast with conventional single-point UV
22	monitors. The 1996 O ₃ AQCD (U.S. EPA, 1996a) provided a brief discussion of DOAS
23	for O ₃ measurements and cited references to document the sensitivity (1.5 ppb for a 1-
24	minute averaging time), correlation ($r = 0.89$), and agreement (on the order of 10%) with
25	UV O ₃ monitors (Stevens, 1993). The 2006 O ₃ AQCD (U.S. EPA, 2006b) provided an
26	update on DOAS where a positive interference due to an unidentified absorber was noted
27	(<u>Reisinger, 2000</u>).
28	More recent study of the accuracy of UV absorbance monitors by Williams et al. (2006)
29	compared UV and DOAS measurements at two urban locations. In order to compare the
30	open path measurements and UV, the data sets were averaged to 30-minute periods and
31	only data when the boundary layer was expected to be well mixed (between 10:00 a.m.
32	and 6:00 p.m. CST) were evaluated. The comparisons showed variations of no more
33	than \pm 7% (based on the slope of the linear least squares regression over a concentration
34	range from about 20 to 200 ppb) and good correlation ($R^2 = 0.96$ and 0.98). Lee et al.

1	(2008b) evaluated DOAS and UV O_3 measurements in Korea and found the average
2	DOAS concentration to be 8.6% lower than the UV point measurements with a good
3	correlation ($\mathbf{R}^2 = 0.94$).
4	DOAS has also been used for the measurement of HNO_2 (or HONO). DOAS was
5	compared to chemical point-measurement methods for HONO. Acker et al. (2006)
6	obtained good results when comparing wet chemical and DOAS during well mixed
7	atmospheric conditions (wet chemical = $0.009 + 0.92 \times \text{DOAS}$; r = 0.7). <u>Kleffmann and</u>
8	Wiesen (2008) noted that interferences with the HONO wet chemical methods can affect
9	results from inter-comparison studies if not addressed. In an earlier study, Kleffmann et
10	al. (2006) demonstrated that when the interferences were addressed, excellent agreement
11	with DOAS can be obtained. Stutz et al. (2009) found good agreement (15% or better)
12	between DOAS and a wet chemical method (Mist Chamber/Ion Chromatography) in
13	Houston, TX except generally during mid-day when the chemical method showed a
14	positive bias that may have been related to concentrations of O ₃ . DOAS remains
15	attractive due to its sensitivity, speed of response, and ability to simultaneously measure
16	multiple pollutants; however, further inter-comparisons and interference testing are
17	recommended.

3.5.5.5 Satellite Remote Sensing

18	Satellite observations for O_3 are growing as a resource for many purposes, including
19	model evaluation, assessing emissions reductions, pollutant transport, and air quality
20	management. Satellite remote sensing instruments do not directly measure the
21	composition of the atmosphere. Satellite retrievals are conducted using the solar
22	backscatter or thermal infrared emission spectra and a variety of algorithms. Most
23	satellite measurement systems have been developed for stratospheric measurement of the
24	total O ₃ column. Mathematical techniques have been developed and must be applied to
25	derive information from these systems about tropospheric O ₃ (Tarasick and Slater, 2008;
26	Ziemke et al., 2006). Direct retrieval of global tropospheric O_3 distributions from solar
27	backscattered UV spectra have been reported from OMI and the Global Ozone
28	Monitoring Experiment (GOME) (Liu et al., 2006). Another satellite measurement
29	system, Tropospheric Emission Spectrometer (TES), produces global-scale vertical
30	concentration profiles of tropospheric O ₃ from measurements of thermal infrared
31	emissions. TES has been designed specifically to focus on mapping the global
32	distribution of tropospheric O ₃ extending from the surface to about 10-15 km altitude
33	(<u>Beer, 2006</u>).

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1	In order to improve the understanding of the quality and reliability of the data, satellite-
2	based observations of total column and tropospheric O ₃ have been validated in several
3	studies using a variety of techniques, such as aircraft observations, ozonesondes, CTMs,
4	and ground-based spectroradiometers. Antón et al. (2009) compared satellite data from
5	two different algorithms (OMI-DOAS and OMI-TOMS) with total column O_3 data from
6	ground-based spectroradiometers at five locations. The satellite total column O_3 data
7	underestimated ground-based measurements by less than 3%. Richards et al. (2008)
8	compared TES tropospheric O_3 profiles using airborne differential absorption lidar
9	(DIAL) and found TES to have a 7 ppbv positive bias relative to DIAL throughout the
10	troposphere. Nassar et al. (2008) compared TES O ₃ profiles and ozonesonde coincidences
11	and found a positive bias of 3-10 ppbv for TES. Worden et al. (2007a) also compared
12	TES with ozonesondes and found TES O_3 profiles to be biased high in the upper
13	troposphere (average bias of 16.8 ppbv for mid-latitudes and 9.8 ppbv for the tropics) and
14	biased low in the lower troposphere (average bias of -2.6 ppbv for mid-latitudes and -
15	7.4 ppbv for the tropics). Comparisons of TES and OMI with ozonesondes by Zhang et
16	al. (2010b) showed a mean positive bias if 5.3 ppbv (10%) for TES and 2.8 ppbv (5%) for
17	OMI at 500 hPa. In addition, Zhang et al. (2010b) used a CTM (GEOS-Chem) to
18	determine global differences between TES and OMI. They found differences between
19	TES and OMI were generally \pm 10 ppbv except at northern mid-latitudes in summer and
20	over tropical continents. Satellite observations have also been combined (e.g., OMI and
21	TES) to improve estimates of tropospheric O_3 (Worden et al., 2007b).

3.5.6 Ambient Ozone Network Design

3.5.6.1 Monitor Siting Requirements

22	To monitor compliance with the NAAQS, state and local monitoring agencies operate O_3
23	monitoring sites at various locations depending on the area size (population and
24	geographic characteristics ¹) and typical peak concentrations (expressed in percentages
25	below, or near the O_3 NAAQS). SLAMS make up the ambient air quality monitoring
26	sites that are primarily needed for NAAQS comparisons, but may also serve some other
27	basic monitoring objectives that include: providing air pollution data to the general public
28	in a timely manner; emissions strategy development; and support for air pollution
29	research. SLAMS include National Core (NCore), Photochemical Assessment
30	Monitoring Stations (PAMS), and all other State or locally-operated stations except for
31	the monitors designated as special purpose monitors (SPMs).

¹ Geographic characteristics such as complexity of terrain, topography, land use, etc.

1	The SLAMS minimum monitoring requirements to meet the O ₃ design criteria are
2	specified in 40 CFR Part 58, Appendix D. Although NCore and PAMS are a subset of
3	SLAMS, the monitoring requirements for those networks are separate and discussed
4	below. The minimum number of O ₃ monitors required in a Metropolitan Statistical Area
5	(MSA) ranges from zero for areas with a population of at least 50,000 and under 350,000
6	with no recent history of an O ₃ design value ¹ greater than 85 percent of the NAAQS, to
7	four for areas with a population greater than 10 million and an O_3 design value greater
8	than 85 percent of the NAAQS. Within an O_3 network, at least one site for each MSA, or
9	Combined Statistical Area (CSA) if multiple MSAs are involved, must be designed to
10	record the maximum concentration for that particular metropolitan area. More than one
11	maximum concentration site may be necessary in some areas. The spatial scales for O_3
12	sites are neighborhood, urban and regional.
13	 Neighborhood scale: represents concentrations within some extended area of
14	the city that has relatively uniform land use with dimensions in the 0.5-4.0 km
15	range. The neighborhood and urban scales listed below have the potential to
16	overlap in applications that concern secondary or homogeneously distributed
17	primary air pollutants.
18	 Urban scale: represents concentrations within an area of city-like dimensions,
19	on the order of 4-50 km. Within a city, the geographic placement of sources
20	may result in there being no single site that can be said to represent air quality
21	on an urban scale.
22	 Regional scale: usually defines a rural area of reasonably homogeneous
23	geography without large sources, and extends from tens to hundreds of
24	kilometers.
25	Since O_3 concentrations decrease appreciably in the colder parts of the year in many
26	areas, O ₃ is required to be monitored at SLAMS monitoring sites only during the "ozone
27	season." Table D-3 of 40 CFR Part 58, Appendix D lists the beginning and ending month
28	of the ozone season for each U.S. state or territory. Most operate O ₃ monitors only during
29	the ozone season. Those that operate some or all of their O_3 monitors on a year-round
30	basis include Arizona, California, Hawaii, Louisiana, Nevada, New Mexico, Puerto Rico,
31	Texas, American Samoa, Guam and the Virgin Islands.
32	The total number of SLAMS O3 sites needed to support the basic monitoring objectives
33	includes more sites than the minimum numbers required in 40 CFR Part 58, Appendix D.
34	In 2010, there were 1250 O_3 monitoring sites reporting values to the EPA AQS database

¹ A design value is a statistic that describes the air quality status of a given area relative to the level of the NAAQS. Design values are typically used to classify nonattainment areas, assess progress towards meeting the NAAQS, and develop control strategies. See http://epa.gov/airtrends/values.html (U.S. EPA, 2010a) for guidance on how these values are defined.

1	(Figure 3-21). Monitoring site information for EPA's air quality monitoring networks is
2	available in spreadsheet format (CSV) and keyhole markup language format (KML or
3	KMZ) that is compatible with Google Earth [™] and other software applications on the
4	AirExplorer website (U.S. EPA, 2011d). States may operate O_3 monitors in non-urban or
5	rural areas to meet other objectives (e.g., support for research studies of atmospheric
6	chemistry or ecosystem impacts). These monitors are often identified as SPMs and can be
7	operated up to 24 months without being considered in NAAQS compliance
8	determinations. The current monitor and probe siting requirements have an urban focus
9	and do not address the siting for SPMs or monitors in non-urban, rural areas to support
10	ecosystem impacts and the secondary standards.

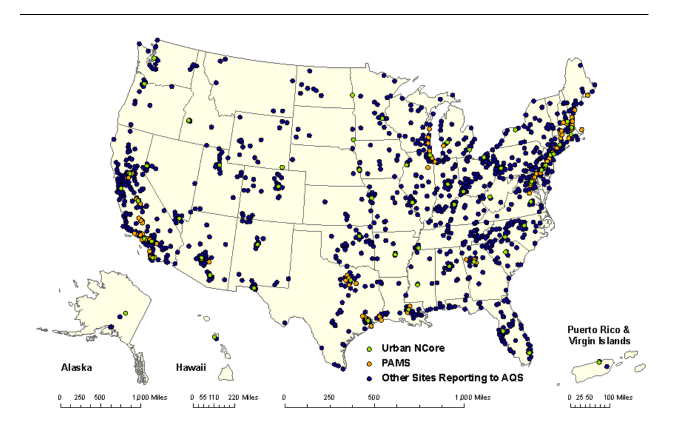


Figure 3-21 U.S. ozone sites reporting data to AQS in 2010.

11	NCore is a new multipollutant monitoring network implemented to meet multiple
12	monitoring objectives. Those objectives include: timely reporting of data to the public
13	through AirNow (U.S. EPA, 2011a); support for the development of emission reduction
14	strategies; tracking long-term trends of criteria pollutants and precursors; support to
15	ongoing reviews of the NAAQS and NAAQS compliance; model evaluation; support for

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1	scientific research studies; and support for ecosystem assessments. Each state is required
2	to operate at least one NCore site. The NCore monitoring network began January 1, 2011
3	at about 80 stations (about 60 urban and 20 rural sites). NCore has leveraged the use of
4	sites in existing networks; for example, some IMPROVE sites also serve as rural NCore
5	sites. In addition to O_3 , other components including CO, NO_X , NO_Y , SO ₂ , and basic
6	meteorology are also measured at NCore sites. The spatial scale for urban NCore stations
7	is urban or neighborhood; however, a middle-scale ¹ site may be acceptable in cases
8	where the site can represent many such locations throughout a metropolitan area. Rural
9	NCore sites are located at a regional or larger scale, away from any large local emission
10	sources so that they represent ambient concentrations over an extensive area. Ozone
11	monitors at NCore sites are operated year round.
12	PAMS provides more comprehensive data on O_3 in areas classified as serious, severe, or
13	extreme nonattainment for O ₃ . In addition to O ₃ , PAMS provides data for NO _X , NO _Y ,
14	VOCs, carbonyls, and meteorology. The PAMS network design criteria are based on
15	locations relative to O ₃ precursor source areas and predominant wind directions
16	associated with high O ₃ concentrations. The overall network design is location specific
17	and geared toward enabling characterization of precursor emission sources in the area, O ₃
18	transport, and photochemical processes related to O ₃ nonattainment. Minimum
19	monitoring for O ₃ and its precursors is required annually during the months of June, July,
20	and August when peak O ₃ concentrations are expected. In 2006, the EPA reduced the
21	minimum PAMS monitoring requirements (71 FR 61236). There were a total of 92
22	PAMS sites reporting values to the AQS data base in 2010.
23	CASTNET is a regional monitoring network established to assess trends in acidic
24	deposition due to emission reduction regulations. CASTNET also provides concentration
25	measurements of air pollutants involved in acidic deposition, such as sulfate and nitrate,
26	in addition to the measurement of O ₃ . CASTNET O ₃ monitors operate year round and are
27	primarily located in rural areas. In 2010, there were 80 CASTNET sites located in, or
28	near, rural areas. As part of CASTNET, the National Park Service (NPS) operates 23
29	sites located in national parks and other Class-i areas. Ozone data collected at the 23 NPS

sites is compliant with the SLAMS QA requirements in 40 CFR Part 58, Appendix A.

Ozone measurements at the remaining CASTNET sites were not collected with the QA

requirements for SLAMS outlined in 40 CFR Part 58, Appendix A, and therefore, these

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O₃ data cannot be used for NAAQS compliance purposes. The SLAMS QA requirements and procedures are currently being implemented at the remaining sites. The NPS also operates a Portable Ozone Monitoring Systems (POMS) network. The

POMS couples the small, low-power O_3 monitor with a data logger, meteorological

¹ Middle scale defines an area up to several city blocks in size with dimensions ranging from about 100 to 500 m.

- 1 measurements, and solar power in a self contained system for monitoring in remote 2 locations. Typical uses for the POMS data include research projects, survey monitoring, 3 and assessments of spatial O₃ distribution. The portable O₃ monitor in use by the NPS 4 was recently designated as an equivalent method for O₃ (75 FR 22126). Seventeen NPS 5 POMS monitors were operating in 2010 (NPS, 2011). A map of the rural NCore sites, 6 along with the CASTNET, and the NPS POMS sites are shown in Figure 3-22. As can be 7 seen from Figure 3-21 and Figure 3-22, vast rural areas of the country still exist without 8 any monitor coverage. Monitoring opportunities exist in these areas where relatively few 9 and easily characterized precursor sources dominate and could be used to improve 10 understanding of O₃ formation.
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- 12

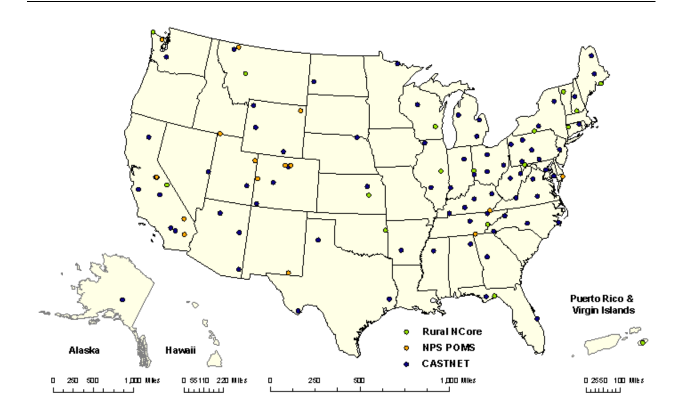


Figure 3-22 U.S. Rural NCore, CASTNET and NPS POMS ozone sites in 2010.

3.5.6.2 Probe/Inlet Siting Requirements

1	Probe and monitoring path siting criteria for ambient air quality monitoring are contained
2	in 40 CFR Part 58, Appendix E. For O_3 , the probe must be located between 2 and
3	15 meters above ground level and be at least 1 meter away (both in the horizontal and
4	vertical directions) from any supporting structure, walls, etc. If it is located on the side of
5	a building, it must be located on the windward side, relative to prevailing wind direction
6	during the season of highest potential O ₃ concentration. Ozone monitors are placed to
7	determine air quality in larger areas (neighborhood, urban, or regional scales) and
8	therefore, placement of the monitor probe should not be near local, minor sources of NO,
9	O ₃ -scavenging hydrocarbons, or O ₃ precursors. The probe or inlet must have unrestricted
10	air flow in an arc of at least 180 degrees and be located away from any building or
11	obstacle at a distance of at least twice the height of the obstacle. The arc of unrestricted
12	air flow must include the predominant wind direction for the season of greatest O_3
13	concentrations. Some exceptions can be made for measurements taken in street canyons
14	or sites where obstruction by buildings or other structures is unavoidable. The scavenging
15	effect of trees on O_3 is greater than other pollutants and the probe/inlet must be located at
16	least 10 meters from the tree drip line to minimize interference with normal air flow.
17	When siting O ₃ monitors near roadways, it is important to minimize the destructive
18	interferences from sources of NO, since NO reacts readily with O ₃ . For siting
19	neighborhood and urban scale O ₃ monitors, guidance on the minimum distance from the
20	edge of the nearest traffic lane is based on roadway average daily traffic count (40 CFR
21	Part 58, Appendix E, Table E-1). The minimum distance from roadways is 10 meters
22	(average daily traffic count \leq 1,000) and increases to a maximum distance of 250 meters
23	(average daily traffic count \geq 110,000).

3.6 Ambient Concentrations

24	This section investigates spatiotemporal variability in ambient O3 concentrations and
25	associations between O_3 and copollutants. To set the stage for the rest of the section,
26	common O_3 measurement units, metrics, and averaging times are described and
27	compared in Section $3.6.1$. Spatial variability is covered in Section $3.6.2$ and is divided
28	into urban-focused variability and rural-focused variability. Urban-focused variability is
29	organized by scale, extending from national-scale down to neighborhood-scale and the
30	near-road environment. Rural-focused variability is organized by region and includes
31	observations of ground-level vertical O3 gradients where available. Temporal variability
32	is covered in Section $3.6.3$ and is organized by time, extending from multiyear trends
33	down to hourly (diel) variability. In many instances, spatial and temporal variability are

1	inseparable (e.g., seasonal dependence to spatial variability), resulting in some overlap
2	between Section $3.6.2$ and Section $3.6.3$. Finally, Section $3.6.4$ covers associations
3	between O_3 and co-pollutants including CO, SO ₂ , NO ₂ , PM _{2.5} and PM ₁₀ .
4	As noted in the 2006 O_3 AQCD (U.S. EPA, 2006b), O_3 is the only photochemical oxidant
5	other than nitrogen dioxide (NO ₂) that is routinely monitored and for which a
6	comprehensive database exists. Data for other photochemical oxidants (e.g., PAN, H ₂ O ₂ ,
7	etc.) typically have been obtained only as part of special field studies. Consequently, no
8	data on nationwide patterns of occurrence are available for these other oxidants; nor are
9	extensive data available on the relationships of concentrations and patterns of these
10	oxidants to those of O_3 . As a result, this section focuses solely on O_3 , the NAAQS
11	indicator for photochemical oxidants. The majority of ambient O3 data reported in this
12	section were obtained from AQS, EPA's repository for detailed, hourly data that has been
13	subject to EPA quality control and assurance procedures (the AQS network was
14	described in Section 3.5).

3.6.1 Measurement Units, Metrics, and Averaging Times

15	Several approaches are commonly used for reporting O_3 data. In atmospheric sciences
16	and epidemiology, O_3 is frequently reported as a concentration, expressed as a volume-to-
17	volume mixing ratio, commonly measured in ppm or ppb. In human exposure, O_3 is
18	frequently reported as a cumulative exposure, expressed as a mixing ratio times time
19	(e.g., ppm-h). In ecology, cumulative exposure indicators are frequently used that extend
20	over longer time periods, such as growing season or year. This section focuses on
21	ambient concentrations derived primarily from hourly average O ₃ measurements and
22	concentrations are reported in ppb wherever possible. Further details on human and
23	ecological exposure metrics can be found in Chapter $\underline{4}$ and Chapter $\underline{9}$, respectively.
24	As discussed in Section 3.5 , most continuous O ₃ monitors report hourly average
25	concentrations to AQS with a required precision of 10 ppb and LDL of 10 ppb (see
26	Table 3-4). This data can be used as reported (1-h avg), or further summarized in one of
27	several ways to focus on important aspects of the data while simultaneously reducing the
28	volume of information. Three common daily reporting metrics include: (1) the average of
29	the hourly observations over a 24-h period (24-h avg); (2) the maximum hourly
30	observation occurring in a 24-h period (1-h daily max); and (3) the maximum 8-h running
31	average of the hourly observations occurring in a 24-h period (8-h daily max) ¹ .
32	Throughout this ISA and the literature, O_3 concentrations are reported using different

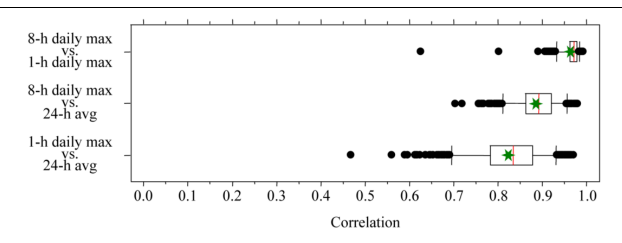
¹ For O_3 regulatory monitoring purposes, the 8-h daily max is calculated by first generating all 8-h running averages and storing these averages hourly by the first hour in the 8-h period. The 8-h daily max is then set equal to the maximum of the 24 individual 8-h avg occurring in a given day.

averaging times as appropriate, making it important to recognize the differences between these metrics.

3 Nation-wide, year-round 1-h avg O₃ data reported to AQS from 2007-2009 was used to 4 compare these different daily metrics. Correlations between the 24-h avg, 1-h daily max 5 and 8-h daily max metrics were generated on a site-by-site basis. Figure 3-23 contains 6 box plots of the distribution in correlations from all sites. The top comparison in 7 Figure 3-23 is between 8-h daily max and 1-h daily max O_3 . Not surprisingly, these two 8 metrics are very highly correlated (median r = 0.97, IQR = 0.96-0.98). There are a couple 9 outlying sites, with correlations between these two metrics as low as 0.63, but 95% of 10 sites have correlations above 0.93. The middle comparison in Figure 3-23 is between 8-h 11 daily max and 24-h avg O_3 . For these metrics, the distribution in correlations is shifted 12 down and broadened out (median r = 0.89, IQR = 0.86-0.92). Finally, the bottom 13 comparison in Figure 3-23 is between 1-h daily max and 24-h avg O_3 . Again, for these 14 metrics the distribution in correlations is shifted down and broadened out relative to the 15 other two comparisons (median r = 0.83, IQR = 0.78-0.88). The correlation between the 16 two daily-maximum metrics (1-h daily max and 8-h daily max) are quite high for most 17 sites, but correlations between the daily maximum metrics and the daily average metric 18 (24-h avg) are lower. This illustrates the influence of the overnight period on the 24-h avg 19 O₃ concentration. In contrast, the 1-h daily max and 8-h daily max are more indicative of 20 the daytime, higher O_3 periods. The correlation between these metrics, however, can be 21 very site-specific, as is evident from the broad range in correlations in Figure 3-23 for all 22 three comparisons. Therefore, understanding which O_3 metric is being used in a given 23 study is very important since they capture different aspects of O_3 temporal variability.

1

2



Note: Shown are the median (red line), mean (green star), inner-quartile range (box), 5th and 95th percentiles (whiskers), and extremes (black circles).

Figure 3-23 Distribution in nation-wide year-round site-level correlations between daily ozone metrics including 24-h avg, 1-h daily max and 8-h daily max using AQS data, 2007-2009.

1	The median 1-h daily max, 8-h daily max, and 24-h avg O_3 concentrations across all sites
2	included in the 3-year nation-wide data set were 44, 40, and 29 ppb, respectively.
3	Representing the upper end of the distribution, the 99th percentiles of these same metrics
4	across all sites were 94, 80, and 60 ppb, respectively. While the ratio of these metrics will
5	vary by location, typically the 1-h daily max will be the highest value representing peak
6	concentrations and the 24-h avg will be considerably lower representing daily average
7	concentrations incorporating the overnight period. The 8-h daily max typically represents
8	the higher mid-day concentrations and will generally lie somewhere between the other
9	two metrics ¹ .

¹ The 8-h daily max is not strictly limited to lie between the 1-h daily max and the 24-h avg since the 8-h averaging period used to calculate the 8-h daily max can extend into the morning hours of the subsequent day. However, the 8-h daily max typically incorporates the middle of the day when O_3 concentrations are at their highest, resulting in an 8-h daily max somewhere between the 1-h daily max and the 24-h avg calculated for that day.

3.6.2 Spatial Variability

3.6.2.1 Urban-Focused Variability

National-Scale Variability

1	AQS contains a large depository of national O ₃ data collected to meet the monitoring
2	objectives described in Section $3.5.6$. In many areas, O ₃ concentrations decrease
3	appreciably during months with lower temperatures and decreased sunlight. As a result,
4	year-round O ₃ monitoring is only required in certain areas. Table D-3 of 40 CFR Part 58,
5	Appendix D lists the beginning and ending month of the ozone season (defined in
6	Section 3.5.6.1) by geographic area and Figure 3-24 illustrates these time periods on a
7	monitor-by-monitor basis. Monitoring is optional outside the ozone season and many
8	states elect to operate their monitors year-round or for time periods outside what is
9	strictly mandated.

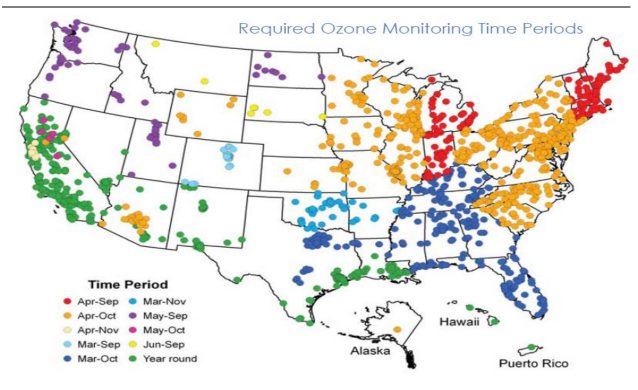




Figure 3-24 Required ozone monitoring time periods (ozone season) identified by monitoring site.

1	Hourly FRM and FEM O ₃ data reported to AQS for the period 2007-2009 were used to
2	investigate national-scale spatial variability in O ₃ concentrations. Given the variability in
3	O ₃ monitoring time periods available in AQS as a result of the regionally-varying ozone
4	seasons, the analyses in this section were based on two distinct data sets:
5	• a year-round data set: data only from monitors reporting year-round;
6	• a warm-season data set: data from all monitors reporting May through
7	September.
8	The warm-season data set was used to capture the majority of ozone season data while
9	providing a consistent time-frame for comparison across states. All available monitoring
10	data including data from year-round monitors was included in the warm-season data set
11	after removing observations outside the 5-month window. Data were retrieved from AQS
12	on February 25, 2011 for these two data sets, and all validated data was included
13	regardless of flags or regional concurrence ¹ . A summary of the two O ₃ data sets including
14	the applied completeness criteria is provided in Table 3-5. Figure 3-25 and Figure 3-26
15	show the location of the 457 year-round and 1,064 warm-season monitors meeting the
16	completeness criteria for all three years (2007-2009).

Table 3-5Summary of ozone data sets originating from AQS

	Year-Round Data Set	Warm-Season Data Set					
Years	2007-2009	2007-2009					
Months	January - December (12 mo)	May - September (5 mo)					
Completeness Criteria	75% of hours in a day	75% of hours in a day					
	75% of days in a calendar quarter	75% of days between May - September					
	All 4 quarters per year						
Number of monitors meeting completeness criteria	618 containing at least one valid year in 2007-2009	1,267 containing at least one valid year in 2007-2009					
	550 containing at least two valid years in 2007-2009	1,169 containing at least two valid years in 2007-2009					
	457 containing all three valid years in 2007-2009	1,064 containing all three valid years in 2007-2009					

¹ Concentrations that might have been affected by exceptional events (and contribute to a violation of the NAAQS) can be flagged in the Air Quality System (AQS) by the reporting organization. Exceptional events are defined as unusual or naturally occurring events that can affect air quality but are not reasonably controllable using techniques that tribal, state or local air agencies may implement in order to attain and maintain the National Ambient Air Quality Standards (NAAQS). The corresponding EPA Regional Office is responsible for reviewing the data and evidence of the event, and deciding whether to concur with the flag. Flagged data that has been concurred by the Regional office is typically excluded for regulatory purposes.

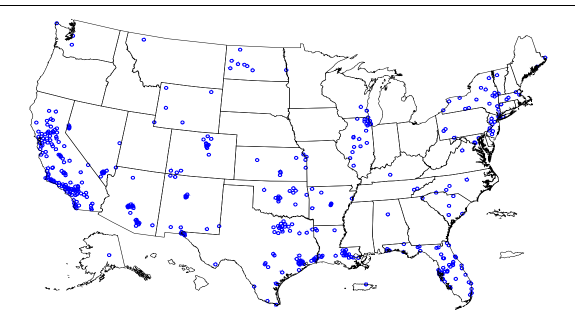


Figure 3-25 Location of the 457 ozone monitors meeting the year-round data set completeness criterion for all 3 years between 2007 and 2009.

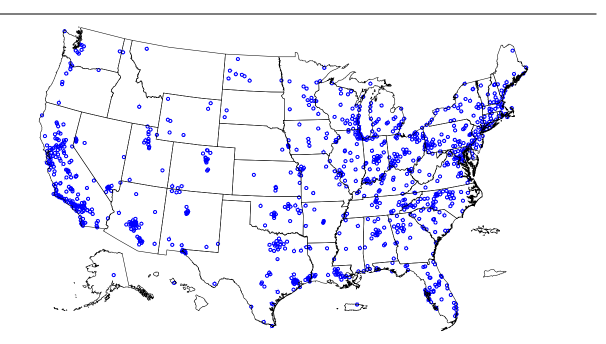


Figure 3-26 Location of the 1,064 ozone monitors meeting the warm-season data set completeness criteria for all 3 years between 2007 and 2009.

1	Tabulated statistics generated from the year-round and warm-season data sets are
2	included in <u>Table 3-6</u> and
3	Table 3-7, respectively. This information was used to compare (1) the year-round and
4	warm-season data sets; (2) the O_3 distribution variability across years (2005-2009); and
5	(3) four different averaging times (1-h avg, 24-h avg, 1-h daily max, and 8-h daily max).
6	Summary statistics for 2005 and 2006 were added to these tables in order to gain a
7	broader view of year-to-year variability, but the year-round and warm-season data sets
8	used for analyses in the rest of this section are limited to 2007-2009 as described above
9	and in Table 3-5. The 8-h daily max pooled by site was also included in these tables to
10	show the distribution of the annual and 3-year (2007-2009) site-averages of the 8-h daily
10	max statistic.
11	max statistic.
12	The year-round data set includes data from roughly half the number of monitors as the
13	warm-season data set and a larger fraction of the year-round monitors are located in the
14	southern half of the U.S. due to extended monitoring requirements in these areas. Despite
15	these differences, the mean, SD and percentiles of the nation-wide O ₃ concentrations
16	were quite similar for the year-round data presented in Table 3-6 and the warm-season
17	data presented in
18	Table 3-7. In both data sets, there was very little variability across years in the central
19	statistics; for example, the median 1-h avg concentrations between 2005 and 2009 ranged
20	from 28 to 29 ppb for the year-round data and from 29 to 30 ppb for the warm-season
21	data. The 8-h daily max showed similar uniformity in median across the five years, with
22	concentrations ranging from 39 to 41 ppb for the year-round data and from 40 to 43 for
23	the warm-season data. The upper percentiles (95th and above) showed a general
24	downward trend from 2005 to 2009 in both nation-wide data sets. For example, the 99th
25	percentile of the 8-h daily max observed in the warm-season data dropped from 85 ppb in
26	2005 to 75 ppb in 2009. Trends in O_3 concentrations investigated over a longer time
27	period are included in Section $3.6.3.1$.
_ ·	r

Time Period	N Monitors	N Obs	Mean	SD	Min	1	5	10	25	50	75	90	95	98	99	Max	Max Site ID ^b
1-h avg ^a																	
2005	499	4,284,219	29	18	2	2	2	2	15	28	41	53	61	71	78	182	060710005
2006	532	4,543,205	30	18	2	2	2	5	16	29	42	54	61	71	78	175	060370016
2007	522	4,547,280	29	18	2	2	2	5	16	29	41	52	60	68	75	237	450790021
2008	520	4,470,065	30	17	2	2	2	6	17	29	41	52	59	67	74	222	450210002
2009	551	4,716,962	29	16	2	2	2	6	17	29	40	50	56	64	70	188	720770001
2007-2009	599	13,734,307	29	17	2	2	2	6	17	29	40	51	58	67	73	237	450790021
24-h avg ^a																	
2005	504	183,815	29	13	2	4	9	13	20	28	37	46	51	57	61	103	060719002
2006	536	194,884	30	13	2	5	10	14	21	29	38	47	52	58	62	102	061070009
2007	531	194,873	29	12	2	5	11	14	20	29	37	45	50	56	60	96	060651016
2008	528	191,875	30	12	2	5	11	14	21	29	38	46	50	56	61	98	060710005
2009	556	202,142	29	11	2	6	11	14	21	28	37	44	48	53	57	95	060710005
2007-2009	611	588,890	29	12	2	5	11	14	21	29	37	45	49	55	60	98	060710005
1-h daily ma	ax ^a																
2005	504	183,815	48	18	2	11	21	26	35	46	58	71	80	91	100	182	060710005
2006	536	194,884	48	18	2	13	23	28	36	46	58	71	80	91	100	175	060370016
2007	531	194,873	47	17	2	14	23	28	36	45	57	69	77	87	94	237	450790021
2008	528	191,875	47	17	2	14	23	27	35	45	56	67	76	87	96	222	450210002
2009	556	202,142	45	15	2	14	22	27	35	44	54	64	72	83	91	188	720770001
2007-2009	611	588,890	46	16	2	14	23	27	35	44	55	67	75	86	94	237	450790021
8-h daily ma	ax ^a																
2005	504	183,279	42	16	2	7	16	21	30	40	52	63	70	78	84	145	060710005
2006	536	194,285	42	16	2	9	18	23	31	41	52	63	70	79	85	142	060710005
2007	528	194,266	41	15	2	10	19	23	31	40	51	61	68	75	81	137	060710005
2008	528	191,283	41	15	2	11	19	23	31	40	51	60	66	75	82	172	450210002
2009	556	201,536	40	14	2	11	18	23	30	39	49	57	63	71	77	128	060712002
2007-2009	608	587,085	41	15	2	10	19	23	31	40	50	60	66	74	80	172	450210002
8-h daily ma	ax (pooled b	y site) ^a															
2005	508	508	42	6	23	27	32	34	38	42	45	48	51	53	55	61	060710005
2006	538	538	42	6	12	28	31	34	38	43	46	50	52	54	55	61	060719002
2007	538	538	41	6	17	27	31	34	38	41	45	49	51	54	55	63	060719002
2008	529	529	41	6	20	28	31	34	37	40	45	50	52	55	57	61	060719002
2009	558	558	40	6	20	26	30	33	36	39	44	48	50	53	54	60	060719002
2007-2009	457	457	41	6	19	29	32	34	38	40	45	49	51	54	55	61	060719002

Table 3-6Nationwide distributions of ozone concentrations (ppb) from the
year-round data set.

^aIncludes all validated data regardless of flags or regional concurrence and therefore may differ from data used for regulatory purposes

^bAQS Site ID corresponding to the observation in the Max column

Time Period	N Monitors	N Obs	Mean	SD	Min	1	5	10	25	50	75	90	95	98	99	Max	Max Site ID ^b
1-h avg ^ª																	
2005	1,023	7,455,018	30	19	2	2	2	5	16	29	43	55	64	73	79	182	060710005
2006	1,036	7,590,796	31	18	2	2	2	6	17	30	43	55	62	71	77	175	060370016
2007	1,021	7,711,463	31	18	2	2	2	6	18	30	43	55	63	71	77	237	450790021
2008	1,034	7,701,597	31	17	2	2	2	7	18	30	42	53	60	68	74	222	450210002
2009	1,029	7,835,074	29	16	2	2	2	7	17	29	40	50	56	63	69	259	311090016
2007-2009	1,103	23,248,134	30	17	2	2	2	7	18	30	42	53	60	68	74	259	311090016
24-h avg ^a																	
2005	1,103	319,410	30	12	2	5	10	14	22	30	39	46	51	57	61	103	060719002
2006	1,110	324,993	31	12	2	6	12	15	22	30	39	47	52	58	61	102	061070009
2007	1,100	330,197	31	12	2	6	12	16	23	31	39	47	51	57	61	96	060651016
2008	1,120	329,918	31	12	2	6	12	16	22	30	38	46	50	56	60	98	060710005
2009	1,141	335,669	29	11	2	6	12	15	21	29	37	44	48	53	56	95	060710005
2007-2009	1,197	995,784	30	12	2	6	12	16	22	30	38	45	50	55	59	98	060710005
1-h daily max ^a	1																
2005	1,103	319,410	50	18	2	12	23	28	38	49	61	74	81	91	99	182	060710005
2006	1,110	324,993	50	17	2	15	25	29	38	48	60	72	80	90	98	175	060370016
2007	1,100	330,197	50	17	2	16	25	30	38	48	60	72	80	88	95	237	450790021
2008	1,120	329,918	48	16	2	16	25	29	37	47	58	69	76	86	93	222	450210002
2009	1,141	335,669	46	15	2	15	23	28	36	45	54	64	71	80	87	259	311090016
2007-2009	1,197	995,784	48	16	2	16	24	29	37	47	58	68	76	85	93	259	311090016
8-h daily max ^a	1																
2005	1,104	318,771	44	16	2	9	18	23	32	43	55	66	72	79	85	145	060710005
2006	1,112	324,327	44	16	2	11	20	25	33	43	54	64	70	78	84	142	060710005
2007	1,097	329,482	44	15	2	12	20	25	33	43	54	65	71	78	82	137	060710005
2008	1,120	329,223	43	15	2	12	20	25	33	42	52	61	67	74	80	172	450210002
2009	1,141	334,972	40	13	2	12	19	24	31	40	49	57	63	69	75	128	060712002
2007-2009	1,194	993,677	42	15	2	12	20	24	32	42	52	61	67	75	80	172	450210002
8-h daily max	(pooled by s	site) ^a															
2005	1,141	1,141	45	6	14	28	34	36	41	46	49	52	54	56	57	61	040139508
2006	1,152	1,152	44	6	12	29	34	37	41	45	48	51	54	58	59	65	060170020
2007	1,164	1,164	45	7	17	28	34	36	40	45	50	54	56	58	59	64	471550102
2008	1,163	1,163	43	6	20	29	33	36	39	44	48	50	53	56	58	61	060719002
2009	1,173	1,173	41	5	20	28	32	35	38	41	44	47	50	53	55	63	060651016
2007-2009	1,064	1,064	43	6	19	29	34	36	39	43	47	50	52	55	57	61	060719002

Table 3-7Nationwide distributions of ozone concentrations (ppb) from the
warm-season data set.

^aIncludes all validated data regardless of flags or regional concurrence and therefore may differ from data used for regulatory purposes.

^bAQS Site ID corresponding to the observation in the Max column.

1Given the strong diurnal pattern in O3 concentrations, the selection of averaging time has2a substantial effect on the magnitude of concentration reporting. The nation-wide median31-h avg, 24-h avg, 1-h daily max, and 8-h daily max concentrations for the year-round4data set in 2009 were 29, 28, 44 and 39 ppb, respectively. The median concentrations for

1	the warm-season data set in 2009 were: 29, 29, 45 and 40 ppb, respectively. The 1-h avg
2	and 24-h avg both include the lowest concentrations typically observed in the overnight
3	period which lowers their values relative to the daily maximum statistics.
4	A strong seasonal pattern in O ₃ concentrations can also be seen in the year-round data.
5	<u>Table 3-8</u> shows the 8-h daily max stratified by season, with the seasons defined as:
6	 winter: December-February;
7	 spring: March-May;
8	 summer: June-August; and
9	 fall: September-November.
10	In addition, warm-season (May-Sept) and cold-season (Oct-Apr) stratifications of the
11	year-round data set are included in the table for comparison with the four seasonal
12	stratifications. Substantial seasonal variability in the 8-h daily max concentration for the
13	period 2007-2009 was evident with lower concentrations present in fall
14	(median = 36 ppb) and winter (median = 32 ppb) and higher concentrations in spring
15	(median = 47 ppb) and summer (median = 46 ppb). The seasonal differences were even
16	more pronounced in the upper percentiles. For example, the 99th percentile in the 8-h
17	daily max over the 2007-09 time period ranged from 52 ppb in winter to 90 ppb in
18	summer. The distribution in 8-h daily max O ₃ during the warm-season (as defined above)
19	and during summer were very similar, which is not surprising given their close overlap in
20	months. The distribution during the cold-season (as defined above) is shifted toward
21	higher 8-h daily max O ₃ concentrations compared with the distribution during winter.
22	This is a result of including the four transition months (Oct, Nov, Mar and Apr) in the
23	cold-season when high O ₃ concentrations can occur. Further investigation of temporal
24	variability including multiyear trends and diel behavior is included in Section <u>3.6.3</u> .

Table 3-8Seasonally stratified distributions of 8-h daily max ozone
concentrations (ppb) from the year-round data set (2007-2009).

Time Period	N Monitors	N Obs	Mean	SD	Min	1	5	10	25	50	75	90	95	98	99	Max	Max Site ID ^b
8-h daily max (2007-2009) ^a																
Year-round	608	587,085	41	15	2	10	19	23	31	40	50	60	66	74	80	172	450210002
8-h daily max b	y season (2	2007-2009)) ^a														
Winter (Dec-Feb)	608	143,855	31	10	2	6	14	18	25	32	38	43	46	49	52	172	450210002
Spring (Mar-May)	612	148,409	47	12	2	20	28	33	40	47	55	62	67	72	77	118	060370016
Summer (Jun-Aug)	613	148,280	47	16	2	16	22	26	35	46	57	67	75	84	90	137	060710005
Fall (Sep-Nov)	608	146,541	37	13	2	10	17	21	28	36	45	54	61	68	75	116	060370016
Warm-season (May-Sep)	616	246,233	47	16	2	16	22	27	35	46	57	66	73	81	87	137	060710005
Cold-season (Oct-Apr)	608	340,852	36	12	2	8	16	21	28	36	44	52	57	63	67	172	450210002

^aIncludes all validated data regardless of flags or regional concurrence and therefore may differ from data used for regulatory purposes.

^bAQS Site ID corresponding to the observation in the Max column.

1	A national picture of AQS O ₃ concentrations was generated from the year-round and
2	warm-season data sets by aggregating the 8-h daily max observations by U.S. county. For
3	this purpose, the 8-h daily max concentrations at each site were averaged over one or
4	more calendar years and then the highest site in each county was selected for that county.
5	Figure 3-27 contains the county-scale 8-h daily max O_3 concentrations from the year-
6	round data set for 2007-2009 (top map) with seasonal stratification (bottom four maps).
7	Figure 3-28 contains the county-scale 8-h daily max O_3 concentrations from the warm-
8	season data set for 2007-2009 (top map) along with individual maps for each calendar
9	year between 2007 and 2009 (bottom three maps). These maps are meant to illustrate the
10	general national-scale distribution in long-term average 8-h daily max O3 concentrations
11	and are not representative of O_3 concentrations at all locations or times within the
12	counties shown; considerable spatial variability can exist within a county. This is
13	particularly important in the West where counties are larger on average than in the East.
14	These maps are limited by monitor availability, resulting in the majority of U.S. counties
15	not having available data (the white regions in Figure 3-27 and Figure 3-28).

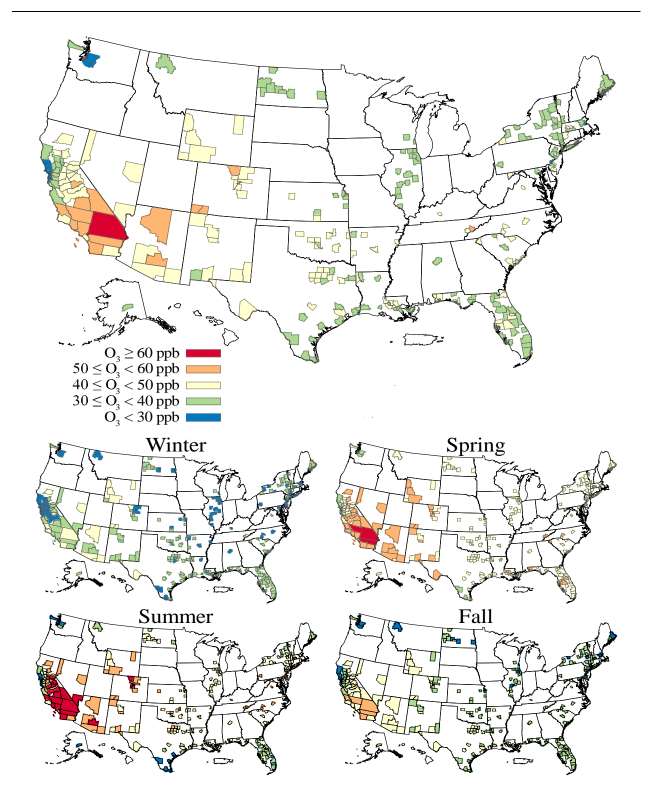


Figure 3-27 Highest monitor (by county) 3-year avg (2007-2009) of the 8-h daily max ozone concentration based on the year-round data set (top map) with seasonal stratification (bottom 4 maps).

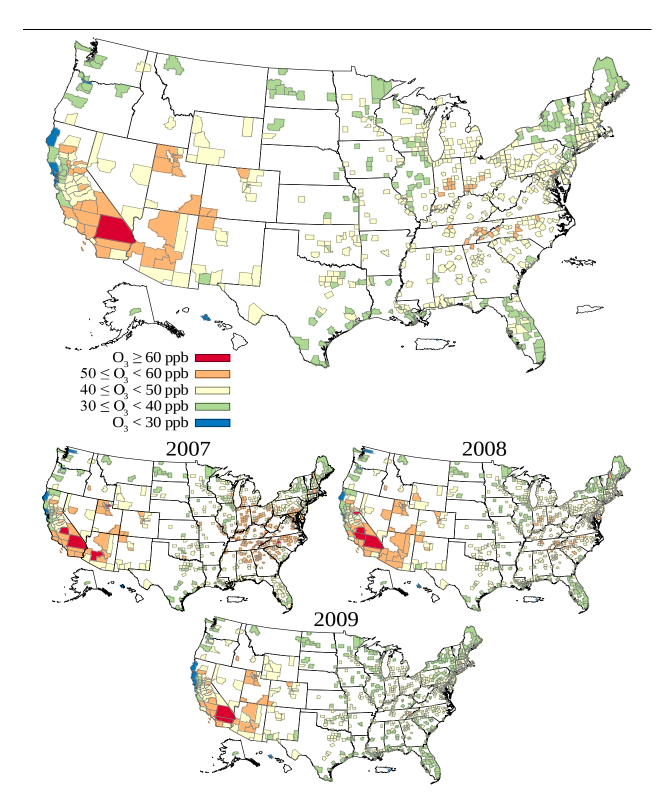


Figure 3-28 Highest monitor (by county) 3-year avg (2007-2009) of the 8-h daily max ozone concentration based on the warm-season data set (top map) with annual stratification (bottom 3 maps).

1	As shown in the top county-scale map generated from the 2007-2009 year-round data set
2	in Figure 3-27, the highest 3-year avg 8-h daily max O_3 concentrations (\geq 50 ppb) occur
3	in counties in central and southern California, Arizona, Colorado and high elevation
4	counties in Tennessee. The highest year-round average concentration of 61 ppb over this
5	period comes from Site #060719002 located at an elevation of 1,244 meters in Joshua
6	Tree National Monument, San Bernardino County, CA. The lowest 3-year avg 8-h daily
7	max O ₃ concentrations (<30 ppb) occur in Pacific Coast counties in northern California
8	and Washington as well as in two northeastern counties in Pennsylvania and
9	Massachusetts. The seasonally-stratified county-scale maps in Figure 3-28 reinforce the
10	strong seasonality in 8-h daily max O_3 concentrations shown in <u>Table 3-8</u> . The highest
11	wintertime concentrations (\geq 40 ppb) occur in the West with the highest 3-year
12	wintertime avg of 46 ppb calculated for Site #080690007 located at an elevation of
13	2,743 meters near Rocky Mountain National Park, Larimer County, CO. In spring and
14	summer, the concentrations increase considerably across all counties, with the highest
15	concentrations (\geq 60 ppb) occurring during the summer in 15 counties in California, 3
16	counties in Colorado and 1 county each in Nevada and Arizona. Many counties in rural
17	Wyoming, Montana, North Dakota, Maine, and along the Gulf Coast peak in the spring
18	instead of the summer. In the fall, 8-h daily max O_3 concentrations drop back down
19	below their spring and summer concentrations.
20	The top county-scale map in Figure 3-28 based on the 2007-2009 warm-season data set
21	looks similar to the corresponding map in Figure 3-27 based on the year-round data set.
22	The warm-season map, however, incorporates approximately twice as many monitors
23	across the U.S., providing more spatial coverage. Several counties in Utah, New Mexico,
24	Indiana, Ohio, Maryland, North Carolina, and Georgia in addition to California, Arizona,
25	Colorado and Tennessee identified above have 3-year avg (2007-2009) 8-h daily max O_3
26	concentrations \geq 50 ppb based on the warm-season data set. The individual yearly
27	average county-maximum 8-h daily max O ₃ concentrations in the lower half of
28	Figure 3-27 show a general decrease in most counties from 2007 to 2009. The number of
29	counties containing a monitor reporting an annual average 8-h daily max O_3
30	concentration above 50 ppb dropped from 230 counties in 2007 to 30 counties in 2009.
31	This is consistent with the general decrease across these years shown in <u>Table 3-6</u> and
32	Table 3-7 for the upper percentiles of the 8-h daily max O_3 concentration.

Urban-Scale Variability

33Statistical analysis of the human health effects of airborne pollutants based on aggregate34population time-series data have often relied on ambient concentrations of pollutants35measured at one or more central monitoring sites in a given metropolitan area. The

1	validity of relying on central monitoring sites is strongly dependent on the spatial
2	variability in concentrations within a given metropolitan area. To investigate urban-scale
3	variability, 20 focus cities were selected for closer analysis of O ₃ concentration
4	variability; these cities are listed in <u>Table 3-9</u> and were selected based on their
5	importance in O ₃ epidemiology studies and on their geographic distribution across the
6	U.S. In order to provide a well-defined boundary around each city, the combined
7	statistical area (CSA) encompassing each city was used. If the city was not within a CSA,
8	the smaller core-based statistical area (CBSA) was selected. The CSAs/CBSAs are
9	defined by the U.S. Census Bureau $(2011)^1$ and have been used to establish analysis
10	regions around cities in previous ISAs for particulate matter (U.S. EPA, 2009d) and
11	carbon monoxide (U.S. EPA, 2010c).

¹A CBSA represents a county-based region surrounding an urban center of at least 10,000 people determined using 2000 census data and replaces the older Metropolitan Statistical Area (MSA) definition from 1990. The CSA represents an aggregate of adjacent CBSAs tied by specific commuting behaviors. The broader CSA definition was used when selecting monitors for the cities listed above with the exception of Phoenix and San Antonio, which are not contained within a CSA. Therefore, the smaller CBSA definition was used for these metropolitan areas.

Focus City	Short Name	CSA/CBSA Name ^a	Year-Round O ₃ Monitoring Sites ^b	Warm-Season O ₃ Monitoring Sites ^c	Included in Prior ISAs ^d	
Atlanta, GA	tlanta, GA Atlanta CSA Atlanta-Sandy Springs-Gainesville		0	11	CO, PM, SO _X , NO _X	
Baltimore, MD	Baltimore CSA	Washington- Baltimore-northern VA	9	19	NO _X	
Birmingham, AL	Birmingham CSA	Birmingham-Hoover- Cullman	1	9	РМ	
Boston, MA	Boston CSA	Boston-Worcester- Manchester	3	18	CO, PM, NO _X	
Chicago, IL	Chicago CSA	Chicago-Naperville- Michigan City	11	15	PM, NO _X	
Dallas, TX	Dallas CSA	Dallas-Fort Worth	19	0		
Denver, CO	Denver CSA	Denver-Aurora- Boulder	12	3	CO, PM	
Detroit, MI	Detroit CSA	Detroit-Warren-Flint	0	9	PM	
Houston, TX	Houston CSA	Houston-Baytown- Huntsville	21	0	CO, PM, NO _X	
Los Angeles, CA	Los Angeles CSA	Los Angeles-Long Beach-Riverside	47	3	CO, PM, SO _X , NO _X	
Minneapolis, MN	Minneapolis CSA	Minneapolis-St. Paul- St. Cloud	2	6		
New York, NY	New York CSA	New York-Newark- Bridgeport	20	10	CO, PM, SO _x , NO _x	
Philadelphia, PA	Philadelphia CSA	Philadelphia-Camden- Vineland	9	8	PM, NO _X	
Phoenix, AZ	Phoenix CBSA	Phoenix-Mesa- Scottsdale	14	17	CO, PM	
Pittsburgh, PA	Pittsburgh CSA	Pittsburgh-New Castle	2	12	CO, PM	
Salt Lake City, UT	Salt Lake City CSA	Salt Lake City-Ogden- Clearfield	2	10		
San Antonio, TX	San Antonio CBSA	San Antonio	5	0		
San Francisco, CA	San Francisco CSA	San Jose- San Francisco- Oakland	25	6		
Seattle, WA	Seattle CSA	Seattle-Tacoma- Olympia	5	5	CO, PM	
St Louis, MO	St Louis CSA	St. Louis-St. Charles- Farmington	3	13	CO, PM, SO _X	

Table 3-9	Focus cities used in this and previous assessments
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^aDefined based on 2000 Census data from the <u>U.S. Census Bureau (2011</u>). ^bThe number of sites within each CSA/CBSA with AQS monitors meeting the year-round data set inclusion criteria.

°The number of sites within each CSA/CBSA with AQS monitors meeting the warm-season data set inclusion criteria; the warmseason data set includes May - September data from both the warm-season and year-round monitors meeting the warm-season data set inclusion criteria.

^dBoundaries for the 2010 CO ISA (U.S. EPA, 2010c) and 2009 PM ISA (U.S. EPA, 2009d) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA definitions; boundaries for the 2008 SO_X ISA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA (U.S. EPA, 2008c) and 2008 NO_X ISA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA (U.S. EPA, 2008b) focus cities were based on CSA/CBSA (U.S. EPA, 2008b) focus cities were based (U.S. EPA, 2008b) focus cities were ba similar metropolitan statistical area (MSA) definitions from the 1990 U.S. Census.

- 1The distribution of the 8-h daily max O_3 concentrations from 2007-2009 for each of the220 focus cities is included in Table 3-10. These city-specific distributions were extracted3from the warm-season data set and can be compared to the nationwide warm-season 8-h4daily max distribution for 2007-2009 in
- 5 Table 3-7 (and repeated in the first line of Table 3-10 for reference). The median 8-h 6 daily max concentration in these focus cities was 41 ppb, similar to the nationwide 7 median of 42 ppb. Seattle had the lowest median (31 ppb) and Salt Lake City had the 8 highest median (53 ppb) of the 20 cities investigated. The 99th percentile of the 8-h daily 9 max concentration in the focus cities was 84 ppb; similar once again to the nationwide 10 99th percentile of 80 ppb. Seattle had the lowest 99th percentile (64 ppb) and 11 Los Angeles had the highest 99th percentile (98 ppb) of the 20 cities investigated. In 12 aggregate, the 20 focus cities selected are similar in distribution to the nationwide data 13 set, but there is substantial city-to-city variability in the individual distributions of the 8-h 14 daily max concentrations based on the warm-season data set.
- 15 Maps showing the location of central monitoring sites with O_3 monitors reporting to AOS 16 for each of the 20 focus cities are included as supplemental material in Section 3.9.1, 17 Figure 3-76 through Figure 3-95; examples for Atlanta, Boston and Los Angeles are 18 shown in Figure 3-29 through Figure 3-31. The sites are delineated in the maps as year-19 round or warm-season based on their inclusion in the year-round data set and the warm-20 season data set (the warm-season data set includes May-September data from both the 21 warm-season monitors and the year-round monitors meeting the warm-season data 22 inclusion criteria). The maps also include the CSA/CBSA boundary selected for monitor 23 inclusion, the location of urban areas and water bodies, the major roadway network, as 24 well as the population gravity center based on the entire CSA/CBSA and the individual 25 focus city boundaries. Population gravity center is calculated from the average longitude 26 and latitude values for the input census tract centroids and represents the mean center of 27 the population in a given area. Census tract centroids are weighted by their population 28 during this calculation.

						_	_										
Time Period	N Monitors	N Obs	Mean	SD	Min	1	5	10	25	50	75	90	95	98	99	Max	Max Site ID ^b
8-h daily ma	x (2007-200	9) ^a															
Nationwide	1,194	993,677	42	15	2	12	20	24	32	42	52	61	67	75	80	172	450210002
8-h daily ma	x by CSA/C	BSA (200	7-2009)	a													
Atlanta CSA	11	7,844	47	16	2	15	22	27	36	47	58	67	72	81	87	124	130890002
Baltimore CSA	28	20,999	43	16	2	9	18	23	31	43	54	64	70	78	83	118	240030014
Birmingham CSA	10	7,676	44	15	2	14	21	25	34	44	54	63	68	76	83	108	010732006
Boston CSA	21	12,603	41	14	2	13	21	25	31	40	49	59	67	75	81	104	250270015
Chicago CSA	27	20,764	37	14	2	9	15	19	27	37	47	57	62	69	74	108	170310042
Dallas CSA	19	19,858	41	15	2	11	20	24	31	39	50	61	67	74	79	121	484390075
Denver CSA	15	12,217	44	15	2	8	18	24	34	44	55	63	68	72	76	98	080590006
Detroit CSA	9	5,016	45	14	2	15	23	28	35	44	52	62	69	77	83	100	260990009
Houston CSA	21	22,305	36	15	2	8	15	19	25	34	46	57	64	72	78	110	482011034
Los Angeles CSA	49	49,295	47	18	2	10	20	26	35	45	58	72	81	91	98	137	060710005
Minneapolis CSA	8	5,315	40	12	2	15	21	25	31	40	48	54	58	63	67	86	270031002
New York CSA	21	26,304	39	16	2	6	15	20	28	37	47	59	68	77	83	123	090050005
Philadelphia CSA	14	12,673	41	17	2	8	17	21	29	39	52	64	70	78	83	125	240150003
Phoenix CBSA	22	26,129	49	12	2	18	27	32	41	50	58	65	68	72	75	85	040137021
Pittsburgh CSA	13	9,814	43	15	2	12	19	24	32	43	53	62	68	74	78	100	420050001
Salt Lake City CSA	12	5,146	51	14	2	8	23	32	44	53	61	67	71	77	80	96	490353008
San Antonio CSA	5	4,701	39	13	2	13	20	23	29	37	46	56	62	67	72	90	480290032
San Francisco CSA	31	28,325	34	12	2	8	16	20	26	33	41	48	55	63	68	110	060010007
Seattle CSA	5	6,148	31	12	2	4	12	17	23	31	39	46	51	59	64	91	530330023
St Louis CSA	19	11,569	43	15	2	12	19	23	32	43	53	61	68	76	81	113	295100086
All CSAs/CBSAs listed	360	314,701	42	16	2	9	18	22	31	41	52	63	69	78	84	137	060710005

Table 3-10City-specific distributions of 8-h daily max ozone concentrations
(ppb) from the warm-season data set (2007-2009).

^aIncludes all validated data regardless of flags or regional concurrence and therefore may differ from data used for regulatory purposes.

^bAQS Site ID corresponding to the observation in the Max column.

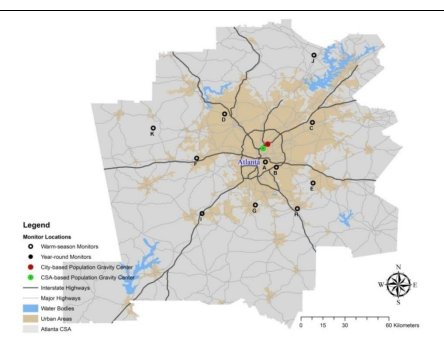


Figure 3-29 Map of the Atlanta CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

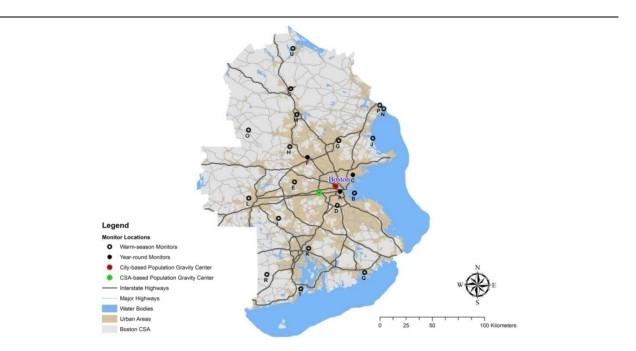


Figure 3-30 Map of the Boston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

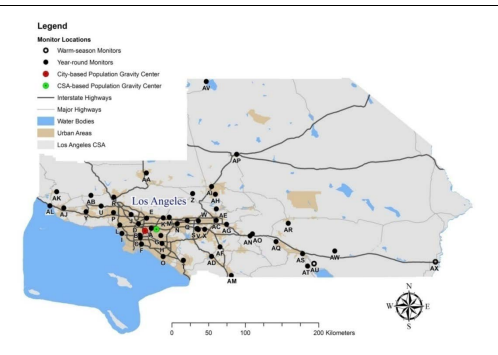


Figure 3-31 Map of the Los Angeles CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

1	The Atlanta CSA contains 11 warm-season monitors distributed evenly yet sparsely
2	around the city center (Figure 3-29). The population gravity center for the city and the
3	larger CSA are only separated by 4 km, indicating that the majority of the population
4	lives within or evenly distributed around the city limits. Atlanta is landlocked with a
5	radial network of interstate highways leading to the city center. The Boston CSA contains
6	3-year-round and 18 warm-season monitors spread evenly throughout the CSA. Boston is
7	a harbor city with the Atlantic Ocean to the east, resulting in the city-based population
8	gravity center being located 17 km east of the CSA-based population gravity center. The
9	Los Angeles CSA contains the largest number of monitors of the 20 CSA/CBSAs
10	investigated with 47 year-round and 3 warm-season monitors. These monitors are
11	primarily concentrated in the Los Angeles urban area with relatively few monitors
12	extending out to the northern and eastern reaches of the CSA. These unmonitored areas
13	are very sparsely populated, resulting in only 15 km separating the city-based and the
14	CSA-based population gravity centers despite the vast area of the Los Angeles CSA.
15	Other CSAs/CBSAs (see Section $3.9.1$) with monitors concentrated within the focus city
16	limits include Birmingham, Chicago, Denver, Houston, Phoenix, San Antonio, and Salt
17	Lake City. The remaining CSAs/CBSAs have monitors distributed more evenly
18	throughout the CSA/CBSA area. Baltimore is contained within the same CSA as

1Washington DC and suburbs, resulting in a 50-km separation (the largest of the focus2cities investigated) between the city-based population gravity center for Baltimore and3the CSA-based population gravity center for the Washington-Baltimore-Northern4Virginia CSA.

5 Box plots depicting the distribution of 2007-2009 warm-season 8-h daily max O_3 data 6 from each individual monitor in the 20 focus cities are included as supplemental material 7 in Section 3.9.2, Figure 3-96 through Figure 3-115; examples for Atlanta, Boston and 8 Los Angeles are shown in Figure 3-32 through Figure 3-34. The Atlanta CSA has little 9 spatial variability in 8-h daily max O_3 concentrations with median concentrations ranging 10 from 47 ppb at Sites I and J located far from the city center to 54 ppb at Site A located 11 closest to the city center. The variation in warm-season 8-h daily max concentrations are 12 also relatively similar across monitors with IQRs ranging from 17 ppb at Site J to 23 ppb 13 at Site B. The Boston CSA has more spatial variability in 8-h daily max O₃ 14 concentrations than the Atlanta CSA with median concentrations ranging from 33 ppb at 15 Site A nearest to the city center to 46 ppb at Site L located 84 km west of the city center. 16 For monitors located within and just adjacent to the Boston city limits (Sites A-D), the O_3 17 concentrations can vary over relatively short distances owing to differing degrees of NO_x 18 titration and influence from the local topography. Like the Atlanta CSA, the variation in 19 warm-season 8-h daily max concentrations are relatively similar across monitors within 20 the Boston CSA with IQRs ranging from 15 ppb at Site U to 21 ppb at Site K. The 21 Los Angeles CSA exhibits the most variability in O_3 concentrations between monitors of 22 all the CSAs/CBSAs investigated. The median 8-h daily max O₃ concentration in the 23 Los Angeles CSA ranged from 20 ppb at Site AM in the south-central extreme of the 24 CSA to 80 ppb at Site AE near Crestline, CA in the San Bernardino National Forest just 25 north of San Bernardino, CA. These two sites are at approximately the same longitude 26 and are separated by only 85 km, but the Crestline site is downwind of the Los Angeles 27 basin, resulting in substantially higher O₃ concentrations. Site AM also contains data for 28 only 2009, which could explain some of the deviation when comparing this site with 29 others in the Los Angeles CSA. Sites AM and AE also had the lowest (8 ppb) and highest 30 (28 ppb) IQR, respectively. The remaining focus cities exhibited spatial variability 31 ranging from uniform as in the Atlanta CSA to non-uniform as observed in the 32 Los Angeles CSA (see supplemental figures in Section 3.9.2).

Atlanta CSA

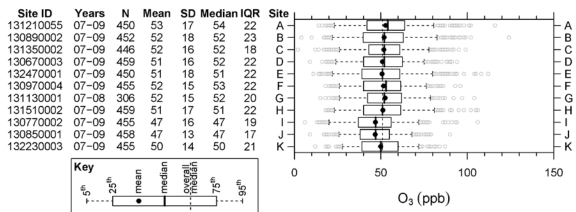


Figure 3-32 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Atlanta CSA.

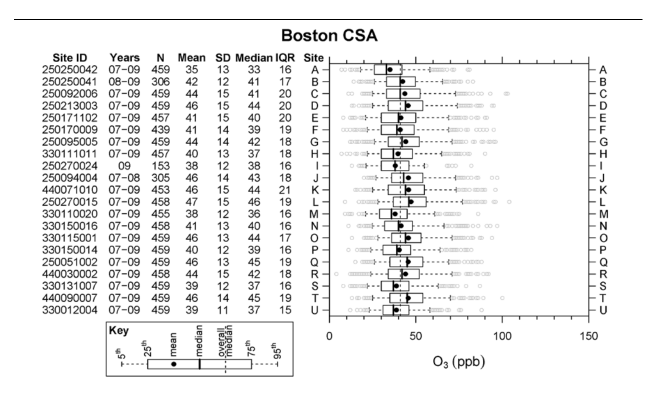


Figure 3-33 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Boston CSA.

					L	os /	Ange	eles CSA	
Site ID	Years	Ν	Mean		Mediar		Site	<u> </u>	4
060371602		458	48	13	47	17	A –		- A
060371301		306	36	9	34	10	В-		⊢ B
060371302		152	44	10	44	12	<u> </u>		ΗC
060371103		457	46	12	45	14	D –		ΡD
060372005		459	54	15	53	18	E-		ΗĒ
060374002		459	38	10	37	11	F -		F
060595001		459	50	12	49	14	G –		⊢ G
060590007		459	48	10	47	12	н –		Η
060375005		459	45	9	45	12	!-		<u>⊢!</u>
060371002		459	56	14	55	19	J –		- J
060370002		459	57	17	56	22	К-		ŀκ
060370113		459	48	10	47	13	L -		⊢ L
060370016		458	64	18	63	23	M –		- M
060371701		459	61	16	60	20	N -		- N
060591003		459	45	9	44	12	0-		⊢ o
060371201		459	61	14	60	19	P –		⊢ P
060711004		457	66 68	19	66 60	23	Q -		EB
060376012		457	68	18 19	69 65	27	R –		⊢ R - S
060650004 060592022		127 457	69 52	18 13	65 50	23 15	S – T –		FT
		457	62	12	62		U –		Γú
061112002		455 276	62 65	12		16 18	v –		Fv
060658005		459	68	19	64 67	24	w-		Fŵ
060658001		439	69	16	68	2 4 18	X –		Fx
061110007		440	54	10	54	12	Ŷ-		FŶ
060710012		456	67	13	67	18	z –		-ż
060379033		452	67	13	66	19	AA -		
061110009		458	58	11	58	14	AB -		
060719004		457	70	19	70	26	AC -		- AC
060659001		453	68	16	67	21	AD -	• @@@@@\$+	- AD
060710005		459	79	19	80	28	AE -		- AE
060656001		459	72	17	73	24	AF -		AF
060714003		459	73	18	73	25	AG -		AG
060714001		455	68	14	68	21	AH -	· · · · · · · · · · · · · · · · · · ·	
060710306		459	64	12	64	17	AI –	o @ami}tanno @m	
061113001		453	44	9	43	11	ÁJ –		- AJ
061111004		458	57	11	57	14	AK –	· · · · · · · · · · · · · · · · · · ·	- AK
061112003		457	41	9	40	12	AL -		- AL
060650009		153	22	8	20	8	ÁM –		
060650012		457	73	15	71	22	AN -		- AN
060651016		459	73	16	73	23	AO -		- AO
060710001		455	61	11	60	15	AP -		- AP
060655001		459	69	14	68	21	AQ -		- AQ
060719002		452	73	13	73	18	AR -	· · · · · · · · · · · · · · · · · · ·	- AR
060652002		448	62	13	61	18	AS -		- AS
060651999		283	49	17	50	22	AT -		- AT
060651010		153	59	10	59	15	AU -		- AU
060711234		453	59	10	58	13	AV-	o annami 🚺 + ann cano o o	- AV
060650008		265	58	10	57	14	AW-		- AV
060659003		444	42	10	42	13	AX -	o ww ww	- AX
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	+ ⁶			0,2		6 - 1		O ₃ (ppb)	
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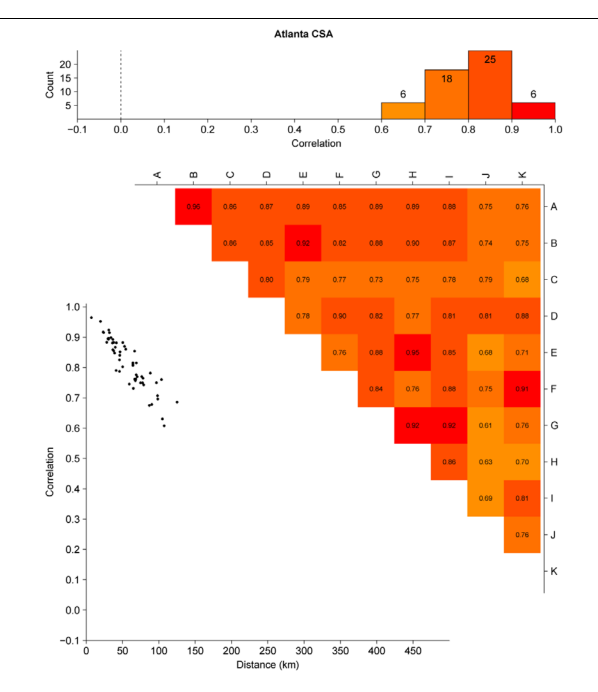
Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion Figure 3-34 criteria within the Los Angeles CSA.

1	Pair-wise monitor comparisons were used to further evaluate spatial variability between
2	monitors within the 20 focus cities. In the particular case of ground-level O ₃ , central-site
3	monitoring has been justified as a regional measure of exposure mainly on the grounds
4	that correlations between concentrations at neighboring sites measured over time are
5	usually high. In areas with multiple monitoring sites, averages over the monitors have
6	often been used to characterize population exposures. However, substantial differences in
7	concentrations between monitors can exist even though concentrations measured at the
8	monitoring sites are highly correlated, thus leading to the potential for exposure
9	misclassification error. Therefore, both the Pearson correlation coefficient and the
10	coefficient of divergence (COD) were calculated for each monitor pair within the
11	CSA/CBSAs using the 8-h daily max O ₃ data. The correlation provides an indication of
12	temporal linear dependence across sites while the COD provides an indication of the
13	variability in absolute concentrations across sites. The COD is defined as follows:

$$COD_{jk} = \sqrt{\frac{1}{p} \sum_{i=1}^{p} \left(\frac{X_{ij} - X_{ik}}{X_{ij} + X_{ik}}\right)^2}$$

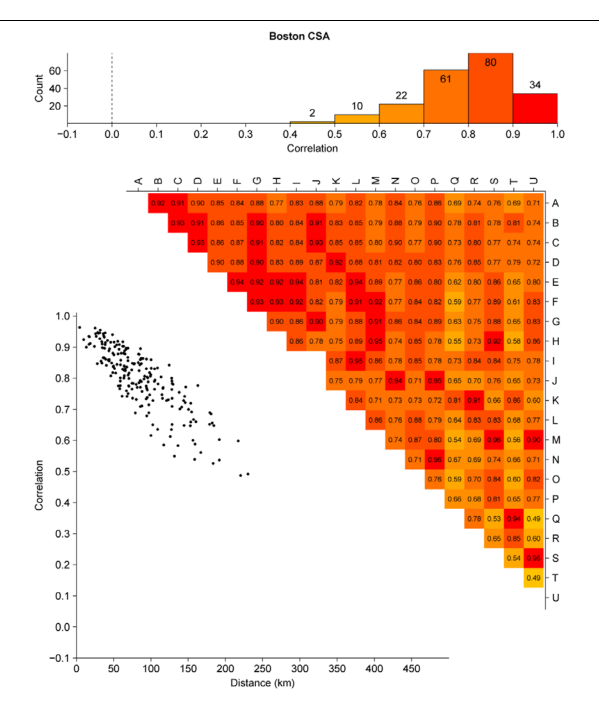
Equation 3-1

- 14 where X_{ij} and X_{ik} represent observed concentrations averaged over some measurement 15 averaging period *i* (hourly, daily, etc.) at sites *j* and *k*, and *p* is the number of paired 16 observations. A COD of 0 indicates there are no differences between concentrations at 17 paired sites (spatial homogeneity), while a COD approaching 1 indicates extreme spatial 18 heterogeneity. These methods for analysis of spatial variability follow those used in 19 previous ISAs for CO, PM, SO_X and NO_X as well as those used in Pinto et al. (2004) for 20 PM_{2.5}. 21 Histograms and contour matrices of the Pearson correlation coefficient between 8-h daily 22 max O_3 concentrations from each monitor pair are included as supplemental material in 23 Section 3.9.3, Figure 3-116 through Figure 3-135; examples for Atlanta, Boston and 24 Los Angeles are shown in Figure 3-35 through Figure 3-37. Likewise, histograms, 25 contour matrices, and scatter plots of the COD between 8-h daily max O₃ concentrations 26 from each monitor pair are included as supplemental material in Section 3.9.3, 27 Figure 3-136 through Figure 3-155; examples for Atlanta, Boston and Los Angeles are 28 shown in Figure 3-38 through Figure 3-40. These figures also contain scatter plots of
- 29 correlation and COD as a function of straight-line distance between monitor pairs.



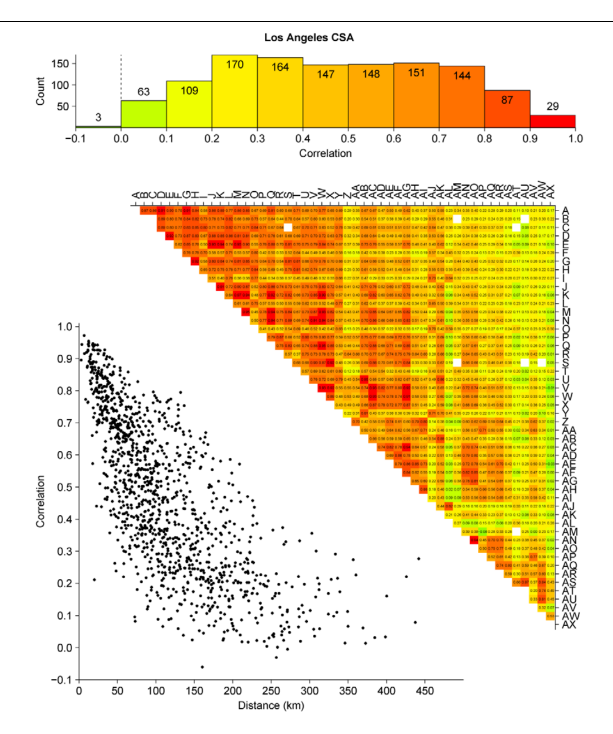
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the correlations.

Figure 3-35 Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.



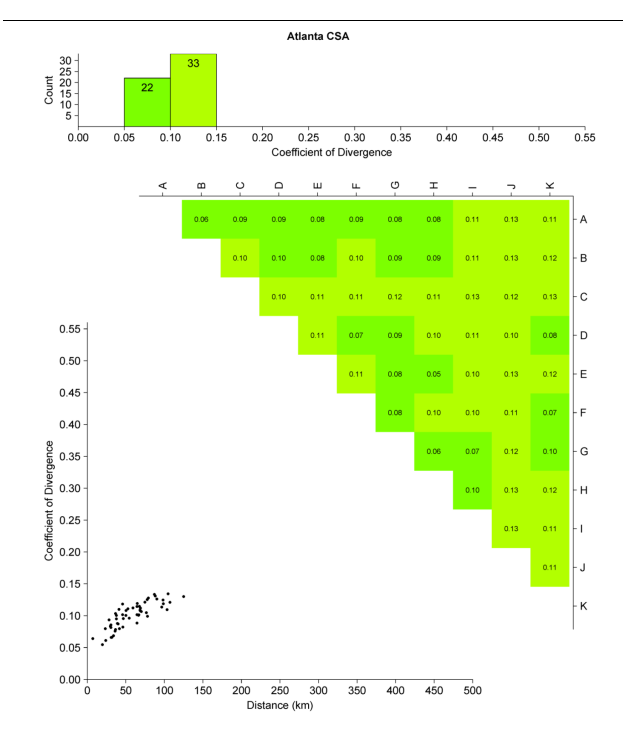
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the correlations.

Figure 3-36 Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.



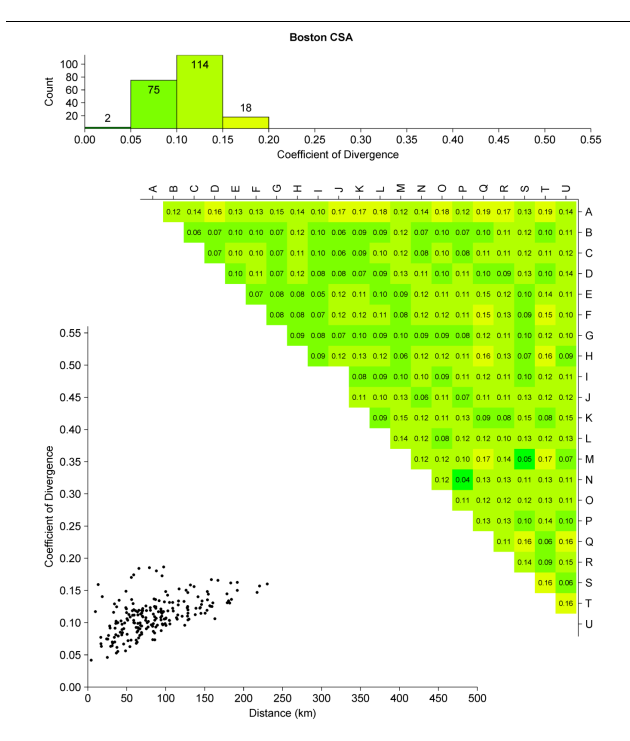
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the correlations.

Figure 3-37 Pair-wise monitor correlations expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.



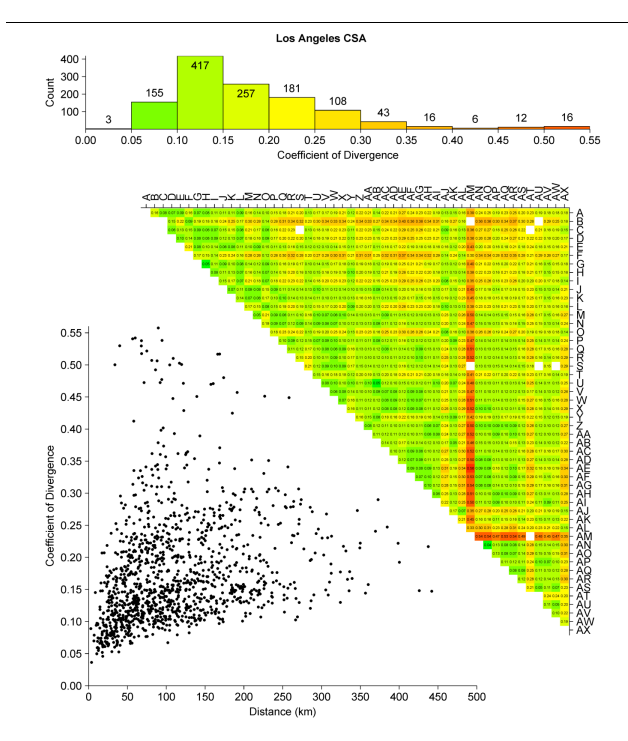
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the CODs.

Figure 3-38 Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the CODs.

Figure 3-39 Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the CODs.

Figure 3-40 Pair-wise monitor COD expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.

1	The monitor pairs within the Atlanta CSA (Figure 3-35) were generally well correlated
2	with correlations between 8-h daily max O_3 concentrations ranging from 0.61 to 0.96.
3	The correlations shown in the scatter plot were highest for close monitor pairs and
4	dropped off with distance in a near-linear form. At a monitor separation distance of
5	50 km or less, the correlations ranged from 0.79 to 0.96. The monitor pairs within the
6	Boston CSA (Figure 3-36) were also generally well correlated with correlations ranging
7	from 0.49 to 0.96. Again, the correlations shown in the scatter plot were highest for close
8	monitor pairs, but there was slightly more scatter in correlation as a function of distance
9	in the Boston CSA compared with the Atlanta CSA. At a monitor separation distance of
10	50 km or less, the correlations ranged from 0.81 to 0.96. The monitor pairs within the
11	Los Angeles CSA (Figure 3-37) showed a much broader range in correlations, extending
12	from -0.06 to 0.97. At a monitor separation distance of 50 km or less, the correlations
13	shown in the scatter plot ranged from 0.21 to 0.97. The negative and near-zero
14	correlations were between monitors with a relatively large separation distance (>150 km),
15	but even some of the closer monitor pairs were not very highly correlated. For example,
16	Site AL located at Emma Wood State Beach in Ventura and Site AK situated in an
17	agricultural valley surrounded by mountains 20 km inland (see map in Figure 3-41) had a
18	correlation coefficient of only 0.21 over the 2007-2009 warm-season time period. This
19	was slightly lower than the correlation between Site AL and Site AX on the Arizona
20	border, 441 km away ($R = 0.28$). San Francisco and Seattle (<u>Figure 3-133</u> and
21	Figure 3-134 in Section 3.9.3) also showed a broad range in pair-wise correlations, likely
22	resulting from their similar geography where background air coming in from the Pacific
23	Ocean rapidly mixes with urban pollutants such as NO_X and VOCs from coastal cities
24	and is transported downwind into diversified terrain to create spatially and temporally
25	varying O ₃ concentrations.

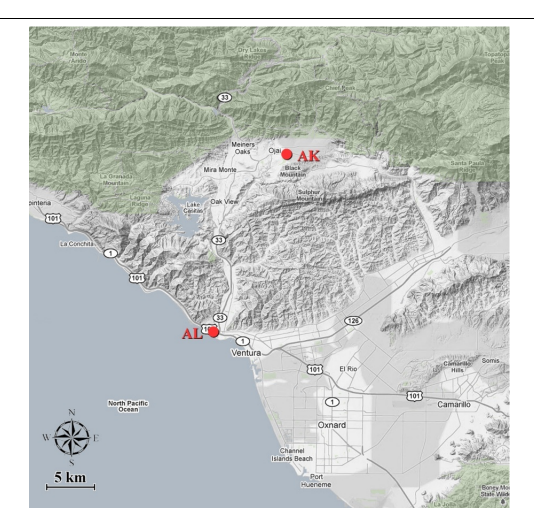


Figure 3-41	Terrain map showing the location of two nearby AQS ozone monitoring sites (red dots) along the western edge of the
	Los Angeles CSA. Site AL is near shore, 3 meters above sea level, while Site AK is in an agricultural valley surrounded by mountains,
	262 meters above sea level.

1	The COD between 8-h daily max O ₃ measured at paired monitors in all CSAs/CBSAs
2	(Figure 3-136 through Figure 3-155 in Section 3.9.3) were generally low, with values
3	similar to those shown in Figure 3-38 and Figure 3-39 for Atlanta and Boston. This
4	suggests a generally uniform distribution in the 8-h daily max O_3 concentration across
5	monitors within these cities and is consistent with the uniformity observed in the box
6	plots (e.g., Figure 3-32, Figure 3-33, and Figure 3-96 through Figure 3-115 in
7	Section 3.9.2). Los Angeles (Figure 3-34) and San Francisco (Figure 3-153 in
8	Section $3.9.3$), however, had several monitor pairs with COD >0.30 indicating greater
9	spatial heterogeneity. This is consistent with the variability observed in the box plots for
10	these two CSAs (Figure 3-34 and Figure 3-113 in Section 3.9.2). In particular, Site AM

1	in the Los Angeles CSA had consistently lower concentrations (median = 20 ppb,
2	IQR = 17-25 ppb) relative to other sites in the CSA (Figure 3-31), resulting in high CODs
3	with other monitors as shown in Figure 3-40. The O_3 monitor at Site AM is located on the
4	Pechanga Tribal Government Building in Temecula, CA, and began collecting data on
5	June 9, 2008. It is located in a suburban setting and is classified as a general background
6	monitor. Another close by site (site $ID = 060731201$) located in the Pala Reservation,
7	9.5 km south of this one (just outside the boundary of the Los Angeles CSA) reported
8	similarly low 2009 8-h daily max O_3 concentrations (median = 28 ppb, IQR = 23-32 ppb)
9	between May-June, 2009 (the only warm-season months with available data from this site
10	between 2007 and 2009).



Figure 3-42 Terrain map showing the location of four AQS ozone monitoring sites (red dots) located in or near the city limits in the center of the Boston CSA. Site characteristics range from Site A near downtown at 6 meters above sea level to Site D in a forested area on Blue Hill at 192 meters above sea level.

1 There are instances where sites in an urban area may exhibit substantial of	differences in
2 median concentrations, but still be moderately well correlated in time. For	For example, Sites
A and D in Boston (see terrain map in Figure 3-42) have an 11 ppb differ	erence in median
4 8-h daily max O_3 concentration (COD = 0.16), but a high correlation (R =	k = 0.90). In this
5 example, Site A is located in the Boston city limits at an elevation of 6 m	meters while Site
6 D is located 13 km to the south in a forested area on Blue Hill, the higher	est point in
7 Norfolk County (elevation = 192 meters). The difference in median O_3 co	concentration at
8 these two sites can be attributed to differing degrees of NO _X titration betw	tween the
9 neighborhood scale site (Site A) and the regional scale site (Site D) and t	to the influence
10 of local topography.	

11 Comparison of monitoring data within the selected focus cities has demonstrated 12 considerable variability between cities in the behavior of the O_3 concentration fields. 13 Median O_3 concentrations vary more within certain urban areas than others. Likewise, 14 pair-wise monitor statistics (R and COD) are dependent on the urban area under 15 investigation. These conclusions are consistent with those drawn in the 2006 O_3 AQCD 16 (U.S. EPA, 2006b) where a subset of these focus cities were investigated using similar 17 statistics. As a result, caution should be observed in using data from a sparse network of 18 ambient O₃ monitors to approximate community-scale exposures.

Neighborhood-Scale Variability and the Near-Road Environment

- 19 Ozone is a secondary pollutant formed in the atmosphere from precursor emissions and 20 therefore is generally more regionally homogeneous than primary pollutants emitted from 21 stationary or mobile point sources. However, O₃ titration from primary NO emissions 22 does result in substantial localized O₃ gradients. This is evident in the near-road 23 environment where fresh NO emissions from motor vehicles titrate O₃ present in the 24 urban background air, resulting in an O_3 gradient down-wind from the roadway. Ozone 25 titration occurring in street canyons where NO emissions are continuously being 26 generated is more efficient because of inhibited transport away from the source of NO. 27 Several studies have reported O_3 concentrations that increase with increasing distance
- 28 from the roadway, both upwind and downwind of the road. Beckerman et al. (2008) 29 measured O_3 profiles in the vicinity of heavily traveled roadways with Annual Average 30 Daily Traffic (AADT) >340,000 vehicles in Toronto, Canada. Ozone was observed to 31 increase with increasing distance from the roadway, both upwind and downwind of the 32 road. This is consistent with scavenging of O_3 in the near-road environment by reaction 33 with NO to form NO₂. Upwind of the road, concentrations were >75% of the maximum 34 observed value at >100 meters from the road: downwind, concentrations were 35 approximately 60% of the maximum within 200-400 meters of the road. The O₃

1	concentration adjacent to the road on the upwind side was approximately 40% of the
2	maximum value observed at the site. Concentrations measured with Ogawa passive
3	samplers over a 1-week period ranged from 7.3-19.4 ppb with the mean at the two sites
4	ranging from 13.0-14.7 ppb. In a study of patrol cars during trooper work shifts, <u>Riediker</u>
5	et al. (2003) made simultaneous 9-h O_3 measurements inside patrol cars, at the roadside,
6	and at a centrally-located ambient monitoring site. The roadside concentrations were
7	approximately 81% of the ambient values (mean of 22.8 ppb versus 28.3 ppb). Wind
8	direction relative to the roadway was not reported.
9	Johnson (1995) measured O ₃ , NO, and CO concentrations at upwind and downwind
10	locations near a variety of roadways in Cincinnati, OH. The effects of O3 scavenging by
11	NO were apparent in the O ₃ reduction in the interval between 9 meters upwind and
12	82 meters downwind of the road. A similar effect was observed by Rodes and Holland
13	(1981) during an earlier study in which outdoor O_3 concentrations were monitored
14	downwind of a freeway in Los Angeles, CA. In this study, O ₃ concentrations measured
15	near the roadway were approximately 20% of the concentrations measured
16	simultaneously at more distant locations judged to be unaffected by the roadway.
17	Minimal separation distances of the samplers from the roadway to eliminate measurable
18	influence were estimated to be approximately 400-500 meters for NO, NO ₂ , and O ₃ .
19	Similar results have been observed outside the U.S., for example in the city of Daegu,
20	Korea, where the yearly roadside concentrations of CO and SO ₂ showed a well-defined
21	decreasing trend with distance from the roadway, whereas concentrations of NO_2 and O_3
22	exhibited the reverse trend (Jo and Park, 2005). During the peak O_3 month of May, O_3
23	concentrations in a residential neighborhood were approximately 40% higher than
24	concentrations at roadside monitors located 1 meter from the edge of multiple-lane
25	freeways.
	-

3.6.2.2 Rural-Focused Variability and Ground-Level Vertical Gradients

26 AQS O₃ data for monitors located at several rural monitoring sites (e.g., national parks, 27 national forests, state parks, etc.) were used to investigate rural-focused O₃ concentration 28 variability in contrast with the urban-focused variability discussed in Section 3.6.2.1. 29 These rural monitoring sites tend to be less directly affected by anthropogenic pollution 30 sources than urban sites. However, they can be regularly affected by transport of O_3 or O_3 31 precursors from upwind urban areas, or by local anthropogenic sources within the rural 32 areas such as emissions from motor vehicles, power generation, biomass combustion, or 33 oil and gas operations. As a result, monitoring data from these rural locations are not 34 unaffected by anthropogenic emissions.

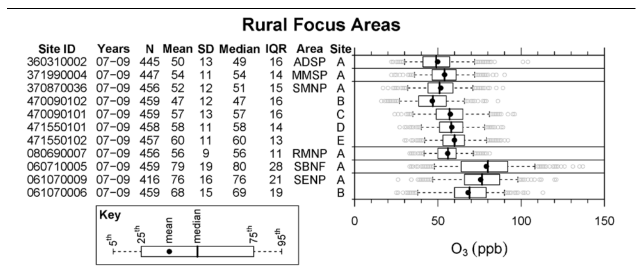
1	Six rural focus areas were selected for their geographic distribution across the U.S. as
2	well as their unique topography and relevance to the ecological assessment in Chapter 9 .
3	Table 3-11 lists the rural focus areas and provides some cursory site information along
4	with the number of available AQS monitors reporting year-round and only during the
5	warm-season. Accompanying box plots depicting the distribution of 2007-2009 warm-
6	season 8-h daily max O_3 data from each individual monitor in the six rural focus areas are
7	included in Figure 3-43. This analysis was restricted to AQS monitors meeting the same
8	data completeness criteria outlined in <u>Table 3-5</u> for a direct comparison with the 20 urban
9	focus areas investigated in Section $3.6.2.1$. Given the population-center emphasis of the
10	AQS network, limited monitoring sites (between one and five) were available for each
11	rural focus area. Expanded analyses of O3 concentrations measured using the more rural-
12	focused CASTNET monitoring network are included in Chapter 9.

Table 3-11Rural focus areas.

Focus Area	Short Name	Year-Round O₃ Monitoring Sites ^ª	Warm-Season O₃ Monitoring Sites ^b	Monitor Elevation (meters)	Site Descriptions
Adirondack State Park, NY	ADSP	1	0	1,483	One site on the summit of Whiteface Mountain in the Adirondack Mountains
Mount Mitchell State Park, NC	MMSP	0	1	1,982	One site near the summit of Mount Mitchell (highest point in the eastern U.S.) in the Appalachian Mountains
Great Smoky Mountain National Park, NC-TN	SMNP	2	3	564-2,021	Five different locations within Great Smoky Mountain National Park in the Appalachian Mountains
Rocky Mountain National Park, CO	RMNP	1	0	2,743	One site in a valley at the foot of Longs Peak in the Rocky Mountains
San Bernardino National Forest, CA	SBNFc	1	0	1,384	One site in Lake Gregory Regional Park (near Crestline, CA) in the San Bernardino Mountains
Sequoia National Park, CA	SENP	2	0	560-1,890	Two contrasting sites at different elevations within Sequoia NP in the Sierra Nevada Mountains

^aNumber of AQS monitors meeting the year-round data set inclusion criteria; the year-round data set is limited to these monitors. ^bNumber of AQS monitors meeting the warm-season data set inclusion criteria; the warm-season data set includes May-September data from both the warm-season and year-round monitors.

^cSame AQS site as Site AE in the Los Angeles CSA shown in Figure 3-31.



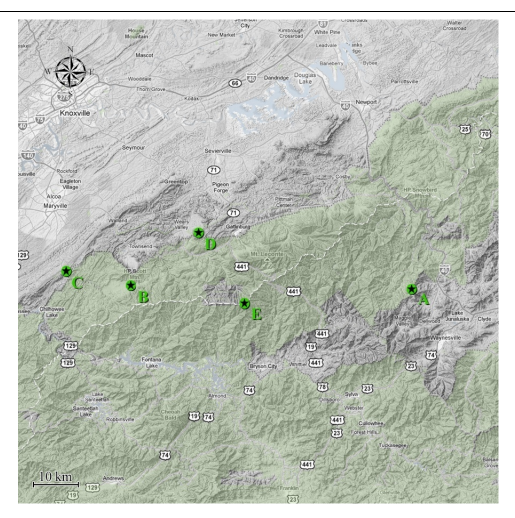
Note:lincludes: Adirondack State Park, NY (ADSP); Mount Mitchell State Park, NC (MMSP); Great Smoky Mountain National Park, NC-TN (SMNP); Rocky Mountain National Park, CO (RMNP); San Bernardino National Forest, CA (SBNF); and Sequoia National Park, CA (SENP).

Figure 3-43 Rural focus area site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the rural focus areas.

Eastern Rural Focus Areas

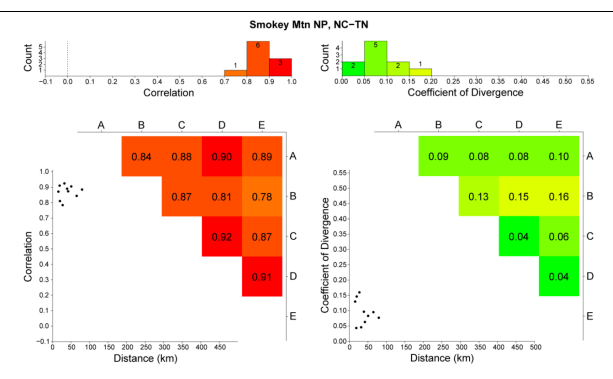
1	In the East, the distribution in warm-season 8-h daily max O_3 concentrations from the
2	Adirondack State Park (ADSP) site on Whiteface Mountain in Upstate NY
3	(median = 49 ppb) (Figure 3-43) was among the lowest of the rural focus monitors
4	investigated, but was still higher than concentration distributions measured in the Boston
5	CSA (medians ranging from 33 to 46 ppb) (Figure 3-33) located 320 km to the southeast.
6	The ADSP AQS site was included in an analysis for the 2006 O_3 AQCD (U.S. EPA,
7	<u>2006b</u>) and had the lowest year-round median hourly O_3 concentration of the rural
8	forested sites investigated (including Yellowstone NP, the Great Smoky Mountains NP,
9	and Shenandoah NP). For the Appalachian Mountain monitors in Mount Mitchell State
10	Park, NC (MMSP) and Great Smoky Mountain National Park, NC-TN (SMNP), there
11	was a fair amount of variability in concentration distribution. Within SMNP, the median
12	warm-season 8-h daily max O_3 concentration ranged from 47 ppb at the lowest elevation
13	site (elevation = 564 meters; site ID = 470090102) to 60 ppb at the highest elevation site
14	(elevation = $2,021$ meters; site ID = 471550102); these sites are shown on the terrain map
15	in Figure 3-44. The warm-season median 8-h daily max O_3 concentration gradient
16	between these two sites located 26.2 km apart in SMNP was 0.9 ppb per 100 meters
17	elevation gain.

1	Data from the five sites within SMNP allowed for further investigation of spatial
2	variability within the park; Figure 3-45 contains histograms, contour plots and scatter
3	plots as a function of distance for the pair-wise correlation and COD (defined in
4	Equation 3-1) for SMNP. The correlations between the five sites ranged from 0.78 to
5	0.92 and the CODs ranged from 0.04 to 0.16. The plots of correlation and COD as a
6	function of distance between SMNP monitor pairs in Figure 3-45 show a large degree of
7	spatial variability between monitors over relatively short distances. A host of factors may
8	contribute to these variations, including proximity to local O ₃ precursor emissions,
9	variations in boundary-layer influences, meteorology and stratospheric intrusion as a
10	function of elevation, and differences in wind patterns and transport behavior due to local
11	topography.



Note: The lowest elevation site (Site B) is 564 meters above sea level, while the highest elevation site (Site E) is 2,021 meters above sea level.

Figure 3-44 Terrain map showing the location of five AQS ozone monitoring sites (green/black stars) in Great Smoky Mountain National Park, NC-TN (SMNP).



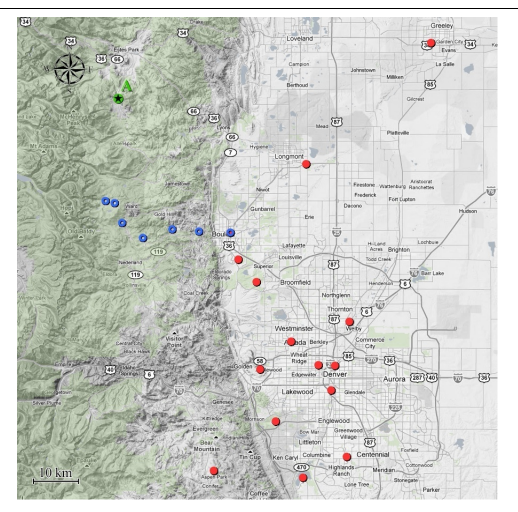
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histograms includes the number of monitor pairs per bin and the contour matrix includes the numeric values of the correlations and CODs.

Figure 3-45 Pair-wise monitor correlations (left) and coefficients of divergence (COD, right) expressed as a histogram (top), contour matrix (middle) and scatter plot vs. distance between monitors (bottom) for Great Smoky Mountain National Park, NC-TN (SMNP).

Western Rural Focus Areas

1	The Rocky Mountain National Park (RMNP) site in Colorado at 2,743 meters in
2	elevation had a warm-season 8-h daily max O ₃ concentration distribution
3	(median = 56 ppb, $IQR = 11 ppb$) (Figure 3-43) that is comparable to the distributions at
4	sites in the Denver CSA located 75 km southeast at elevations around 1,600 meters
5	(medians ranging from 41 to 59 ppb, IQRs ranging from 10 to 16 ppb; see Figure 3-102
6	in Section <u>3.9.1</u>). In nearby Boulder County, CO, a 1-year time-series (Sep 2007-Aug
7	2008) of ambient surface-level O_3 measurements was collected by <u>Brodin et al. (2010</u>)
8	along an elevation gradient ranging from 1,608 meters to 3,528 meters. The 7 sites used
9	in this study are shown in Figure 3-46 along with the RMNP site and the 15 Denver CSA
10	sites. In fall, winter, and spring, they observed a clear monotonic increase in O_3
11	concentration with elevation, with a rate of increase in the mean O_3 concentration of
12	1.5 ppb per 100 meters elevation gain during winter. In summer, the O ₃ gradient was
13	similar in magnitude over the seven-site transect (1.3 ppb per 100 meters), but much less

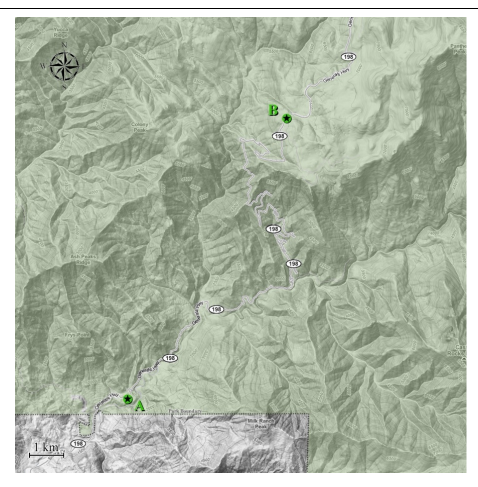
1 monotonic; the majority of the vertical gradient occurred between the lowest two sites 2 (4.5 ppb per 100 meters) and between the highest two sites (5.5 ppb per 100 meters), with 3 the middle five sites all having approximately equal median O₃ concentrations. Ozone 4 concentrations at the lowest site in Boulder were influenced by NO titration as evidenced 5 by traffic-related diel cycles in O_3 concentrations, but the remaining six sites were located 6 at elevation in more rural/remote settings and illustrate a positive surface-level O₃ 7 elevation gradient similar to that seen in SMNP and typical of areas under less direct 8 influence of boundary layer pollution.



Note: Elevations range from approximately 1,600 meters above sea level in Denver and Boulder to 3,528 meters above sea level at the highest mountainous site.

Figure 3-46 Terrain map showing the location of the AQS ozone monitoring site in Rocky Mountain National Park, CO (black/green star) and the Denver CSA (red dots) along with ozone monitoring sites used in the <u>Brodin et al. (2010</u>) study (blue circles).

1	The three sites in California-one in San Bernardino National Forest (SBNF) and two in
2	Sequoia National Park (SENP)-had the highest distribution of 8-h daily max O_3
3	concentrations of the selected rural focus area monitors included in Figure 3-43. The
4	SBNF site had a warm-season 8-h daily max O_3 concentration mean of 80 ppb and a
5	maximum of 137 ppb measured on July 1, 2007. This site is located in Crestline, CA,
6	90 km down-wind of Los Angeles in the San Bernardino Mountains. This site was
7	included in the Los Angeles CSA shown in Figure 3-31 (Site AE) and had the highest
8	median 8-h daily max O_3 concentration of any AQS site in the Los Angeles CSA during
9	this time period (Figure 3-34). This site was also included in an analysis performed for
10	the 2006 O_3 AQCD (U.S. EPA, 2006b) where similarly high O_3 concentrations were
11	observed using 2004 year-round hourly observations.



Note: The lower site (site ID = 061070009) is 560 meters above sea level and the higher site (site ID = 061070006) is 1,890 meters above sea level.

Figure 3-47 Terrain map showing the location of two AQS ozone monitoring sites (black/green stars) in Sequoia National Park, CA.

12

1	The two sites in SENP are located 9.7 km apart at contrasting elevations as is illustrated
2	in the terrain map in Figure 3-47. The correlation in 8-h daily max O_3 between these two
3	sites was 0.86 and the COD was 0.09, which are within the range in correlations and
4	CODs for SMNP (Figure 3-45). The distribution of 8-h daily max O_3 concentrations at
5	the lower elevation site (elevation = 560 meters; site ID = 061070009) is shifted slightly
6	higher with a median of 76 ppb compared to the higher elevation site
7	(elevation = $1,890$ meters; site ID = 061070006) with a median of 69 ppb. The lower
8	elevation site is located at the entrance to the park and is at a low enough elevation to be
9	influenced by boundary layer pollution coming upwind from Fresno and the San Joaquin
10	Valley. The higher elevation site is in the free troposphere above the planetary boundary
11	layer and is less influenced by such pollution. This gives rise to a negative average
12	surface-level elevation gradient of -0.5 ppb per 100 meters elevation gain in SENP,
13	illustrating the location-specific complexities inherent to high-altitude surface-level O ₃
14	concentrations.
15	Since O ₃ produced from emissions in urban areas is transported to more rural downwind
16	locations, elevated O ₃ concentrations can occur at considerable distances from urban
17	centers. In addition, major sources of O ₃ precursors such as highways, power plants,
18	biomass combustion, and oil and gas operations are commonly found in rural areas,
19	adding to the O_3 in these areas. Due to lower chemical scavenging in non-urban areas, O_3
20	tends to persist longer in rural than in urban areas which tends to lead to higher
21	cumulative exposures in rural areas influenced by anthropogenic precursor emissions.
22	The persistently high O ₃ concentrations observed at many of these rural sites investigated
23	here indicate that cumulative exposures for humans and vegetation in rural areas can be
24	substantial and often higher than cumulative exposures in urban areas.

3.6.3 Temporal Variability

3.6.3.1 Multiyear Trends

25	As reported in the 2010 National Air Quality Status and Trends report (U.S. EPA,
26	<u>2010e</u>), nation-wide surface level O_3 concentrations in the U.S. have declined gradually
27	over the last decade. Figure 3-48 shows the downward trend in the annual 4th highest 8-h
28	daily max O_3 concentration from 870 surface level monitors across the U.S. Figure 3-49
29	shows a similar trend in the annual second highest 1-h daily max O_3 concentration from
30	875 surface level monitors. The median annual 4th highest 8-h daily max dropped from
31	88 ppb in 1998 to 71 ppb in 2010. Likewise, the median annual second highest 1-h daily
32	max dropped from 109 ppb in 1998 to 86 ppb in 2010. The large decreases in 2003 and

1	2004 in both figures coincides with NO_X emissions reductions resulting from
2	implementation of the NO _X State Implementation Plan (SIP) Call rule, which began in
3	2003 and was fully implemented in 2004. This rule was designed to reduce NO_X
4	emissions from power plants and other large combustion sources in the eastern U.S.
5	Reductions in mobile NO_X emissions nationwide from the implementation of recent
6	vehicle and fuel standards could also be adding to the gradual decline in nationwide
7	surface level O ₃ concentrations (Dallmann and Harley, 2010).

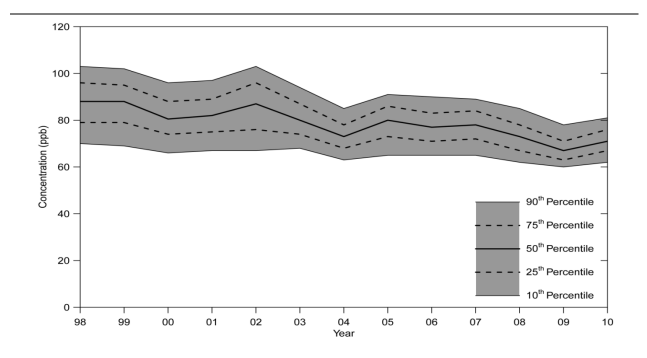
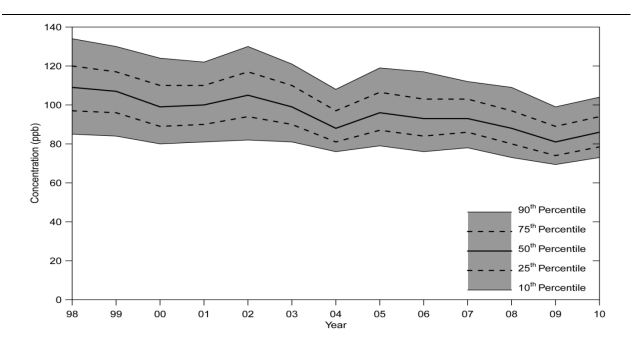
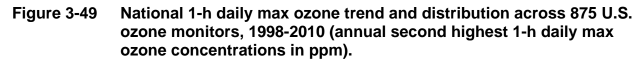


Figure 3-48 National 8-h daily max ozone trend and distribution across 870 U.S. ozone monitors, 1998-2010 (annual 4th highest 8-h daily max ozone concentrations in ppm).





1	The distributional percentiles (10th, 25th, 75th, and 90th) displayed in Figure 3-48 and
2	<u>Figure 3-49</u> reveal a gradual tightening of the O_3 concentration distribution observed
3	across monitors. For the annual 4th highest 8-h daily max O_3 concentration, the IQR
4	decreased from 17 ppb in 1998 to 9 ppb in 2010. Likewise, for the annual second highest
5	1-h daily max O_3 concentration, the IQR decreased from 23 ppb in 1998 to 16 ppb in
6	2010. A similar tightening was observed for the wider percentiles (90th-10th) for both
7	averaging times.
8	Weather can have a strong influence on the O_3 trends shown in Figure 3-48 and
9	<u>Figure 3-49</u> . The number of hot, dry days can substantially alter the number of high O_3
10	days in any given year, even if O ₃ forming emissions do not change. To better evaluate
11	the progress and effectiveness of emissions reduction programs, EPA uses a statistical
12	model to estimate the influence of atypical weather on O_3 formation (U.S. EPA, 2010e).
13	After adjusting for the influence of weather, the downward trend in surface level national
14	8-h daily max O_3 concentrations between 2001 and 2008 increased slightly from an 8%
15	reduction prior to adjustment for weather to an 11% reduction after adjustment for
16	weather (<u>U.S. EPA, 2010e</u>).
17	A regional breakdown of the trend in O ₃ concentrations for the 8-hour and 1-hour metrics
18	is included in Figure 3-50 and Figure 3-51, respectively. In general, the trends are region-

1	specific with a substantial amount of year-to-year variability. The reduction in NO_X and
2	O_3 during the 2003-2004 timeframe is particularly evident in the North Central and
3	Northeastern U.S. where the NO _X SIP Call was focused (<u>U.S. EPA, 2010e</u>). The western
4	region (including Alaska and Hawaii but excluding California) started out with the lowest
5	annual O_3 concentration in 1998 and exhibits the least amount of reduction when
6	compared to 2010 concentrations (11% reduction in the average annual 4th highest 8-h
7	daily max and 13% reduction in the average annual second highest 1-h daily max). In
8	contrast, California-which has some of the highest concentrations of the identified
9	regions—shows a larger downward trend in O3 concentrations over the same time period
10	(19% reduction in the average annual 4th highest 8-h daily max and 22% reduction in the
11	average annual second highest 1-h daily max).

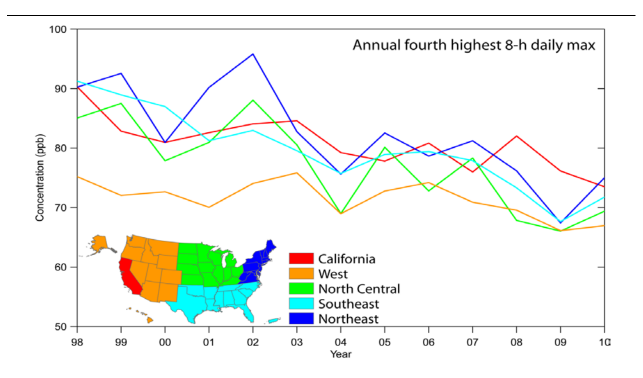


Figure 3-50 Trend in 8-h daily max ozone by region, 1998-2010 (annual 4th highest 8-h daily max ozone concentrations in ppm).

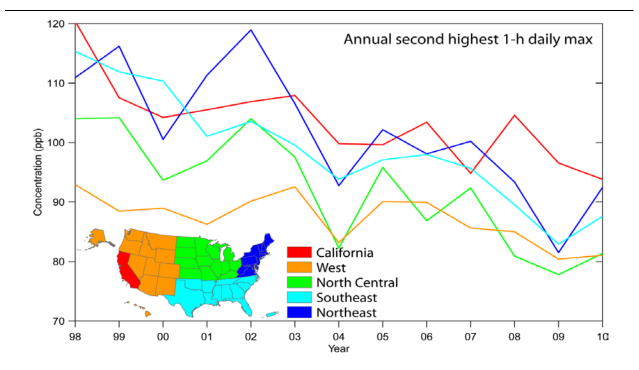


Figure 3-51 Trend in 1-h daily max ozone by region, 1998-2010 (annual second highest 1-h daily max ozone concentrations in ppm).

1	Narrowing the focus to changes in O_3 concentrations at the individual monitor level,
2	Figure 3-52 displays the 8-h O ₃ design value (4th highest 8-h daily max O ₃ concentration
3	occurring within a three-year period) for all available monitors for the 2008-2010 period
4	(Figure 3-52A) as well as the change in this design value between the 2001-2003 period
5	and the 2008-2010 period (Figure 3-52B). Figure 3-53 displays analogous information for
6	a 1-h O_3 design value (4th highest 1-h daily max O_3 concentration occurring within a
7	three-year period). As can be seen in both figures, the majority of monitors recorded a
8	decrease in design values when comparing the 2001-2003 period to the 2008-2010
9	period. Specifically, 699 of 853 sites (82%) included in Figure 3-52B for the 8-h design
10	value and 747 of 869 sites (86%) included in Figure 3-53B for the 1-h design value
11	reported a decrease of at least 6 ppb in the respective design values. The highest density
12	of monitors reporting decreases occurs in the Northeast. Only 8 sites (1%) reported an
13	increase of more than 5 ppb in the 8-h design value and only 16 sites (2%) reported an
14	increase of more than 5 ppb in the 1-h design value. These sites reporting an increase
15	between the 2001-2003 and the 2008-2010 periods were located primarily in the West.

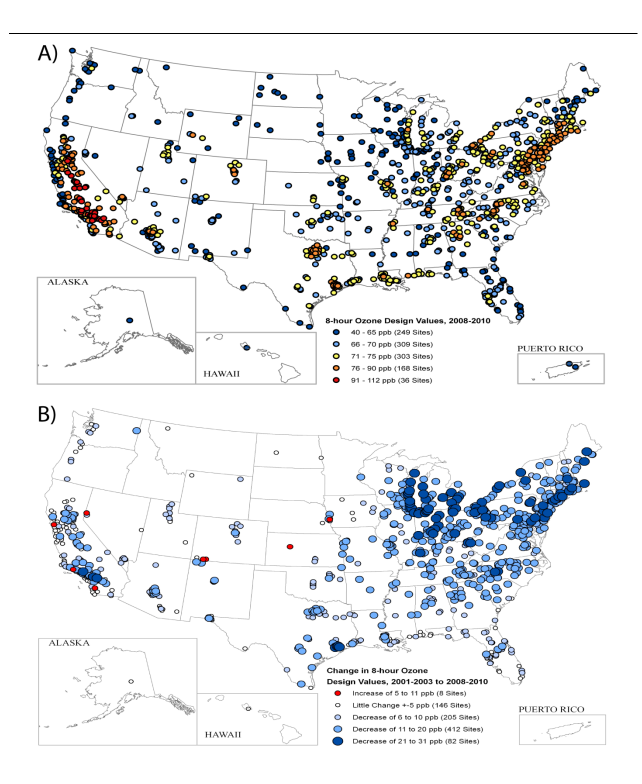


Figure 3-52 Individual monitor 8-h daily max ozone design values displayed A) for the 2008-2010 period and B) as the change since the 2001-2003 period.

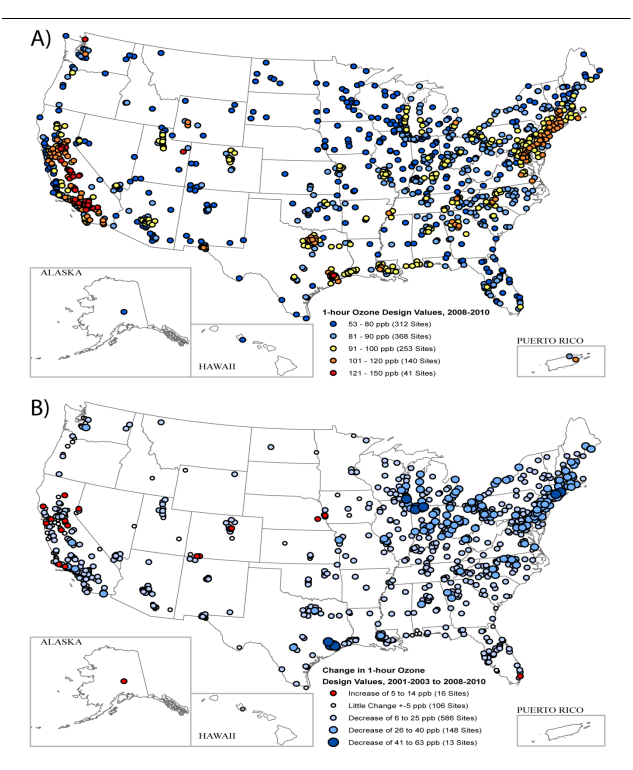


Figure 3-53 Individual monitor 1-h daily max ozone design values displayed A) for the 2008-2010 period and B) as the change since the 2001-2003 period.

1	Similar findings were reported for regional trends in the 4th highest 8-h daily max O_3
2	concentration between 2001 and 2008 in the 2010 National Air Quality Status and Trends
2 3	
4	report (<u>U.S. EPA, 2010e</u>). Individual sites that showed the greatest reduction in O_3 between 2001 and 2008 were in or near the following metropolitan areas: Anderson, IN;
4 5	
6	Chambersburg, PA; Chicago, IL; Cleveland, OH; Houston, TX; Michigan City, IN;
0 7	Milwaukee, WI; New York, NY; Racine, WI; Watertown, NY; and parts of Los Angeles,
8	CA. Individual sites that showed an increase in O_3 over this time period and had measured concentrations above the O_3 standard ¹ during the 2006-2008 time period were
9	located in or near the following metropolitan areas: Atlanta, GA; Baton Rouge, LA;
10	Birmingham, AL; Denver, CO; El Centro, CA; San Diego, CA; Seattle, WA; and parts of
11	Los Angeles, CA.
12	Pegues et al. (2012) investigated changes in 3-year average 8-h daily max O_3 design
13	values between 2003 and 2009 and found reductions at the majority of sites across the
14	U.S.; consistent with the findings in this section and in the 2010 National Air Quality
15	Status and Trends report (U.S. EPA, 2010e). Furthermore, they compared trends in O_3
16	design values between areas that were or were not classified as nonattainment of the
17	84 ppb O ₃ standard in the 2004 designations. Monitors designated nonattainment
18	achieved O ₃ design value reductions of 13.3 ppb on average while monitors designated in
19	attainment achieved reductions of 7.0 ppb on average.
20	Looking further back in time, Leibensperger et al. (2008) included an analysis of June-
21	August 8-h daily max O_3 trends from 1980-2006 using AQS data from over 2000 sites in
22	the contiguous U.S. They created an index for "pollution days" representing days when
23	the 8-h daily max O_3 concentration was greater than 84 ppb. The observed trend in
24	summertime O_3 pollution days over this 27 year period decreased at an average rate of -
25	0.84 days/year. The authors used several methods to deconstruct this trend into a
26	component coming from reductions in O ₃ precursor emissions (-1.50 days/year) and a
27	component coming from climate change (+0.63 days/year). The climate change impact is
28	a result of decreases in frequency of mid-latitude cyclones which serve to ventilate
29	surface air over the U.S. {Leibensperger, 2008, 611799@@author-year} conclude that
30	the reduction in frequency of mid-latitude cyclones over the 1980-2006 time period has
31	offset almost half of the air quality gains in the Northeastern U.S. that should have been
32	achieved from reductions of anthropogenic emissions alone over that period.
33	Averaging time can have an impact on perceived trends in surface level O ₃
34	concentrations. Lefohn et al. (2008) investigated the impact of using different exposure
35	indices on trends in surface level O_3 concentrations in the U.S. by comparing the annual
36	second highest 1-h average concentration, the annual 4th highest daily max 8-h average

¹ On September 16, 2009, EPA announced it would reconsider the 2008 O_3 NAAQS, which, at the time, included primary and secondary standards of 0.075 ppm (8-h daily max).

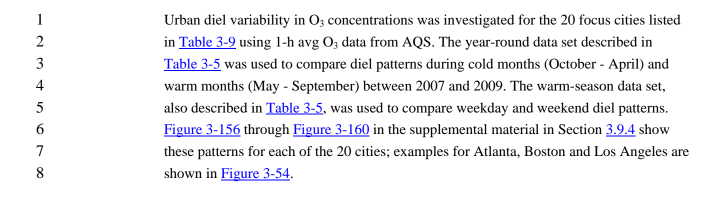
- 1 concentration, and the seasonally corrected 24-h W126 cumulative exposure index. 2 Between 1980 and 2005, most of the urban and rural sites across the U.S. included in this 3 study showed decreasing or zero trend for all three of these metrics. However, the 4 magnitude of this trend varied greatly by exposure index. The largest downward trend in 5 the 1-h and 8-h metrics listed above were observed in Southern California (>2%/yr 6 downward trend) but the W126 cumulative exposure metric showed large (>2%/yr) 7 downward trends in many locations across the U.S. including Southern California, the 8 Midwest and Northeast. By contrasting the 1980 – 2005 trends with more recent 1990 – 9 2005 trends, Lefohn et al. (2008) reported that a large number of sites (44%, 35% and 10 25% of sites for the 1-h, 8-h and W126 metrics, respectively) shifted from a negative 11 trend to no trend. These shifts in trends were attributed to slow changes in mid-level 12 concentrations (i.e., 60-90 ppb) following a more rapid change in peak concentrations in 13 the early years. A similar conclusion was drawn from nationwide O_3 data between 1980 – 14 2008 (Lefohn et al., 2010b), suggesting a shift in the O_3 distribution over this time period.
- 15 In contrast to the mostly urban observations included in the Pegues et al. (2012) study 16 above, several studies focusing on rural western monitors have reported positive trends in 17 O₃ concentrations over the last few decades. Jaffe and Ray (2007) investigated daytime 18 $(10 \text{ a.m.} - 6 \text{ p.m. local time}) O_3$ concentrations at rural sites in the northern and western 19 U.S. between 1987-2004. They found significantly positive trends in seven of the eleven 20 sites selected ranging from 0.19 ppb/yr in Gothic, CO to 0.51 ppb/yr in Rocky Mountain 21 NP, CO (mean trend of 0.26 ppb/yr at these seven sites). No significant trend was 22 observed for the two sites in Alaska and one site each in Wyoming and Montana. 23 Seasonal analyses were conducted on the sites having the longest records in Rocky 24 Mountain NP, Yellowstone, NP and Lassen NP and positive trends were found for all 25 seasons at all sites. As noted in the 2006 O₃ AQCD (U.S. EPA, 2006b), caution should be 26 exercised in using trends calculated at national parks to infer contributions from distant 27 sources either inside or outside of North America because of the influence of regional 28 pollution (see Section 3.4 for a discussion of background O₃ concentrations and 29 international transport).
- 30 Trends in baseline O_3 concentrations, defined as O_3 concentrations at a given site in the 31 absence of strong local influences, were estimated by region and season in the U.S. in 32 Chan and Vet (2010). The temperature-adjusted decadal (1997-2006) trends in estimated 33 baseline O_3 varied substantially by region and season. In the Pacific coastal regions, the 34 trends increased in all seasons except fall, but none of the trends were statistically 35 significant. In the eastern U.S., negative trends were observed in all seasons with the 36 exception of (1) insignificant positive trends in northeast Maine in summer, fall and 37 winter; (2) significant positive trends in the Midwest in winter; and 3) significant positive 38 trends at one site in Vermont in the summer. The density of sites in the central and

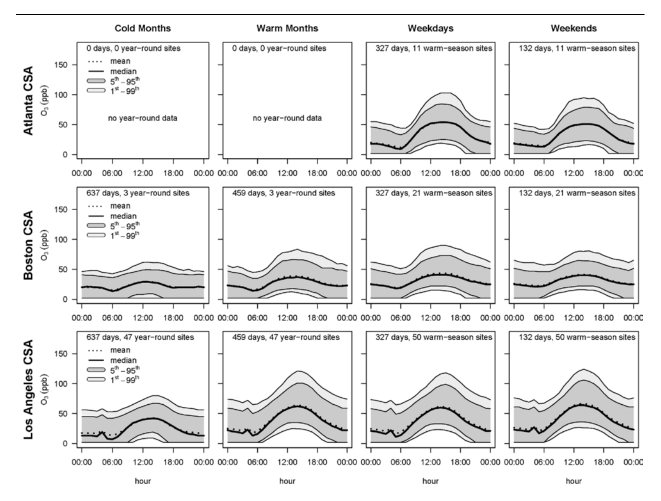
western U.S. were much lower than the coastal and eastern areas, but in general all sites
 showed trends that tended to be negative in the spring and fall but positive in the summer
 and winter.

- 4 Positive trends in marine boundary layer O_3 concentrations at several sites on the Pacific 5 Coast have been reported by other sources in the literature. Parrish et al. (2009) used 6 observations from multiple coastal sites in California and Washington and reported a 7 positive annual mean trend of 0.34 ± 0.09 ppb/yr between the mid-1980s and 2007 (exact 8 dates varied by site depending on available data). A seasonal stratification of the data at
- 9these sites showed the largest positive trend in the spring $(0.46 \pm 0.13 \text{ ppb/yr})$ with a10smaller and non-significant positive trend during fall $(0.12 \pm 0.14 \text{ ppb/yr})$. These results11agree with positive trends in springtime O_3 mixing ratios reported in an earlier study12(Jaffe et al., 2003). Positive trends in O_3 measurements in the free troposphere above13western North America at altitudes of 3-8 km (above sea level) during April and May of
- western North America at altitudes of 3-8 km (above sea level) during April and May of
 14
 1995 to 2008 were reported by <u>Cooper et al. (2010</u>) and discussed in Section <u>3.4.2</u> as they
 relate to intercontinental transport. Comparable trends were observed in the median as
 well as 5th, 33rd, 67th, and 95th percentiles of observations. Note, however, that these
 results relate to O₃ trends above ground level and not to surface O₃.
- 18Extending back to the 19th Century, Volz and Kley (1988) report a series of historic O319measurements from Europe. Comparing these with more contemporary measurements,20Parrish et al. (2009) report a 2 to 3 fold increase in boundary layer O3 mixing ratios over21the last 130 years with no indication of stabilization in recent years. Other long-term22observations of global trends in the burden of tropospheric O3 as they relate to climate23change are discussed in Chapter 10, Section 10.3.3.1.

3.6.3.2 Hourly Variations

24	Ozone concentrations frequently possess a strong degree of diel variability resulting from
25	daily patterns in temperature, sunlight, and precursor emissions. Other factors, such as the
26	relative importance of transport versus local photochemical production and loss rates, the
27	timing for entrainment of air from the nocturnal residual boundary layer, and the diurnal
28	variability in mixing layer height also play a role in daily O_3 patterns. The 2006 O_3
29	AQCD (U.S. EPA, 2006b) looked at composite urban diel variations from April to
30	October 2000 to 2004 and found 1-h maxima to occur in mid-afternoon and 1-h minima
31	to occur in early morning. On a national basis, however, there was a high degree of
32	spread in these times and caution was raised in extrapolating results from one city to
33	another in determining the time of day for O_3 maxima and minima.





Note: Uses the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half). Atlanta had no year-round monitors available for the cold month/warm month comparison.

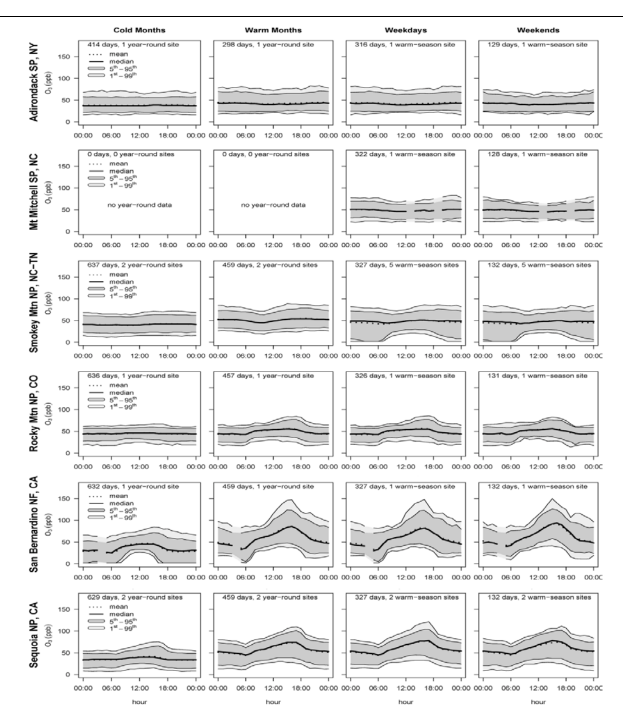
Figure 3-54 Diel patterns in 1-h avg ozone for Atlanta, Boston and Los Angeles between 2007 and 2009.

1	In general, all the urban areas showed 1-h daily max concentrations occurring typically in
2	the early afternoon. In all cities, these afternoon peaks were more pronounced in the
3	warm months than in the cold months. However, a small peak was still present during the
4	cold months. During warm months, the difference between the median daily extrema
5	varied considerably by city. For example, in Los Angeles, the median 1-h daily min
6	(10 ppb) at ~5:00 a.m. was 50 ppb less than the median 1-h daily max (60 ppb) at ~2:00
7	p.m. By contrast, in Boston, the median 1-h daily min (13 ppb) occurred at the same time,
8	but was only 25 ppb less than the median 1-h daily max (38 ppb). Cities with large daily
9	swings (>40 ppb) in median 1-h O ₃ concentrations included Atlanta, Birmingham,
10	Los Angeles, Phoenix, Pittsburgh, and Salt Lake City (Figure 3-156 through Figure 3-160
11	in Section 3.9.4). Cities with small daily swings (<25 ppb) in median 1-h O_3
12	concentrations included Boston, Minneapolis, San Francisco and Seattle (Figure 3-156
13	through <u>Figure 3-160</u> in Section <u>3.9.4</u>). These results are very similar to those found in
14	the 2006 O_3 AQCD (U.S. EPA, 2006b) where many of these same urban areas were
15	investigated. This supports the conclusions drawn in the previous O ₃ review that diel
16	patterns in O ₃ have remained stable over the last 20 years, with times of occurrence of the
17	daily maxima varying by no more than an hour from year to year.

18 Using the warm-season data, there was little difference in the median diel profiles for 19 weekdays compared with weekends across all urban areas. This result stresses the 20 complexity of O₃ formation and the importance of meteorology, entrainment, biogenic 21 precursor emissions, and transport in addition to anthropogenic precursor emissions. 22 There was, however, a subtle deviation between weekdays and weekends in the lower 23 percentiles (1st and 5th) of the distribution. The lower end of the distribution tended to be 24 lower on weekdays relative to weekends. This is consistent with analyses in the 2006 O_3 25 AQCD (U.S. EPA, 2006b) and is a result of lower traffic volumes on weekends relative 26 to weekdays, leading to less NO emissions and O₃ titration on the weekends.

27 Seasonal and site-to-site variations in diel patterns within a subset of the urban focus 28 areas presented here were investigated in the 2006 O₃ AQCD (U.S. EPA, 2006b). In 29 northern cities, there was substantial seasonal variability in the diel patterns with higher 30 extreme values in the O₃ distribution during the warm season than during the cold season. 31 In southern cities, the seasonal differences in extreme O₃ concentrations were much 32 smaller, and some of the highest O₃ concentrations in the Houston CSA were found 33 outside of summer. The general pattern that emerged from investigating site-to-site 34 variability within the urban areas was that peaks in 1-h avg O_3 concentrations are higher 35 and tend to occur later in the day at downwind sites relative to sites located in the urban 36 core. Differences between sites were not only related to the distance between them, but 37 also depend on the presence or absence of nearby O₃ sources or sinks.

1	Rural diel variability in O3 concentrations was investigated for the six rural focus areas
2	listed in Table 3-11 using 1-h avg O_3 data from AQS. As with the urban analysis, the
3	year-round data set described in <u>Table 3-5</u> was used to compare diel patterns during cold
4	months (October - April) and warm months (May - September) between 2007 and 2009.
5	The warm-season data set, also described in Table 3-5, was used to compare weekday
6	and weekend diel patterns. Figure 3-55 shows the diel patterns for each of the rural areas
7	investigated.



Note: Uses the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half). Mt. Mitchell SP, NC had no year-round monitors available for the cold month/warm month comparison.

Figure 3-55 Diel patterns in 1-h avg ozone for six rural focus areas between 2007 and 2009.

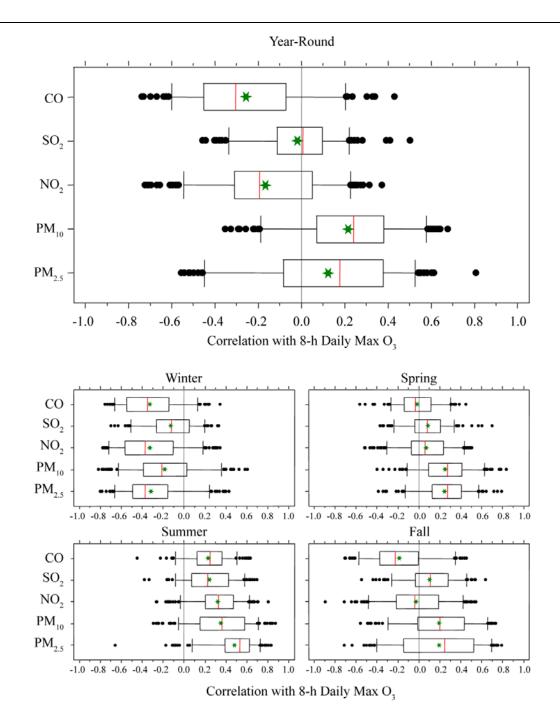
1	There was considerable variability in the diel patterns observed in the six rural focus
2	areas. The selected mountainous eastern sites in ADSP, MMSP, and SMNP exhibited a
3	generally flat profile with little hourly variability in the median concentration and the
4	upper percentiles. In SMNP, there was some diel variability in the lower percentiles, with
5	higher values during the daylight hours in the warm season data. This behavior was not
6	present in the data coming from the two year-round monitors located at lower elevation
7	sites (Sites C and Site D; see map in Figure 3-44), however, possibly resulting from
8	differing impacts from local sources within SMNP. For the western rural areas, there was
9	a clear diel pattern to the hourly O_3 data with a peak in concentration in the afternoon
10	similar to those seen in the urban areas in Figure 3-54 and Figure 3-156 through
11	Figure 3-160 in Section 3.9.4. This was especially obvious at the SBNF site which sits
12	90 km east of Los Angeles in the San Bernardino Mountains at an elevation of
13	1,384 meters. This site was located here to monitor O ₃ transported downwind from major
14	urban areas in the South Coast Air Basin. It had the highest 2007-2009 median 8-h daily
15	max O_3 concentration of any AQS site in the Los Angeles CSA (see Figure 3-34), and is
16	clearly impacted by the upwind urban plume which has sufficient time and sunlight to
17	form O_3 from precursor emissions and concentrate the O_3 in the shallow boundary layer
18	present at this elevation.
19	As with the urban analysis, there was little difference observed in the weekday and
20	weekend diel profiles using the warm season data, even down at the lower percentiles in

20weekend diel profiles using the warm-season data, even down at the lower percentiles in21the distribution. This is consistent with the regional nature of tropospheric O3. Using the22year-round data, there was an upward shift in the distribution going from the cold months23to the warm months, and in some instances the general shape of the distribution changed24considerably as was seen in several urban sites.

3.6.4 Associations with Co-pollutants

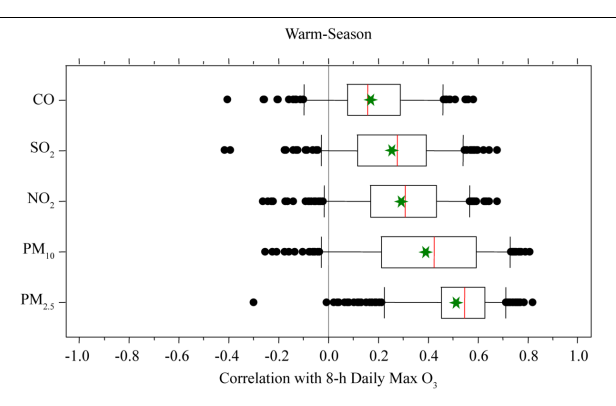
25	Correlations between O ₃ and other criteria pollutants are discussed in this section. Since
26	O_3 is a secondary pollutant formed in the atmosphere from precursor emissions, its
27	correlation with primary pollutants such as CO and NO_X can vary substantially by
28	location. Furthermore, O ₃ formation is strongly influenced by meteorology, entrainment,
29	and transport of both O_3 and O_3 precursors, resulting in a broad range in correlations with
30	other pollutants which can vary substantially with season. This section focuses on
31	correlations between O_3 and other criteria pollutants measured at the mostly urban AQS
32	sites: a more detailed discussion of O_3 and O_3 -precursor relationships is included in
33	Section <u>3.2.4</u> . To investigate correlations with co-pollutants, 8-h daily max O_3 from the
34	year-round and warm-season data sets (Table 3-6 and

1 2 3 4	Table 3-7) were compared with co-located 24-h avg CO, SO ₂ , NO ₂ , PM _{2.5} and PM ₁₀ obtained from AQS for 2007-2009. Figure 3-56 and Figure 3-57 contain co-pollutant box plots of the correlation between co-located monitors for the year-round data set and the warm-season data set, respectively.
5	The year-round 8-h daily max O_3 data (Figure 3-56) had a very wide range in correlations
6	with all the 24-h avg co-pollutants. A clearer pattern emerged when the data were
7	stratified by season (bottom four plots in Figure 3-56) with mostly negative correlations
8	in the winter and mostly positive correlations in the summer for all co-pollutants. In
9	summer, the IQR in correlations is positive for all co-pollutants. However, the median
10	seasonal correlations are still modest at best with the highest positive correlation at 0.52
11	for $PM_{2.5}$ in the summer and the highest negative correlation at -0.38 for $PM_{2.5}$ in the
12	winter. Spring and fall lie in between with spring having a slightly narrower distribution
13	than fall for all copollutants. The warm-season 8-h daily max O_3 data (Figure 3-57)
14	shows a very similar distribution to the summer stratification of the year-round data due
15	to their overlap in time periods (May-Sept and Jun-Aug, respectively).



Note: Year round (Top figure), and with seasonal stratification (Bottom four figures). Shown are the median (red line), mean (green star), inner-quartile range (box), 5th and 95th percentiles (whiskers) and extremes (black circles).

Figure 3-56 Distribution of Pearson correlation coefficients for comparison of 8-h daily max ozone from the year-round data set with co-located 24-h avg CO, SO₂, NO₂, PM₁₀ and PM_{2.5} from AQS, 2007-2009.



Note: Shown are the median (red line), mean (green star), inner-quartile range (box), 5th and 95th percentiles (whiskers), and extremes (black circles).

Figure 3-57 Distribution of Pearson correlation coefficients for comparison of 8-h daily max ozone from the warm-season (May-Sept) data set with co-located 24-h avg CO, SO₂, NO₂, PM₁₀ and PM_{2.5} from AQS, 2007-2009.

1	The seasonal fluctuations in correlations present in Figure 3-56 result in part from the
2	mixture of primary and secondary sources for the co-pollutants. For example, O_3 is a
3	secondary pollutant whereas PM _{2.5} has both primary and secondary origins and these two
4	pollutants show the largest summertime/wintertime swing in correlation distributions.
5	This situation arises because the secondary component to PM _{2.5} is larger during the
6	summer and is formed in conditions conducive to secondary O ₃ formation. This results in
7	positive correlations between O_3 and $PM_{2.5}$ during the summer. During the winter,
8	photochemical production of O ₃ is much smaller than during summer and O ₃ comes
9	mainly from aloft, i.e., the free troposphere (see Section $3.4.1.1$ for further details). In
10	addition, concentrations of PM _{2.5} are much lower aloft. On relatively clean days, this can
11	lead to high concentrations of O ₃ and lower concentrations of primary pollutants such as
12	$PM_{2.5}$ or NO. On relatively dirty days with elevated NO and $PM_{2.5}$, the intruding O ₃ is
13	readily titrated by NO in the boundary layer. These processes result in negative
14	correlations between O_3 and $PM_{2.5}$ during the winter.

3.7 Chapter Summary

1	This section contains a summary of the major topics included in this chapter on the
2	atmospheric chemistry and ambient concentrations of tropospheric O_3 and other related
3	photochemical oxidants. This chapter has built upon information previously reported in
4	the 2006 O ₃ AQCD (U.S. EPA, 2006b) and includes updated material on: (1) physical
5	and chemical processes of O_3 formation and removal; (2) atmospheric modeling;
6	(3) background O ₃ concentrations; (4) monitoring techniques and networks; and
7	(5) ambient concentrations.

3.7.1 Physical and Chemical Processes

8	Ozone in the troposphere is a secondary pollutant; it is formed by photochemical
9	reactions of precursor gases and is not directly emitted from specific sources. Ozone
10	precursor gases originate from both anthropogenic and natural source categories. Ozone
11	attributed to anthropogenic sources is formed in the atmosphere by photochemical
12	reactions involving sunlight and precursor pollutants including VOCs, NO _X , and CO.
13	Ozone attributed to natural sources is formed through similar photochemical reactions
14	involving natural emissions of precursor pollutants from vegetation, microbes, animals,
15	biomass burning, lightning, and geogenic sources. The distinction between natural and
16	anthropogenic sources of O ₃ precursors is often difficult to make in practice, as human
17	activities affect directly or indirectly emissions from what would have been considered
18	natural sources during the preindustrial era. The formation of O ₃ , other oxidants, and
19	oxidation products from these precursors is a complex, nonlinear function of many
20	factors including: (1) the intensity and spectral distribution of sunlight reaching the lower
21	troposphere; (2) atmospheric mixing; (3) concentrations of precursors in the ambient air
22	and the rates of chemical reactions of these precursors; and (4) processing on cloud and
23	aerosol particles.
24	Ozone is present not only in polluted urban atmospheres but throughout the troposphere,
25	even in remote areas of the globe. The same basic processes involving sunlight-driven
24	

reactions of NO_x, VOCs and CO contribute to O₃ formation throughout the troposphere. These processes also lead to the formation of other photochemical products, such as PAN, nitric acid, and sulfuric acid, and to other compounds, such as formaldehyde and other carbonyl compounds. In urban areas, NO_x, VOCs and CO are all important for O₃ formation. In non-urban vegetated areas, biogenic VOCs emitted from vegetation tend to be the most important precursor to O₃ formation. In the remote troposphere, methane– structurally the simplest VOC–and CO are the main carbon-containing precursors to O₃

1	formation. Ozone is subsequently removed from the troposphere through a number of gas
2	phase reactions and deposition to surfaces.
3	Convective processes and turbulence transport O_3 and other pollutants both upward and
4	downward throughout the planetary boundary layer and the free troposphere. In many
5	areas of the U.S., O ₃ and its precursors can be transported over long distances, aided by
6	vertical mixing. The transport of pollutants downwind of major urban centers is
7	characterized by the development of urban plumes. Meteorological conditions, small-
8	scale circulation patterns, localized chemistry, and mountain barriers can influence
9	mixing on a smaller scale, resulting in frequent heterogeneous O ₃ concentrations across
10	individual urban areas.

3.7.2 Atmospheric Modeling

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CTMs have been widely used to compute the interactions among atmospheric pollutants and their transformation products, and the transport and deposition of pollutants. They have also been widely used to improve basic understanding of atmospheric chemical processes and to develop control strategies. The domains of CTMs extend from a few hundred kilometers on a side to the entire globe.

16 Most major regional (i.e., sub-continental) scale air-related modeling efforts at EPA rely 17 on the CMAQ modeling system. The horizontal domain for CMAQ typically extends 18 over North America with efforts underway to extend it over the entire Northern 19 Hemisphere. The upper boundary for CMAQ is typically set at 100 hPa, which is located 20 on average at an altitude of ~16 km. CMAQ is most often driven by the MM5 mesoscale 21 meteorological model, though it may be driven by other meteorological models including 22 the WRF model and the RAMS. Other major air quality systems used for regional scale 23 applications include CAMx and WRF/Chem.

24 Fine scale resolution is necessary to resolve features which can affect pollutant 25 concentrations such as urban heat island circulation; sea breezes; mountain and valley 26 breezes; and the nocturnal low-level jet. Horizontal domains are typically modeled by 27 nesting a finer grid model within a larger domain model of coarser resolution. Caution 28 must be exercised in using nested models because certain parameterizations like those for 29 convection might be valid on a relatively coarse grid scale but may not be valid on finer 30 scales and because incompatibilities can occur at the model boundaries. The use of finer 31 resolution in CTMs will require advanced parameterizations of meteorological processes 32 such as boundary layer fluxes, deep convection, and clouds, and necessitate finer-scale 33 inventories of land use, source locations, and emission inventories.

1	Because of the large number of chemical species and reactions that are involved in the
2	oxidation of realistic mixtures of anthropogenic and biogenic hydrocarbons, condensed
3	mechanisms must be used to simplify atmospheric models. These mechanisms can be
4	tested by comparison with smog chamber data. However, the existing chemical
5	mechanisms often neglect many important processes such as the formation and
6	subsequent reactions of long-lived carbonyl compounds, the incorporation of the most
7	recent information about intermediate compounds, and heterogeneous reactions involving
8	cloud droplets and aerosol particles. As a result, models such as CMAQ have had
9	difficulties with capturing the regional nature of O_3 episodes, in part because of
10	uncertainty in the chemical pathways converting NO_X to isoprene nitrates and recycling
11	of NO _X .

- Errors in photochemical modeling arise from meteorological, chemical, and emissions inputs to the model. Algorithms must be used for simulating meteorological processes that occur on spatial scales smaller than the model's grid spacing and for calculating the dependence of emissions on meteorology and time. Large uncertainties exist in the mechanism for oxidizing compounds of importance for atmospheric chemistry such as isoprene. Appreciable errors in emissions can occur if inappropriate assumptions are used in these parameterizations.
- 19The performance of CTMs must be evaluated by comparison with field data as part of a20cycle of model evaluations and subsequent improvements. Discrepancies between model21predictions and observations can be used to point out gaps in current understanding of22atmospheric chemistry and to spur improvements in parameterizations of atmospheric23chemical and physical processes.

3.7.3 Background Concentrations

- 24 Because the mean tropospheric lifetime of O_3 is on the order of a few weeks, O_3 can be 25 transported from continent to continent. The degree of influence from intercontinental 26 transport varies greatly by location and time. For instance, high elevation sites are most 27 susceptible to the intercontinental transport of pollution, particularly during spring. 28 However, because the atmospheric chemistry of O_3 is quite complex and can be highly 29 non-linear in environments close to sources of precursors, isolating the influence of 30 intercontinental transport of O_3 and O_3 precursors on urban air quality is particularly 31 problematic. 32 A number of recent studies indicate that natural sources such as wildfires and
- 33stratospheric intrusions and the intercontinental transport of pollution can significantly34affect O3 air quality in the United States. Two major modeling/field studies that focused

1	on discerning the contributions of Asian emissions to air quality in California were the
2	IONS-2010 and the CalNex studies conducted in May through June of 2010. Modeling
3	
	and observational components of these studies found evidence for substantive
4	contributions from stratospheric intrusions and Eurasian pollution to boundary layer O ₃ .
5	In particular, one modeling study found evidence of Asian contributions of 8 -15 ppb in
6	surface air during strong transport events in southern California. These contributions are
7	in addition to contributions from dominant local pollution sources. Their results suggest
8	that the influence of background sources on high O ₃ concentrations at the surface is not
9	always confined to high elevation sites. It is not clear to what extent the contributions
10	inferred by these studies are likely to be found in other years, during other seasons, or in
11	other locations. To gain a broader perspective and to isolate the influence of natural or
12	transported O ₃ , estimates from CTMs must be used. This is because observations within
13	the U.S.—even at relatively remote monitoring sites—are impacted by transport from
14	anthropogenic source regions within the U.S. borders.
15	In the context of a review of the NAAQS, it is useful to define background O_3
16	concentrations in a way that distinguishes between concentrations that result from
17	precursor emissions that are relatively less controllable from those that are relatively
18	more controllable through U.S. policies. For this assessment, three definitions of
19	background O_3 concentrations are considered, including (1) NA background (simulated
20	O_3 concentrations that would exist in the absence of anthropogenic emissions from the
21	U.S., Canada and Mexico), (2) U.S. background (simulated O_3 concentrations that would
22	exist in the absence of anthropogenic emissions from the U.S.), and (3) natural
23	background (simulated O_3 concentrations in the absence of all anthropogenic emissions
24	globally). Each definition of background O_3 includes contributions resulting from
25	emissions from natural sources (e.g., stratospheric intrusion, wildfires, biogenic methane
26	and more short-lived VOC emissions) throughout the globe. There is no chemical
27	difference between background O_3 and O_3 attributable to U.S. or North American
28	anthropogenic sources. However, to inform policy considerations regarding the current or
29	potential alternative standards, it is useful to understand how total O_3 concentrations can
	•
30	be attributed to different sources.
31	Since background O_3 concentrations as defined above are a construct that cannot be
32	directly measured, the range of background O ₃ concentrations is estimated using CTMs.
33	For the current assessment, recently published results from Zhang et al. (2011) using the
34	GEOS-Chem model at $0.5^{\circ} \times 0.667^{\circ}$ (~50 km × 50 km) horizontal resolution and Emery
35	et al. (2012) using a GEOS-Chem/CAMx model (hereafter referred to as CAMx) at finer
36	horizontal resolution (12 km \times 12 km) were used. Results from these models represent
37	the latest estimates for background O3 concentrations documented in the peer-reviewed
38	literature.

1	The main results from these modeling efforts can be summarized as follows. Simulated
2	regional and seasonal means of base-case O ₃ using both models generally agree to within
3	a few pbb with observations for most of the U.S. However, neither model is currently
4	capable of simulating day specific base-case O_3 concentrations within reasonable bounds.
5	Both models show background concentrations vary spatially and temporally. NA
6	background concentrations are generally higher in spring than in summer across the U.S.
7	Simulated mean NA background concentrations are highest in the Intermountain West
8	(i.e., at high altitude) in spring and in the Southwest in summer. Lowest estimates of NA
9	background occur in the East in the spring and the Northeast in summer. NA background
10	concentrations tend to increase with total (i.e., base case) O ₃ concentrations at high
11	elevation, but that tendency is not as clear at low elevations. Comparison of NA
12	background and natural background indicate that methane is a major contributor to NA
13	background O ₃ , accounting for slightly less than half of the increase in background since
14	the pre-industrial era; and whose relative contribution is projected to grow in the future.
15	U.S. background concentrations are on average 2.6 ppb higher than NA background
16	concentrations during spring and 2.7 ppb during summer across the U.S. with highest
17	increases above NA background over the Northern Tier of New York State (19.1 ppb
18	higher than NA background) in summer. High values for U.S. background are also found
19	in other areas bordering Canada and Mexico. Contributions to background O ₃ can be
20	episodic or non-episodic; high background concentrations are driven primarily by the
21	episodic events such as stratospheric intrusions and wildfires. The most pronounced
22	differences between these model results and observations are at the upper end of the
23	concentration distribution, particularly at high elevations. In general, these model
24	simulations provide a consistent representation of average background concentrations
25	over seasons and broad spatial areas, but are not able to capture background
26	concentrations at finer spatial (i.e., urban) and temporal (i.e., specific day) scales.
27	Note that the calculations of background concentrations presented in this chapter were
28	formulated to answer the question, "what would O3 concentrations be if there were no
29	anthropogenic sources". This is different from asking, "how much of the O3 measured or
30	simulated in a given area is due to background contributions". Because of potentially
31	strong non-linearities—particularly in many urban areas—these estimates by themselves
32	should not be used to answer the second question posed above. The extent of these non-
33	linearities will generally depend on location and time, the strength of concentrated
34	sources, and the nature of the chemical regime. Further work is needed on how these
35	estimates of background concentrations can be used to help determine the contributions
26	

36 of background sources of O_3 to urban concentrations.

3.7.4 Monitoring

1 2 3 4 5	The FRM for O_3 measurement is the CLM and is based on the detection of chemiluminescence resulting from the reaction of O_3 with ethylene gas. Almost all of the SLAMS that reported data to AQS from 2005 to 2009 used UV absorption photometer FEMs and greater than 96% of O_3 monitors met precision and bias goals during this period.
6 7 8 9 10	State and local monitoring agencies operate O_3 monitors at various locations depending on the area size and typical peak concentrations (expressed in percentages below, or near the O_3 NAAQS). SLAMS make up the ambient air quality monitoring sites that are primarily needed for NAAQS comparisons and include PAMS, NCore, and all other State or locally-operated stations except for the monitors designated as SPMs.
11 12 13 14 15 16 17 18 19	In 2010, there were 1250 SLAMS O ₃ monitors reporting values to the EPA AQS database. Since O ₃ levels decrease appreciably in the colder parts of the year in many areas, O ₃ is required to be monitored at SLAMS monitoring sites only during the "ozone season" which varies by state. PAMS provides more comprehensive data on O ₃ in areas classified as serious, severe, or extreme nonattainment for O ₃ . There were a total of 119 PAMS reporting values to the EPA AQS database in 2009. NCore is a new multipollutant monitoring network currently being implemented to meet multiple monitoring objectives. Each state is required to operate at least one NCore site and the network will consist of about 60 urban and 20 rural sites nationwide.
20 21 22 23 24 25 26 27 28	CASTNET is a regional monitoring network established to assess trends in acidic deposition and also provides concentration measurements of O ₃ . CASTNET O ₃ monitors operate year round and are primarily located in rural areas. At the beginning of 2010, there were 80 CASTNET sites located in, or near, rural areas. The NPS also operates a POMS network. The POMS couples the small, low-power O ₃ monitor with a data logger, meteorological measurements, and solar power in a self contained system for monitoring in remote locations. Twenty NPS POMS reported O ₃ data to AQS in 2010. A map of the current and proposed rural NCore sites, along with the CASTNET, and the NPS POMS sites was shown in Figure 3-22.
29 30 31 32 33 34 35	Satellite observations for O_3 are growing as a resource for many purposes, including model evaluation, assessing emissions reductions, pollutant transport, and air quality management. Satellite retrievals are conducted using the solar backscatter or thermal infrared emission spectra and a variety of algorithms. Most satellite measurement systems have been developed for measurement of the total O_3 column. Mathematical techniques have been developed and must be applied to derive information from these systems about tropospheric O_3 .

3.7.5 Ambient Concentrations

1 2 3	Ozone is the only photochemical oxidant other than NO_2 that is routinely monitored and for which a comprehensive database exists. Other photochemical oxidants are typically only measured during special field studies. Therefore, the concentration analyses
4 5	contained in this chapter have been limited to widely available O_3 data obtained directly from AQS for the period from 2007 to 2009.
6 7	The median 24-h avg, 8-h daily max, and 1-h daily max O_3 concentrations across all U.S. sites reporting data to AQS between 2007 and 2009 were 29, 40, and 44 ppb,
8 9	respectively. Representing the upper end of the distribution, the 99th percentiles of these same metrics across all sites were 60, 80, and 94 ppb, respectively.
10	To investigate urban-scale O ₃ variability, 20 focus cities were selected for closer analysis;
11	these cities were selected based on their importance in O ₃ epidemiologic studies and on
12	their geographic distribution across the U.S. Several of these cities had relatively little
13	spatial variability in 8-h daily max O ₃ concentrations (e.g., inter-monitor correlations
14	ranging from 0.61 to 0.96 in Atlanta) while other cities exhibited considerably more
15	variability in O_3 concentrations (e.g., inter-monitor correlations ranging from -0.06 to
16	0.97 for Los Angeles). The negative and near-zero correlations in Los Angeles were
17	between monitors with a relatively large separation distance (>150 km), but even some of
18	the closer monitor pairs were not very highly correlated. Similar to the correlation, the
19	coefficient of divergence was found to be highly dependent on the urban area under
20	investigation. As a result, caution should be observed in using data from a sparse network
21	of ambient O ₃ monitors to approximate community-scale exposures.
22	To investigate rural-focused O3 variability using AQS data, all monitors located within
23	six rural monitoring areas were examined. These rural monitoring sites are impacted by
24	transport of O ₃ or O ₃ precursors from upwind urban areas, and by local anthropogenic
25	emissions within the rural areas such as emissions from motor vehicles, power
26	generation, biomass combustion, or oil and gas operations. As a result, monitoring data
27	from these rural locations are not unaffected by anthropogenic emissions. The rural area
28	investigated with the largest number of available AQS monitors was Great Smoky
29	Mountain National Park in NC and TN where the median warm-season 8-h daily max O ₃
30	concentration ranged from 47 ppb at the lowest elevation site (elevation = 564 meters;
31	site ID = 470090102) to 60 ppb at the highest elevation site (elevation = $2,021$ meters;
32	site ID = 471550102), with correlations between the 5 sites ranging from 0.78 to 0.92 and
33	CODs ranging from 0.04 to 0.16. A host of factors may contribute to variations observed
34	at these rural sites, including proximity to local O ₃ precursor emissions, variations in
35	boundary-layer influences, meteorology and stratospheric intrusion as a function of

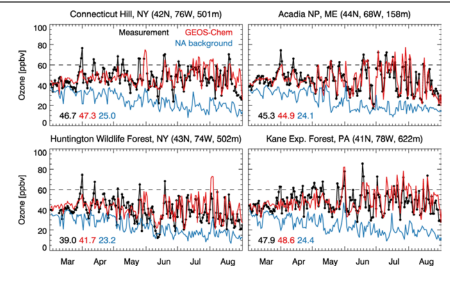
1	elevation, and differences in wind patterns and transport behavior due to local
2	topography.
3	Since O_3 produced from emissions in urban areas is transported to more rural downwind
4	locations, elevated O_3 concentrations can occur at considerable distances from urban
5	centers. In addition, major sources of O ₃ precursors such as highways, power plants,
6	biomass combustion, and oil and gas operations are commonly found in rural areas,
7	adding to the O_3 in these areas. Due to lower chemical scavenging in non-urban areas, O_3
8	tends to persist longer in rural than in urban areas which tends to lead to higher
9	cumulative exposures in rural areas influenced by anthropogenic precursor emissions.
10	The persistently high O ₃ concentrations observed at many of these rural sites investigated
11	here indicate that cumulative exposures for humans and vegetation in rural areas can be
12	substantial and often higher than cumulative exposures to O_3 in urban areas.
13	Nation-wide surface level O ₃ concentrations in the U.S. have declined gradually over the
14	last decade. A noticeable decrease in O_3 concentrations between 2003 and 2004,
15	particularly in the eastern U.S., coincided with NO_X emissions reductions resulting from
16	implementation of the NO_X SIP Call rule, which began in 2003 and was fully
17	implemented in 2004. This rule was designed to reduce NO_X emissions from power
18	plants and other large combustion sources in the eastern U.S. Downward trends in O_3
19	concentrations in the western U.S. have not been as substantial and several individual
20	monitors have reported increases in O_3 concentrations when 2001-2003 design values are
21	compared with 2008-2010 design values. In contrast to the downward regional trends in
22	surface-level O_3 concentrations in the U.S., global scale observations have indicated a
23	general rise in O_3 by a factor of 2 or more since pre-industrial times, as discussed in
24	Chapter <u>10</u> , Section <u>10.3.3.1</u> . Several observational studies investigating O_3
25	concentrations in the marine layer off the Pacific Coast of the U.S. have reported a steady
26	rise in O ₃ concentrations over the last few decades.
27	Urban O_3 concentrations show a strong degree of diel variability resulting from daily
28	patterns in temperature, sunlight, and precursor emissions. Other factors, such as the
29	relative importance of transport versus local photochemical production and loss rates, the
30	timing for entrainment of air from the nocturnal residual boundary layer, and the diurnal
31	variability in mixing layer height also play a role in daily O_3 patterns. Urban diel
32	variations investigated in this assessment show no substantial change in patterns since the
33	2006 O_3 AQCD (U.S. EPA, 2006b). The 1-h max concentrations tend to occur in mid-
34	afternoon and 1-h min concentrations tend to occur in early morning, with more
35	pronounced peaks in the warm months relative to the cold months. There is city-to-city
36	variability in these times, however, and caution is raised in extrapolating results from one
37	city to another in determining the time of day for O_3 maxima and minima.

1	Rural O_3 concentrations show a varying degree of diel variability depending on their
2	location relative to larger urban areas. Three rural areas investigated in the east showed
3	relatively little diel variability, reflecting the regional nature of O ₃ in the east. In contrast,
4	three rural areas investigated in the west did display diel variability resulting from their
5	proximity to fresh urban emissions. These six areas investigated were selected as
6	illustrative examples and do not represent all rural areas in the U.S.
7	Since O ₃ is a secondary pollutant formed in the atmosphere from precursor emissions, its
8	correlation with primary pollutants such as CO and NO_X can vary substantially by
9	location. Furthermore, O ₃ formation is strongly influenced by meteorology, entrainment,
10	and transport of both O_3 and O_3 precursors, resulting in a broad range in correlations with
11	other pollutants which can vary substantially with season. In the co-pollutant analyses
12	shown in Figure 3-56, the year-round 8-h daily max O_3 data exhibited a very wide range
13	in correlations with all the criteria pollutants. A clearer pattern emerged when the data are
14	stratified by season with mostly negative correlations in the winter and mostly positive
15	correlations in the summer for all co-pollutants. The median seasonal correlations are
16	modest at best with the highest positive correlation at 0.52 for $PM_{2.5}$ in the summer and
17	the highest negative correlation at -0.38 for PM _{2.5} in the winter. Therefore, statistical
18	analyses that may be sensitive to correlations between co-pollutants need to take
19	seasonality into consideration, particularly when O ₃ is being investigated.

3.8 Supplemental Information on Ozone Model Predictions

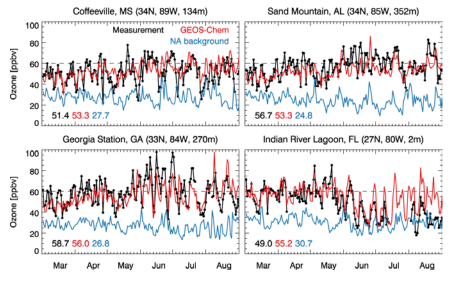
20	This section contains supplemental comparisons between GEOS-Chem simulations of
21	MDA8 O_3 concentrations with observations for 2006 from <u>Zhang et al. (2011</u>) and <u>Emery</u>
22	et al. (2012). Further details on these simulations can be found in Section $3.4.3$.
23	Figure 3-58 through Figure 3-64 show GEOS-Chem predictions for the base model
24	(i.e., model including all anthropogenic and natural sources; labeled as GEOS-Chem in
25	the figure) and the NA background model (i.e., model including natural sources
26	everywhere in the world and anthropogenic sources outside the U.S., Canada, and
27	Mexico; labeled as NA background in the figure) along with measurements obtained
28	from selected CASTNET sites (labeled as Measurement in the figure). Figure 3-65 shows
29	a comparison of GEOS-Chem output with measurements at Mt. Bachelor, OR from
30	March-August, 2006. Figure 3-66 shows a comparison of vertical profiles (mean ± 1
31	standard deviation) calculated by GEOS-Chem with ozonesondes launched at Trinidad
32	Head, CA and Boulder, CO. Figure 3-67 and Figure 3-68 show a comparison of AM3
33	simulations of individual stratospheric intrusions during May-June 2010. Figure 3-69
34	through Figure 3-74 show box plots for measurements at CASTNET sites, GEOS-Chem
35	predictions from Zhang et al. (2011) and CAMx predictions from Emery et al. (2012) for

both the base case and NA background. Figure 3-75 shows time series of AM3 simulations at approximately $2^{\circ} \times 2^{\circ}$ at Gothic CO for 2006.



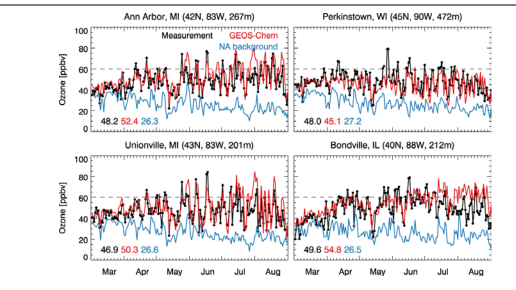
Source: Zhang et al. (2011).

Figure 3-58 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Northeast with GEOS-Chem predictions for the base case and for the North American background case during March-August, 2006.



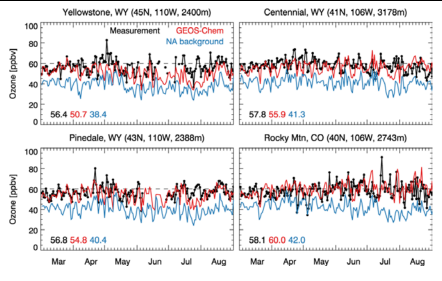
Source: Zhang et al. (2011).

Figure 3-59 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Southeast with GEOS-Chem predictions for the base case and for the North American background case during March-August, 2006.



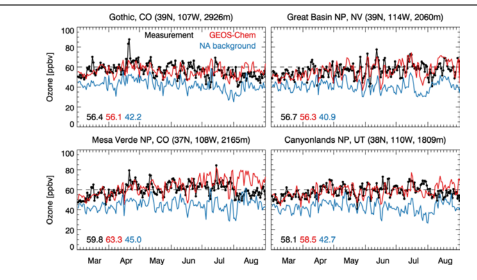
Source: Zhang et al. (2011).

Figure 3-60 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Upper Midwest with GEOS-Chem predictions for the base case and for the North American background case during March-August, 2006.



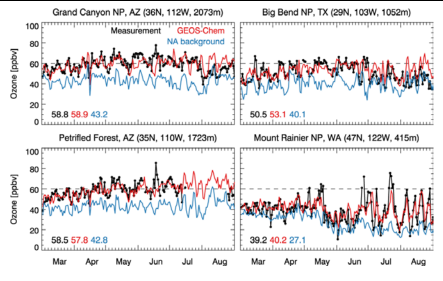
Source: Zhang et al. (2011).

Figure 3-61 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Intermountain West with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.



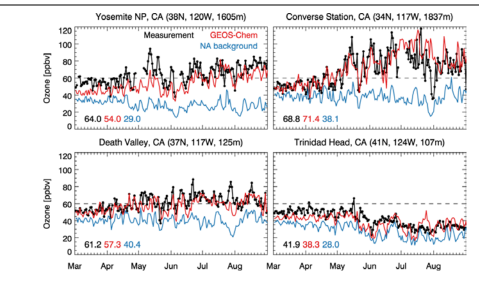
Source: Zhang et al. (2011).

Figure 3-62 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the Intermountain West with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.



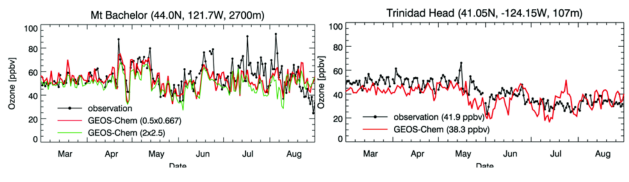
Source: Zhang et al. (2011).

Figure 3-63 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at four CASTNET sites in the West with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.



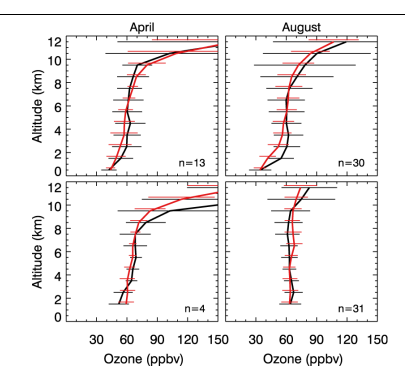
Source: Zhang et al. (2011).

Figure 3-64 Comparison of time series of measurements of daily maximum 8-hour average ozone concentrations at three CASTNET sites and the Trinidad Head site in California with GEOS-Chem predictions for the base case and the North American background case during March-August, 2006.



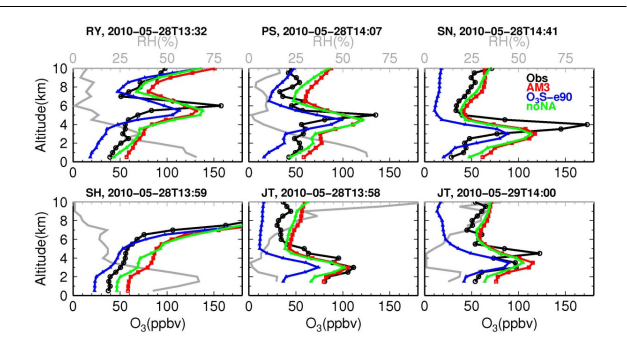
Source: Zhang et al. (2011).

Figure 3-65 Comparison of daily maximum 8-h average ozone predicted using GEOS-Chem at 0.5° × 0.667° (and 2° × 2.5° resolution; left figure only) with measurements at Mount Bachelor, OR (left); and at Trinidad Head, CA (right) from March to August 2006.



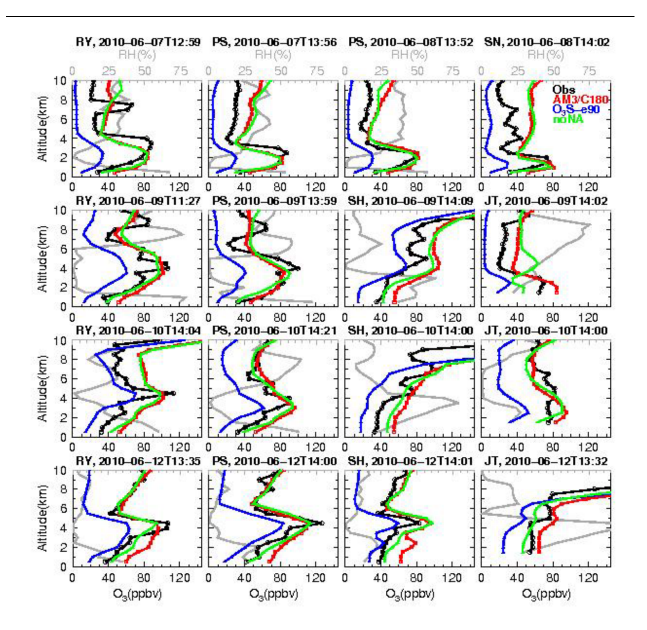
Note: The letter 'n' refers to the number of ozonesonde profiles, and the model was sampled on the same days as the ozonesonde launches. As can be seen from the figure, variability in both model and measurements increases with altitude, but variability in the model results is much smaller at high altitudes than seen in the observations at both sites. Source: Zhang et al. (2011).

Figure 3-66 Comparison of monthly mean (± 1 standard deviation) ozone calculated GEOS-Chem (in red) with ozonesondes (in black) at Trinidad Head, CA (top) and Boulder, CO (bottom) during April and August 2006.



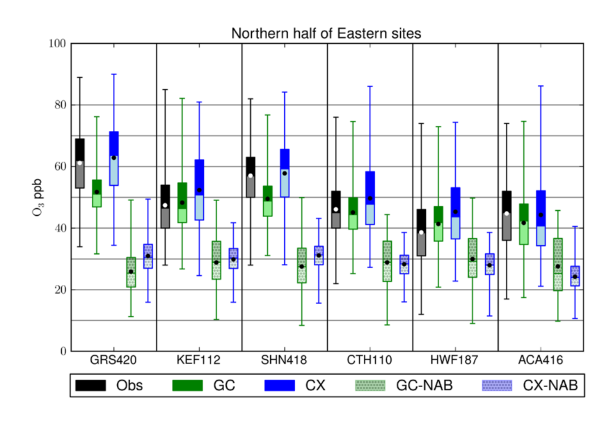
Note: Shows ozone profiles at multiple sites as observed (black) by ozonesondes and simulated (red) by the GFDL AM3 model at ~50 × 50 km resolution. Also shown are observed relative humidity (gray) and AM3 estimates of ozone concentrations in the absence of North American anthropogenic emissions (green) and the stratospheric contribution (blue). Model results have been interpolated to sonde pressure and averaged over 0.5-km altitude bins.

Figure 3-67 A deep stratospheric ozone intrusion over California on May 28-29, 2010.



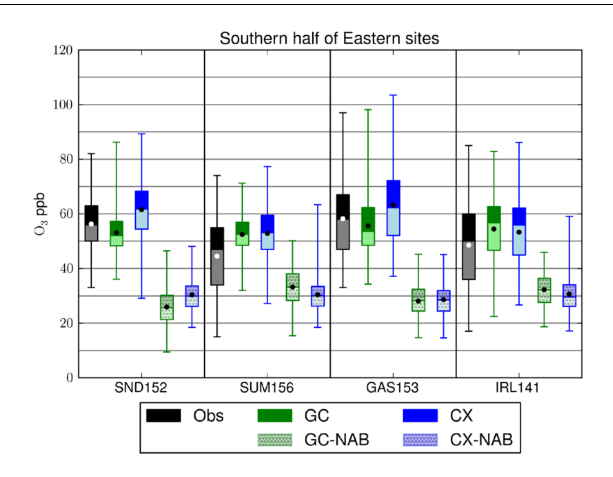
Note: Shows ozone profiles at multiple sites as observed (black) by ozonesondes and simulated (red) by the GFDL AM3 model at ~50 × 50 km resolution. Also shown are observed relative humidity (gray) and AM3 estimates of ozone concentrations in the absence of North American anthropogenic emissions (green) and the stratospheric contribution (blue). Model results have been interpolated to sonde pressure and averaged over 0.5-km altitude bins.

Figure 3-68 A deep stratospheric ozone intrusion over California on June 7-12, 2010.



Note: Stippled boxes indicate North American background. GRS = Great Smoky NP; KEF = Kane Exp. Forest; SHN = Shenandoah NP; CTH = Connecticut Hill; HWF = Huntington Wildlife Forest; ACA = Acadia NP. Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

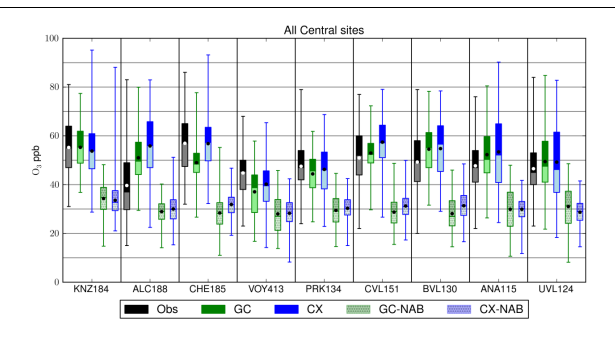
Figure 3-69 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Northeast and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



Note: Stippled boxes indicate North American background. SND = Sand Mountain; SUM = Sumatra; GAS = Georgia Station; IRL = Indian River Lagoon.

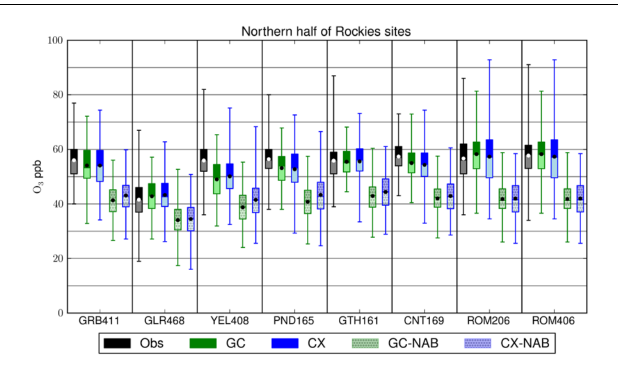
Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

Figure 3-70 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Southeast and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



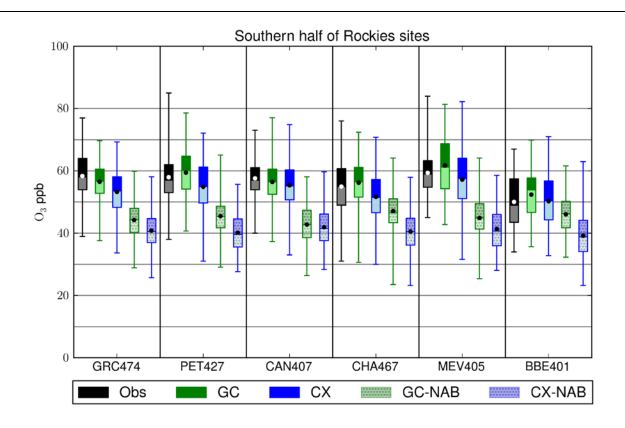
Note: Stippled boxes indicate North American background. KNZ = Konza Prairie; ALC = Alabama-Coushatta; CHE = Cherokee Nation; VOY = Voyageurs NP; PRK = Perkinstown; CVL = Coffeeville; BVL = Bondsville; ANA = Ann Arbor; UVL = Unionville. Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

Figure 3-71 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Central U.S. and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



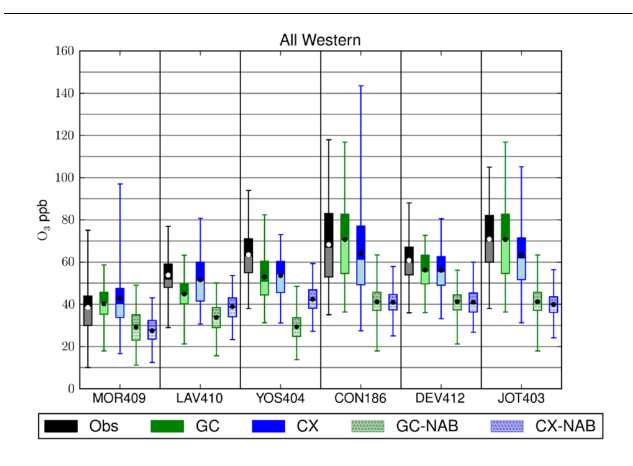
Note: Stippled boxes indicate North American background. GRB = Great Basin NP; GLR = Glacier NP; YEL = Yellowstone NP; PND = Pinedale; GTH = Gothic; CNT = Centennial; ROM = Rocky Mountain NP (co-located sites). Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

Figure 3-72 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Northern Rockies and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



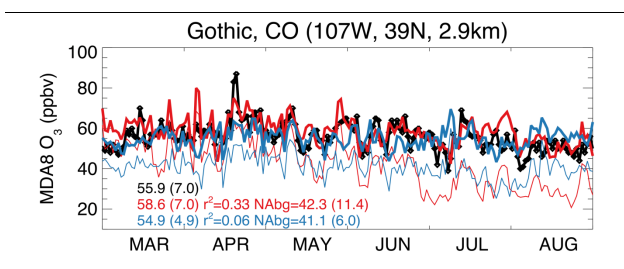
Note: Stippled boxes indicate North American background. GRC = Grand Canyon NP; PET = Petrified Forest; CAN = Canyonlands NP; CHA = Chiracahua NM; MEV = Mesa Verde NP; BBE = Big Bend NP. Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

Figure 3-73 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the Southern Rockies and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



Note: Stippled boxes indicate North American background. MOR = Mount Ranier NP; LAV = Lassen Volcanic NP; YOS = Yosemite NP; CON = Converse Station; DEV = Death Valley NM; JOT = Joshua Tree NM. Source: Adapted from Emery et al. (2012) and Zhang et al. (2011).

Figure 3-74 Box plots showing maximum, interquartile range and minimum ozone concentrations measured at CASTNET sites (black) in the West and predictions from GEOS-Chem at ~50 × 50 km resolution (green) and CAMx at 12 × 12 km resolution (blue) for May-August 2006.



Note: Observed (black) and simulated by the GEOS-Chem (blue; horizontal resolution is $0.5^{\circ} \times 0.667^{\circ}$) and AM3 (red; horizontal resolution is approximately $2^{\circ} \times 2^{\circ}$) global models. Also shown are the model estimates for North American background (thin lines); the spike in mid-April likely corresponds to a stratospheric intrusion. The model correlations with observations, average (over the entire March through August period) total O₃ and North American background (NAbg) O₃ estimates, and their standard deviations (shown in parentheses) are presented in the lower left.

Figure 3-75 Daily maximum 8-hour average (MDA8) ozone in surface air at Gothic, CO for March through August 2006.

3.9 Supplemental Figures of Observed Ambient Ozone Concentrations

3.9.1 Ozone Monitor Maps for the Urban Focus Cities

1	This section contains supplemental maps showing the location of O ₃ monitors reporting
2	to AQS for each of the 20 urban focus cities introduced in Section 3.6.2.1. The monitors
3	are delineated in the maps as year-round or warm-season based on their inclusion in the
4	year-round data set and the warm-season data set discussed in Section 3.6.2.1. The maps
5	also include the CSA/CBSA boundary selected for monitor inclusion, the location of
6	urban areas and water bodies, the major roadway network, as well as the population
7	gravity center based on the entire CSA/CBSA and the individual focus city boundaries.
8	Population gravity center is calculated from the average longitude and latitude values for
9	the input census tract centroids and represents the mean center of the population in a
10	given area. Census tract centroids are weighted by their population during this
11	calculation.

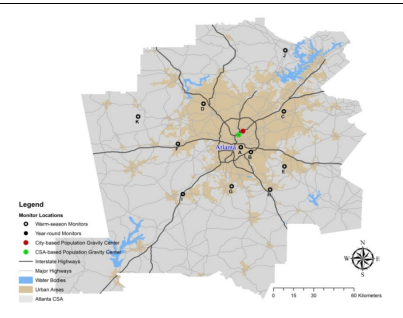


Figure 3-76 Map of the Atlanta CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

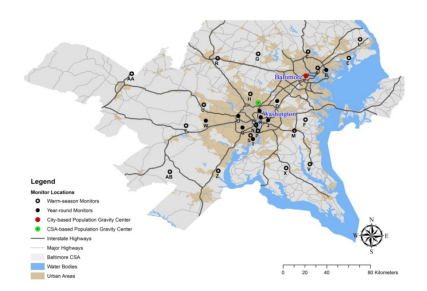


Figure 3-77 Map of the Baltimore CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

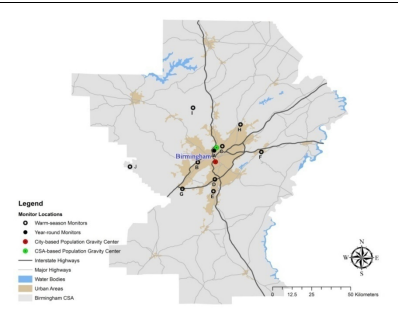


Figure 3-78 Map of the Birmingham CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

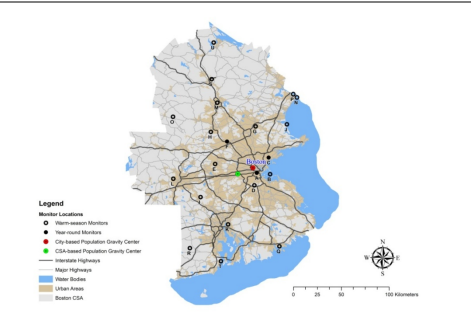


Figure 3-79 Map of the Boston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

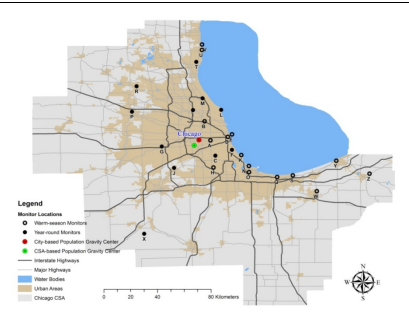


Figure 3-80 Map of the Chicago CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

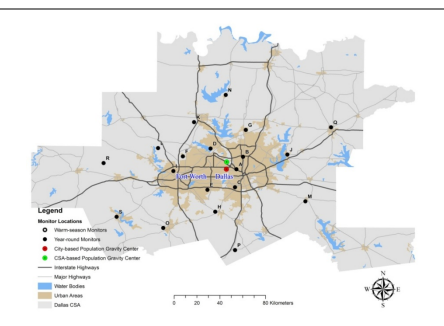


Figure 3-81 Map of the Dallas CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

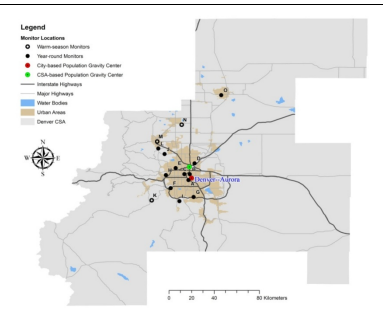


Figure 3-82 Map of the Denver CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

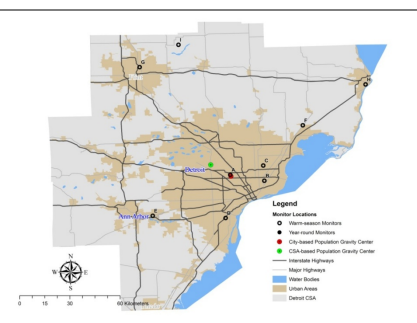


Figure 3-83 Map of the Detroit CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

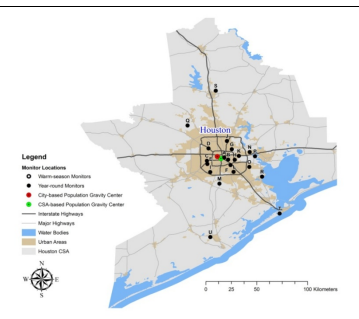


Figure 3-84 Map of the Houston CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

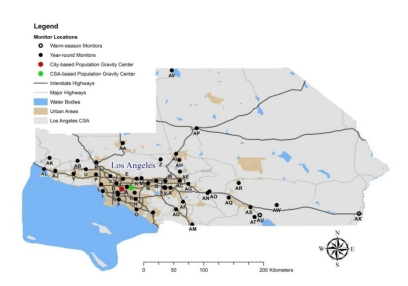


Figure 3-85 Map of the Los Angeles CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

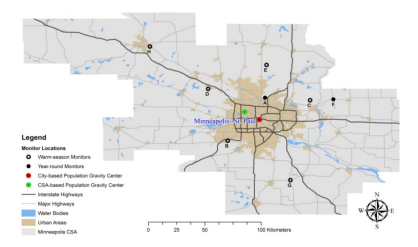


Figure 3-86 Map of the Minneapolis CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

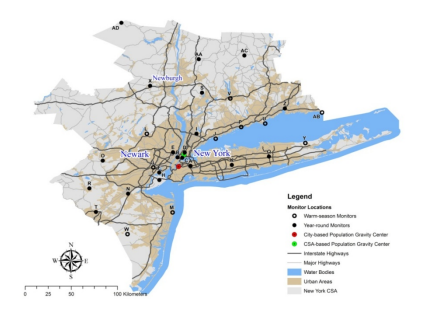


Figure 3-87 Map of the New York CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

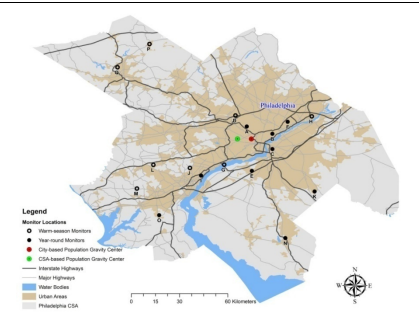


Figure 3-88 Map of the Philadelphia CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

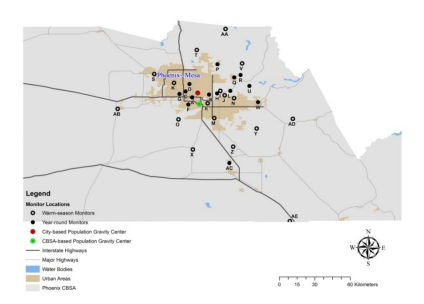


Figure 3-89 Map of the Phoenix CBSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

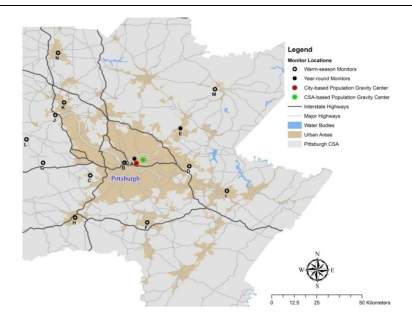


Figure 3-90 Map of the Pittsburgh CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

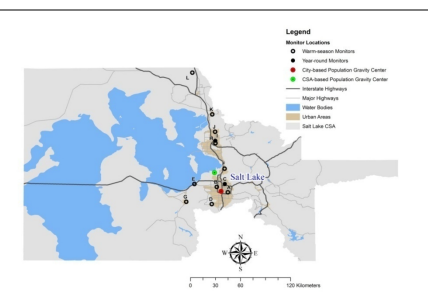


Figure 3-91 Map of the Salt Lake City CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

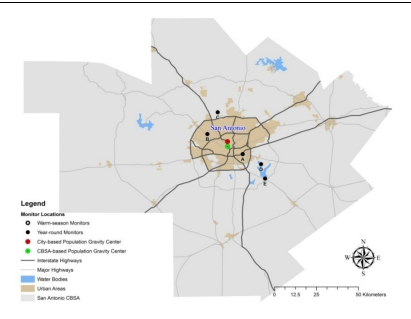


Figure 3-92 Map of the San Antonio CBSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

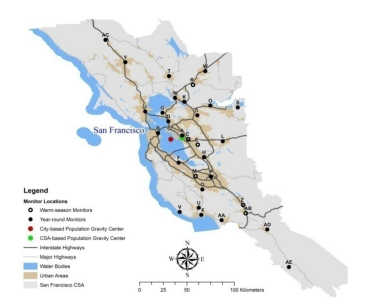


Figure 3-93 Map of the San Francisco CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

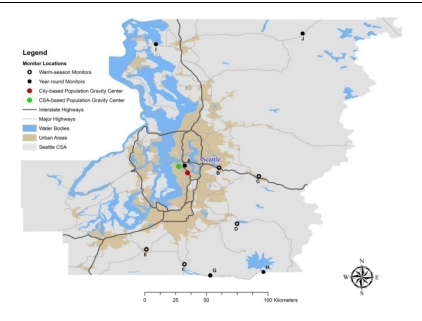


Figure 3-94 Map of the Seattle CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

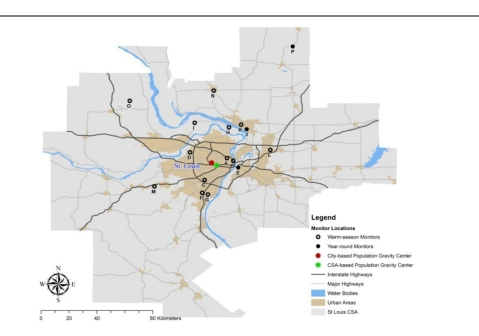


Figure 3-95 Map of the St. Louis CSA including ozone monitor locations, population gravity centers, urban areas, and major roadways.

3.9.2 Ozone Concentration Box Plots for the Urban Focus Cities

1	This section contains box plots depicting the distribution of 2007-2009 warm-season 8-h
2	daily max O ₃ data from each individual monitor in the 20 urban focus cities introduced in
3	Section $3.6.2.1$. Monitor information including the AQS site id, the years containing
4	qualifying data between 2007 and 2009, and the number of 8-h daily max O_3
5	observations included in the data set are listed next to the box plot. Statistics including
6	the mean, standard deviation (SD), median and inner quartile range (IQR) are also shown
7	for each monitor with the site letter corresponding to the sites listed in the figures above.

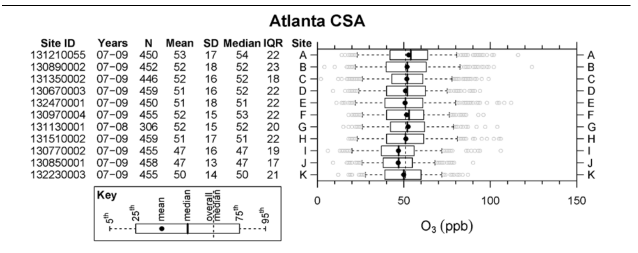


Figure 3-96 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Atlanta CSA.

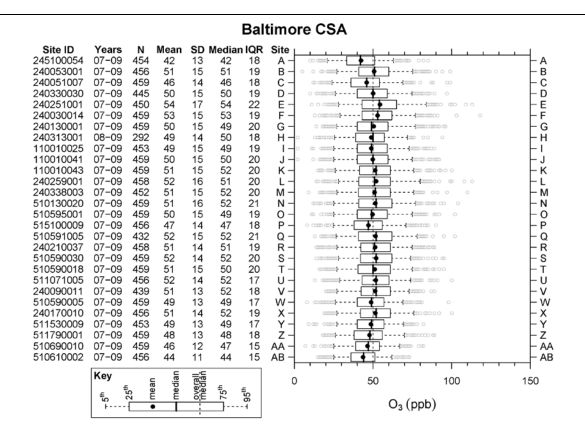


Figure 3-97 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Baltimore CSA.

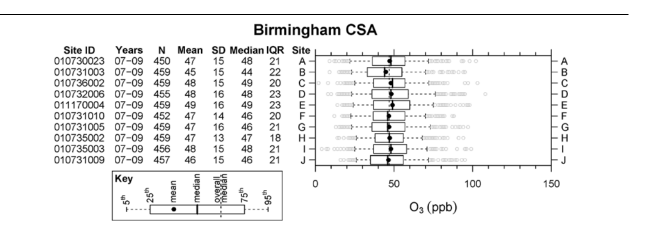


Figure 3-98 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Birmingham CSA.

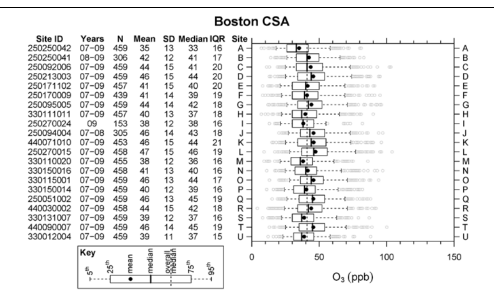


Figure 3-99 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Boston CSA.

						Ch	icag	o CSA				
Site ID	Years	Ν	Mean	SD	Media	n IOR	Site				1	
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170310076	07-09	458	44	14	44	18	č-	0101001	÷			⊢ č
170310042	07-09	412	45	14	44	17	Ď –	a ama	· •	- 10000 0	0	⊢ Ď
170310072	07-09	459	42	12	42	17	Е –	o commit	÷;	00 0000000		ΗE
170310064	07-09	459	41	13	40	18	F	o com;[• • • • • •			⊢ F
170436001	07-09	459	39	12	39	16	G –	annt	.	0 0000		⊢G
170310001	07-09	459	46	14	46	19	н –	o anamati	:•	- impan a		⊢н
170314007	07-09	458	39	13	37	18	1-	00000	•			<u>⊢</u> 1
170311601	07-09	457	48	14	47	19	J –		• •	4 mano a	0 0	- J
170310032	07-09	450	46	13	45	17	к-		:	0 000 0000		⊢ĸ
170317002	07-09	450	43	13	42	17	L-	amami	• • • • • •			⊢L
170314201	07-09	611	42	13	41	17	м –	0 000 1	• · · · ·	-		- M
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180892008	07-09	451	45	13	44	18	0 -	o amani <mark></mark>	:• ···	• 0 000000		⊢o
170890005	07-09	459	44	13	42	16	Р –	commit•••••	•	• cancant•		⊢ P
180890022	07-09	455	42	13	41	15	Q –	0000000	• · · · · ·			⊢Q
171110001	07-09	458	43	12	42	15	R –		• • • • • •			⊢ R
181270024	07-09	456	46	14	44	17	S –		_ ! •			- s
170971002	07-09	459	41	13	39	18	Т –	o annuit	· · · · ·	100 CO (10		⊢ T
170971007	07-09	459	46	13	46	18	U –	o amp[÷.	- param on		- U
550590019	07-09	457	47	14	45	19	V -	· · · · · · · · · · · ·	••••••			⊢v.
181270026	07-09	453	43	13	42	18	w-	• • • • • • • •	▶ ····	• 1 000 0 000		⊢.w
171971011	07-09	458	42	12	41	15	X -		P	100000		⊢×
180910005	07-09	453	42	12	41	15	Y -	· · · · ·	P i			ŀΥ
180910010	07-09	456	44	13	43	17	Z –		₽┤╌╴	-monomp o		Z
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	1			+		'			```	3 (PPD)		

Figure 3-100 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Chicago CSA.

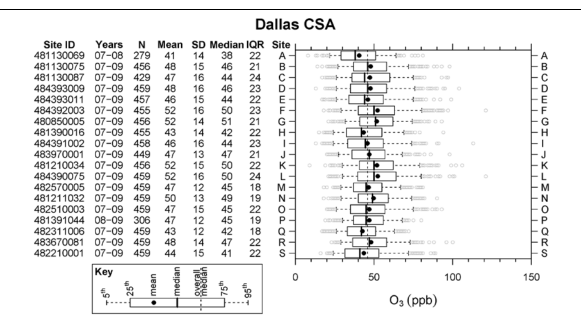


Figure 3-101 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Dallas CSA.

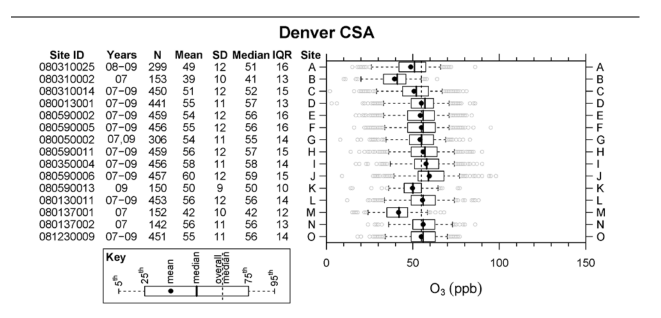


Figure 3-102 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Denver CSA.

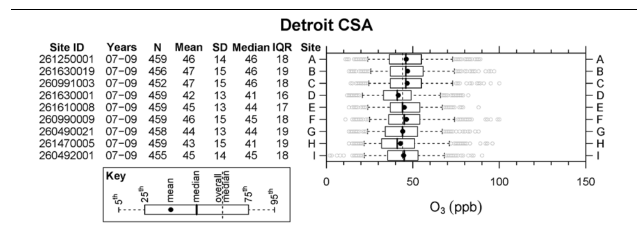


Figure 3-103 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Detroit CSA.

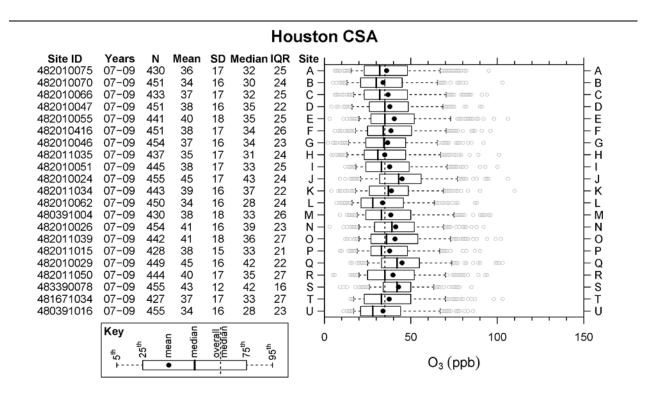


Figure 3-104 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Houston CSA.

					L	.os /	Ange	eles CSA	
Site ID	Years	Ν	Mean	SD	Media	n IQR	Site		
060371602	07-09	458	48	13	47	17	A –		- A
060371301	07-08	306	36	9	34	10	В –		- B
060371302	09	152	44	10	44	12	<u>c</u> –		⊢ c
060371103	07-09	457	46	12	45	14	_ <u>P</u> –		ΗP
060372005 060374002	07-09 07-09	459 459	54 38	15 10	53 37	18	E – F –		⊢E ⊢F
060595001	07-09	459	50	12	49	11 14	Ġ-		ĒĞ
060590007	07-09	459	48	10	47	12	н-		– H
060375005	07-09	459	45	9	45	12	i-		⊢ï
060371002	07-09	459	56	14	55	19	j —	oo aca:	⊢ j
060370002	07-09	459	57	17	56	22	к –	o coco ampi	-κ
060370113	07-09	459	48	10	47	13	L –		-L
060370016	07-09	458	64	18	63	23	М –	an annan	- M
060371701	07-09	459	61	16	60	20	N –	@ @@C@ 	- N
060591003	07-09	459	45	9	44	12	<u>o</u> –		μo
060371201	07-09	459	61	14	60	19	P -		⊢ P
060711004	07-09 07-09	457	66	19	66	23	Q -		- Q
060376012 060650004	07-09	457 127	68 69	18 18	69 65	27 23	R – S –		- R - S
060592022	07-09	457	52	13	50	15	т –	o come[●]tamp@cood	- T
061112002	07-09	455	62	12	62	16	- u –		Ļύ
060658005	08-09	276	65	15	64	18	v –	o occurrente in the second sec	ĻΫ
060712002	07-09	459	68	19	67	24	w-	o	⊢ w
060658001	07-09	440	69	16	68	18	Х –	oo oom+tanoo oo o	-x
061110007	07-09	459	54	10	54	12	Y –		- Υ
060710012	07-09	456	67	13	67	18	Z –	a caana; mana	– Z
060379033	07-09	452	67	13	66	19	AA -		- AA
061110009	07-09	458	58	11	58	14	AB -		- AB
060719004	07-09	457	70	19	70 67	26	AC -		
060659001 060710005	07-09 07-09	453 459	68 79	16 19	67 80	21 28	AD - AE -		⊢ AD ⊢ AE
060656001	07-09	459	72	17	73	24	AF -		
060714003	07-09	459	73	18	73	25	AG -		- AG
060714001	07-09	455	68	14	68	21	AH -	ommmi +mmm m o	- AH
060710306	07-09	459	64	12	64	17	AI –	o maan (- AI
061113001	07-09	453	44	9	43	11	AJ –	0 000 mmm	– AJ
061111004	07-09	458	57	11	57	14	AK –	• • • • • • • • • • • • • • • • • • •	– AK
061112003	07-09	457	41	9	40	12	AL –		– AL
060650009	09	153	22	8	20	8	AM -		- AM
060650012	07-09	457	73	15	71	22	AN -		- AN
060651016	07-09	459	73 61	16 11	73	23 15		o ∞ ∞ ∞ ∞∞m+ 4 <u>;</u> ∞ ∞ ∞∞m+ - ! 4	– AO – AP
060710001 060655001	07-09 07-09	455 459	61 69	11 14	60 68	15 21	AP – AQ –		– AP – AQ
060719002	07-09	452	73	13	73	18	AQ -		- AQ
060652002	07-09	448	62	13	61	18	AS -		- AS
060651999	07-08	283	49	17	50	22	AT-		- AT
060651010	09	153	59	10	59	15	AU -	(m)	- AU
060711234	07-09	453	59	10	58	13	AV –		- AV
060650008	07,09	265	58	10	57	14	AW –		– AW
060659003	07-09	444	42	10	42	13	AX –		
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Figure 3-105 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Los Angeles CSA.

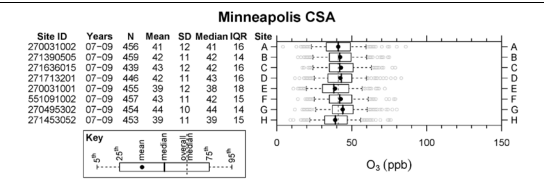


Figure 3-106 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Minneapolis CSA.

New York CSA											
Site ID	Years	Ν	Mean	SD	Mediar	IQR	Site				
360810124	07-09	446	43	15	41	21	A -	0 0000		mm::::::::::::::::::::::::::::::::	
360610135	08-09	298	39	15	38	19	в –	com amp	Þ :		- B
360050110	07-09	457	40	15	39	19	ē-l		•		⊢ē
360050133	07-09	459	41	14	39	18	Ď –		•		- D
340030006	08-09	300	42	15	41	20	E –	com:	A		- E
340170006	07-09	442	45	17	43	20	F				- F
340130003	09	122	36	14	36	19	G –		· · · · · ·	10000	- G
360850067	07,09	298	45	16	43	23	й –				μĤ
361192004	07-09	444	46	17	44	22	1-	o oams[•		- 1
090010017	07-09	447	49	15	47	20	1 – I	o cmmm			- J
361030002	07-09	454	47	15	46	20	к-	@ cmm !			⊢κ
340315001	07-09	445	45	15	43	19	L –	o cammi 	• •	••••• •	<u>-</u> ι
340250005	07-09	458	47	15	45	19	м-	0 00000			⊢м
340230011	07-09	459	48	17	47	22	N -				- N
340273001	07-09	456	48	16	47	22	0-	0 @ 03333			⊢ο
090019003	07-09	457	47	16	44	21	Р-	co (camp	•	(momo (mom) o	- P
361030009	07-09	890	47	16	46	20	Q –	@ C			⊢ Q
340190001	07-09	455	50	16	48	21	R –	(DOD) CHIEF			⊢ R
360790005	07-09	459	44	14	42	19	s –	o amant	• -		– s
340210005	07-09	456	49	16	48	22	т-	(1111111)	- : •		⊢ ⊤
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361030004	07-09	453	48	14	46	18	Y-	Canan			- Y
090090027	07-09	456	41	14	40	17	Z –	· · · · · · · · · · · · · · · · · · ·	•:		– Z
360270007	07-09	456	43	14	41	17	AA -		•	(IIII III CO (III)	- AA
090093002	07-09	459	47	15	45	18	AB –	@ @@	•		- AB
090050005	07-09	446	46	15	43	19	AC –	cumu	•	• • • • • • • • • • • • • • • • • • •	1 /10
361111005	07-09	459	41	12	39	16	AD -	o	•		AD
	Key	25" mean	median	overall mèdián	75 th	2 th	0)	50	100	150
	+⊦ ⊦			0,2	<u> </u>	-1				$O_3(ppb)$	

Figure 3-107 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the New York CSA.

Philadelphia CSA

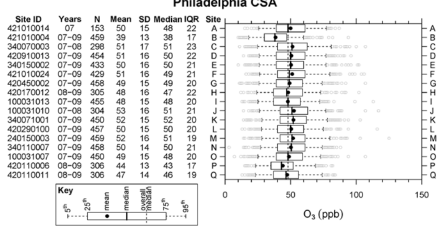


Figure 3-108 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Philadelphia CSA.

						Pho	ceni	k CE	BSA			
Site ID	Years	Ν	Mean	SD	Media	n IQR	Site				1	
040133002	07-09	455	53	9	53	11	A -		o anama [•			<u>-</u>
040133003	07-09	459	57	10	57	12	В –	0	o coo anna			- B
040139997	07-09	455	56	10	56	12	<u>c</u> –		0 00 0000			ΗŪ
040131004	07-09	459	58	10	58	13	Ď –		o opumni [•		μĎ
040134005	07-09	454	55	10	55	13	E –		cmpmmi	10000 00		⊢E
040134003	07-09	459	55	9	55	12	F		o commu s			⊢ F
040130019	07-09	459	55	10	55	12	G –		0 0 mmm) 🚺			G
040137020	07-09	459	56	9	56	12	н –		co mont			⊢н
040137024	07-09	459	56	9	57	12	1-		ampann 	1 00000		<u>⊢</u> ।
040137022	07-09	457	56	10	56	13	J –		aman an			⊢J
040132001	07-09	459	53	10	53	13	К-	0	ooaaa 🚺			⊢ĸ
040137021	07-09	457	59	9	59	12	L-		(1997)	• • • • IIIIDID O		⊢L
040134004	07-09	459	56	9	56	12	м –		• • • • • • • • • • • • • • • • • • •	• • • • • • • • •		- M
040131010	07-09	456	55	9	55	13	N -		accama al []			⊢ N
040137003	07-09	455	52	8	52	12	<u>o</u> –		an commit 🛉			- <u>o</u>
040132005	07-09	459	57	8	57	11	P -		acomi Li	-		- P
040139704	07-09	459	58	9	59	11	Q -		0 0 0 0 0 (• • • • • • • • • • •		⊢ Q
040135100	07-09	453	55	10	57	14	R-			- men o		⊢ R
040134010	07-09	457	48	9	48	13	<u>s</u> –	0		00 00		- <u>s</u>
040134008	07-09	459	58 53	9 9	57 53	14						
040139702 040139706	07-09 07-09	451 448	53 58	9 11	53 58	10 14	U – V –	0				Fv
040139708	07-09	448 459	58 59	9	58 59	14	w-		· · · · · · · · · · · · · ·			Εŵ
040213001	07-09	459	45	9	46	11	x ¬					Fx
040213009	07-09	459	48	9	40	12	Ŷ					ΕŶ
040213003	07-09	459	52	9	51	11	ż-			1		- ż
040139508	07-09	459	57	8	56	11	AĀ -		· · · · · ·			- ÃA
040134011	07-09	459	46	9	46	12	AB -					- AB
040213003	07-09	459	52	ğ	52	11	AC -					
040218001	07-09	459	59	9	59	12	AD -			• - · · · · · · · · · · · · · · · · · ·		- AD
040213007	07-09	459	50	8	50	10	AE -		amanni 🚺	- 1000		- AE
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	Key	5	lian	llan			C)	50		100	150
	±2+	- 25"		overall median	175 th	- 95 th				O ₃ (ppb)		

Figure 3-109 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Phoenix CBSA.

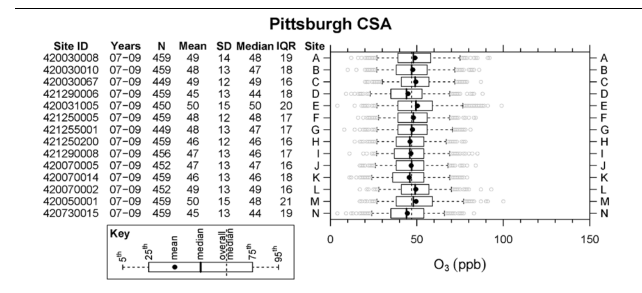


Figure 3-110 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Pittsburgh CSA.

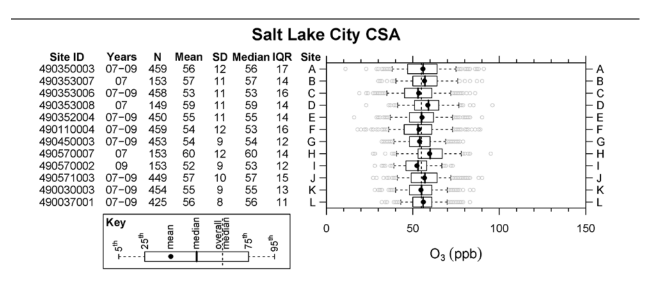


Figure 3-111 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Salt Lake City CSA.

San Antonio CBSA

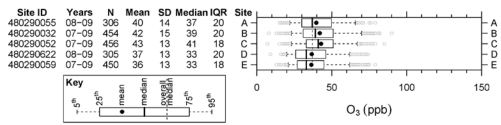


Figure 3-112 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the San Antonio CBSA.

					Sa	an F	rand	isco CSA
Site ID	Years	Ν	Mean	SD	Media	n IQR	Site	
060010009	08-09	306	29	9	28	12	A –	
060750005	07-09	458	28	8	27	10	В –	
060010006	07-08	303	31	9	30	12	ē-	
060012004	08-09	306	25	7	24	10	Ď-	
060012001	07-09	459	35	10	33	12	Ē-	
060811001	07-09	459	31	9	29	11	Ē-	
060131004	07-08	306	31	8	29	12	Ġ –	·
060011001	07-09	456	34	10	33	12	н –	
060130002	07-09	458	42	13	40	18		
060410001	07-09	458	29	8	28	10	j –	
060950006	07-08	306	40	11	39	13	к–	
060010007	07-09	459	43	14	41	18	ĩ –	
060852007	07-08	306	34	10	33	13	м-	
060950004	07-09	459	35	10	34	12	N -	mi
060133001	07-08	306	41	10	41	12	0 -	
060850005	07-09	456	36	10	35	13	Р –	• • • • • • • • • • • • • • • • • • •
060851001	07-09	459	39	12	37	16	Q –	cm+[]• tmmsss@c c o
060950005	07-09	459	39	11	37	12	R –	- R
060131002	07-09	459	47	12	45	15	s –	
060550003	07-09	459	37	9	35	10	Т –	ami[•]tamooacoo o
060870006	07-08	306	38	9	37	11	U –	····
060870003	07-09	456	31	8	30	10	V –	ocmat [●] +mmooo − V
060953003	07-09	455	44	13	43	17	w –	o @md ₽tmmoccom o −W
060870007	07-09	456	33	8	32	10	X –	oo œ≡ [€] tœ≡ œo o — X
060970003	07-09	459	31	8	31	10	Y –	amit
060852006	07-09	458	44	11	43	14	Z –	○ @ D
060870004	07-09	459	33	8	32	10	AA -	
060850002	07-09	459	44	11	42	15	AB –	
060971003	08-09	306	34	10	33	13	AC -	
060690002	07-09	456	42	10	40	12	AD -	
060690003	07-09	457	54	12	54	16	AE -	
	14			-			1	· · · · · · · · · · · · · · · · · · ·
	Key		median	overall mèdiăñ	75 th	£	C	50 100 150
	1.0°#		ĔĔ	3Ĕ	- 175	+ 95 th		O ₃ (ppb)
			-	- 1				

Figure 3-113 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the San Francisco CSA.

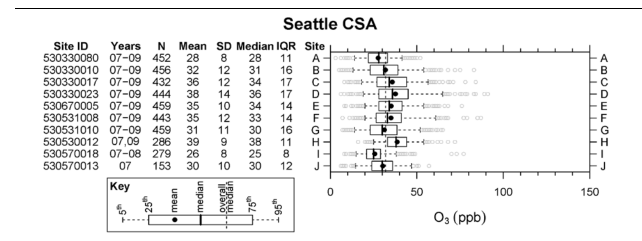


Figure 3-114 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the Seattle CSA.

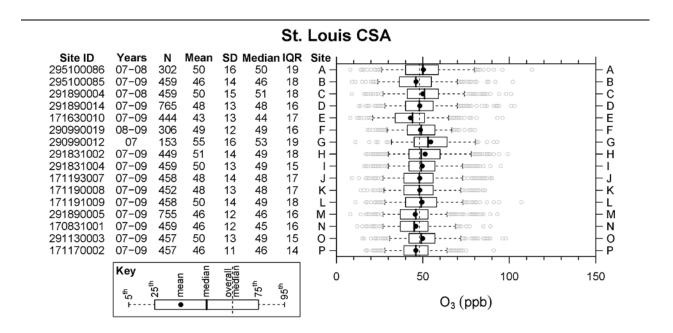


Figure 3-115 Site information, statistics and box plots for 8-h daily max ozone from AQS monitors meeting the warm-season data set inclusion criteria within the St. Louis CSA.

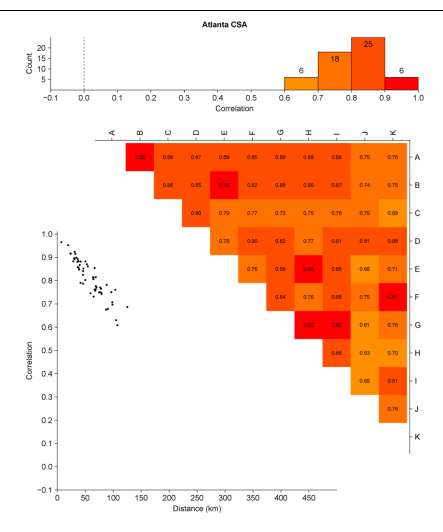
3.9.3 Ozone Concentration Relationships for the Urban Focus Cities

2 3 4

5

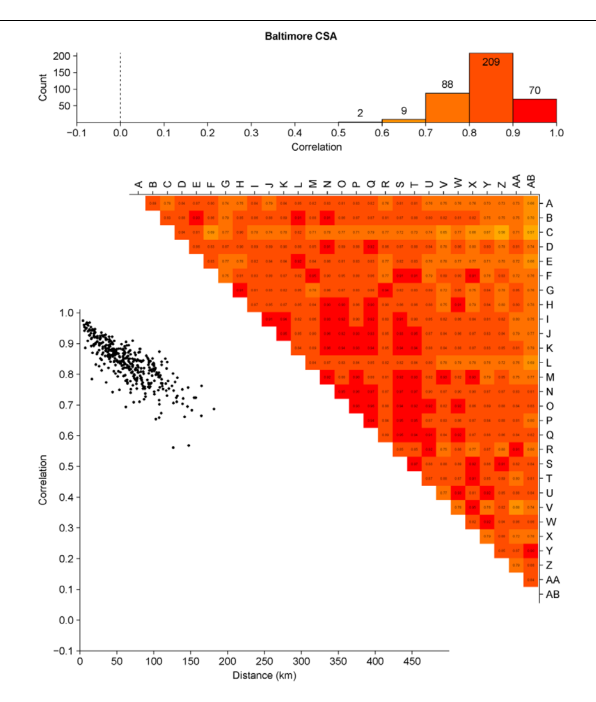
1

This section contains histograms and contour matrices of the Pearson correlation coefficient (R) and the coefficient of divergence (COD) between 8-h daily max O_3 concentrations from each monitor pair within the 20 urban focus cities discussed in Section <u>3.6.2.1</u>. These figures also contain scatter plots of R and COD as a function of straight-line distance between monitor pairs.



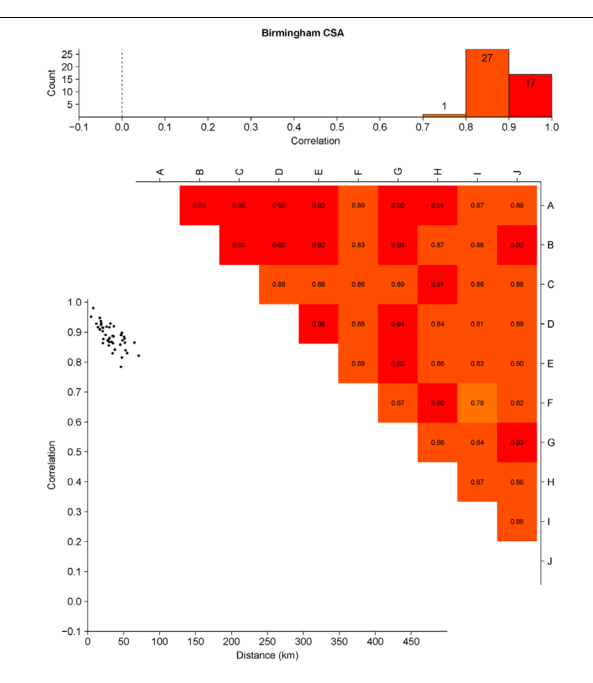
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-116 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.



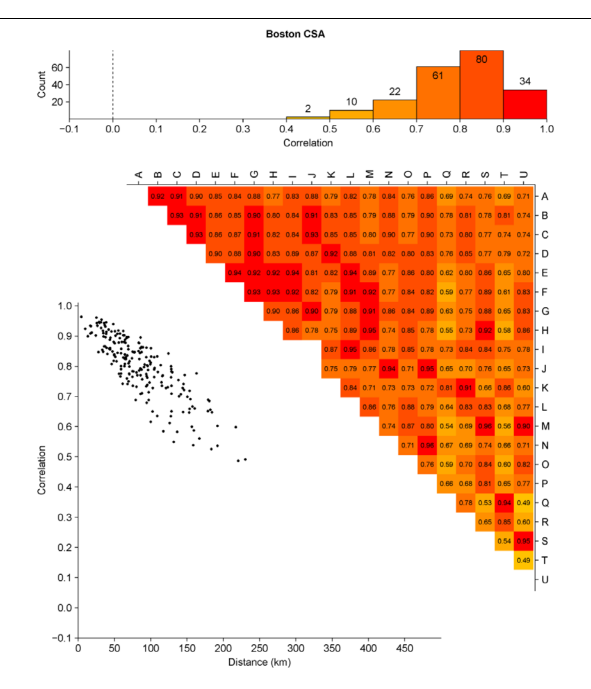
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-117 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Baltimore CSA.



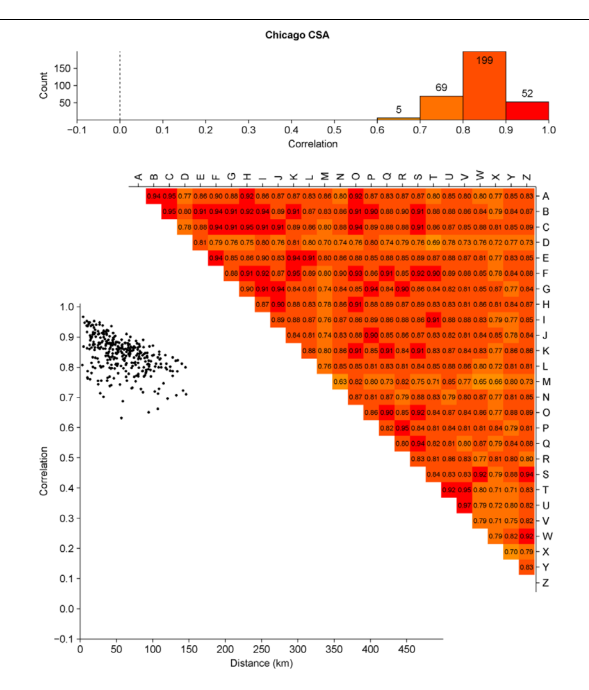
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-118 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Birmingham CSA.



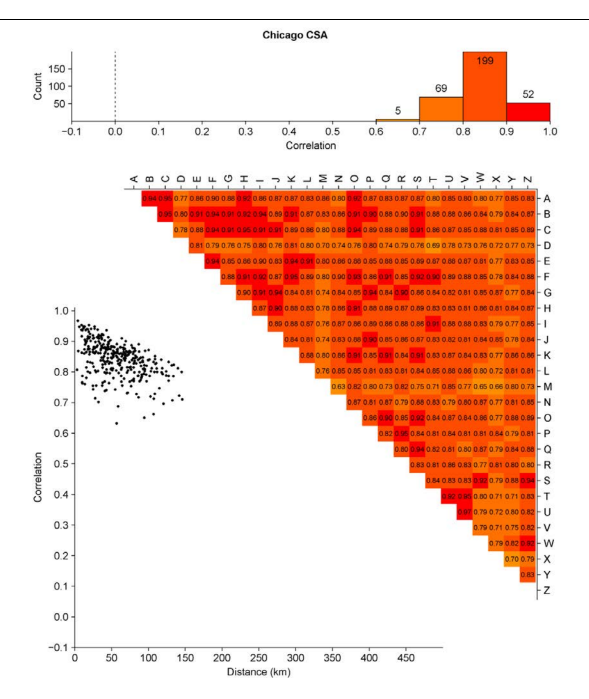
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-119 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.



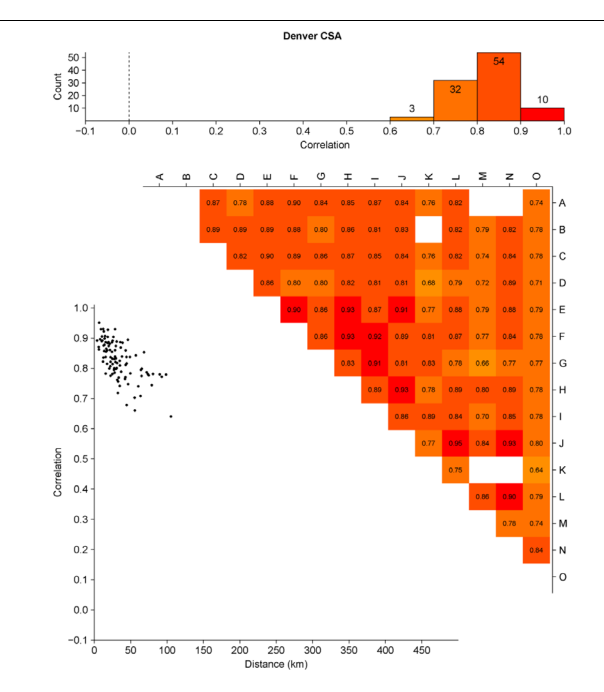
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-120 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Chicago CSA.



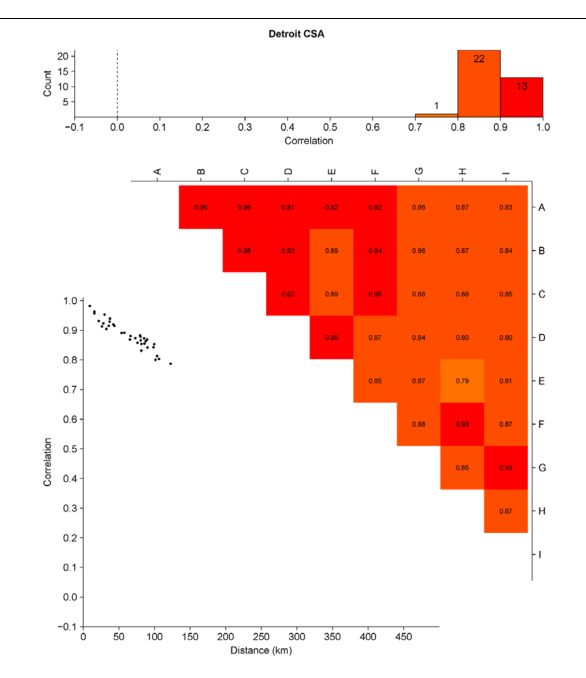
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-121 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Dallas CSA.



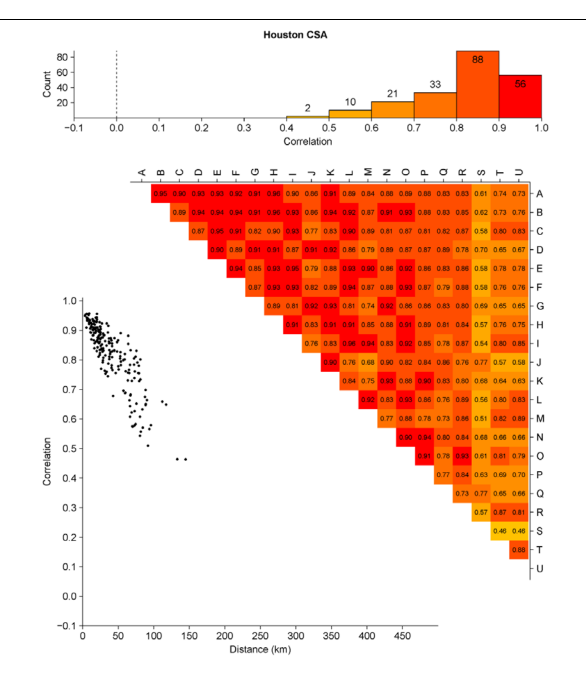
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-122 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Denver CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-123 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Detroit CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-124 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Houston CSA.

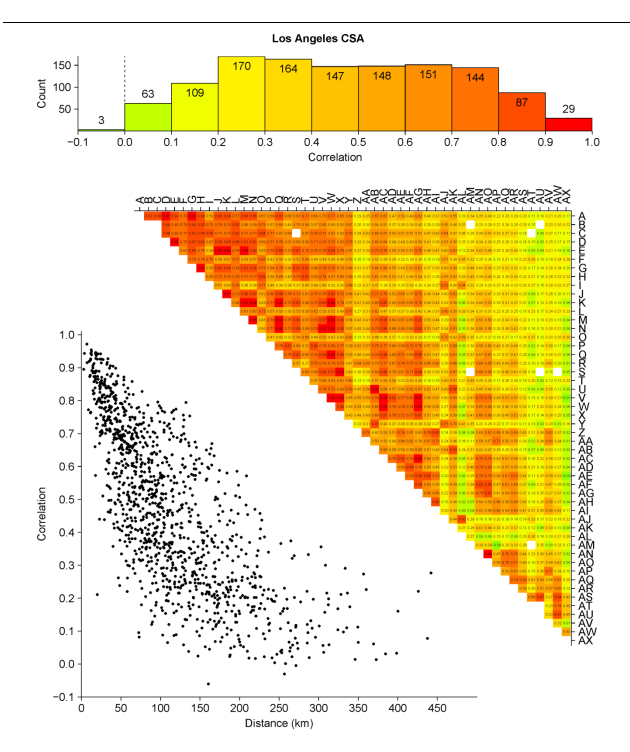


Figure 3-125 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.

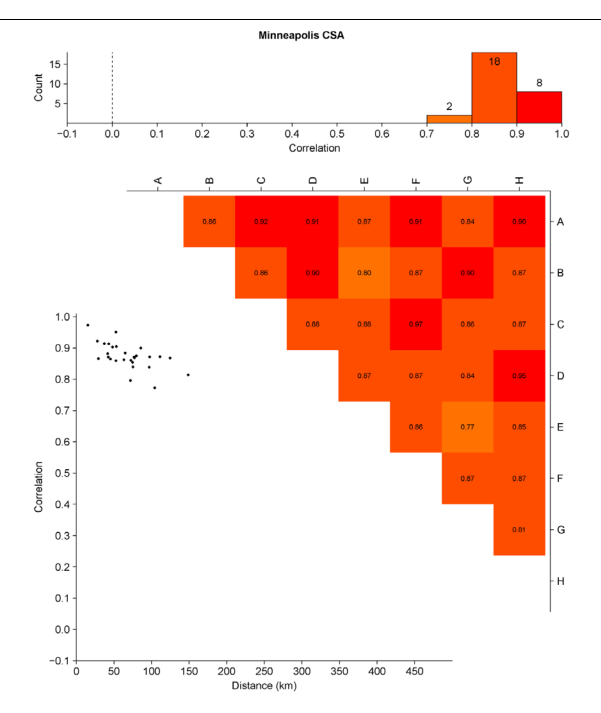
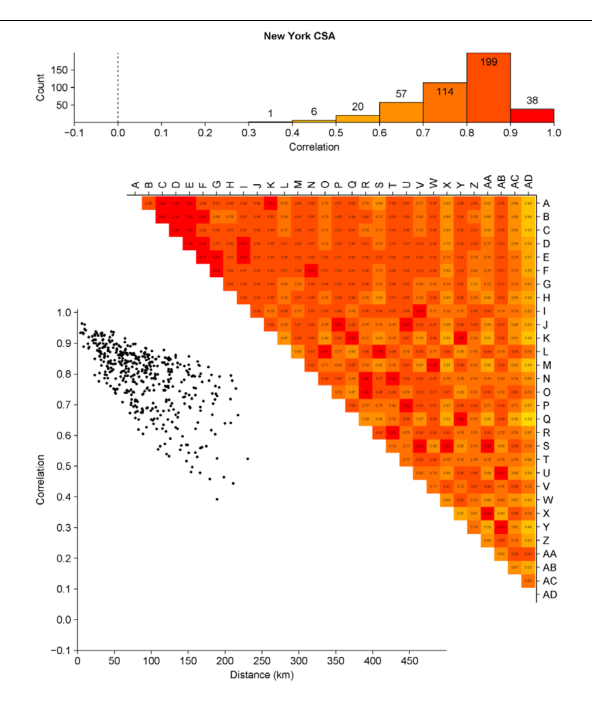
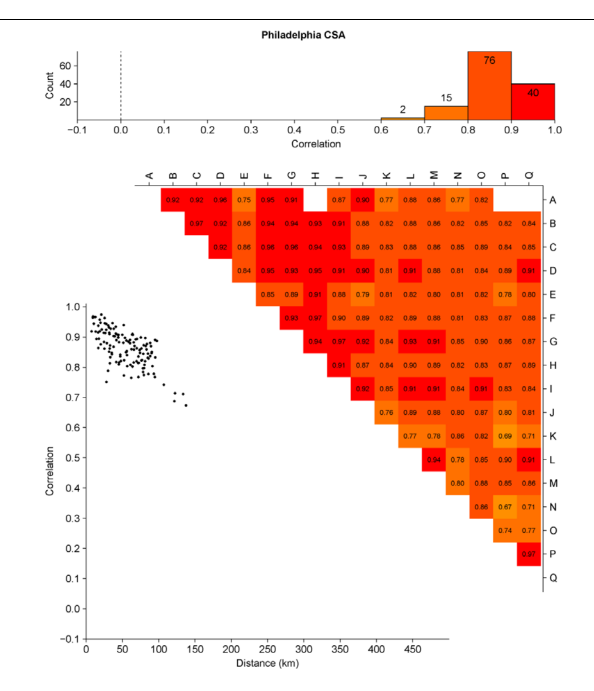


Figure 3-126 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Minneapolis CSA.



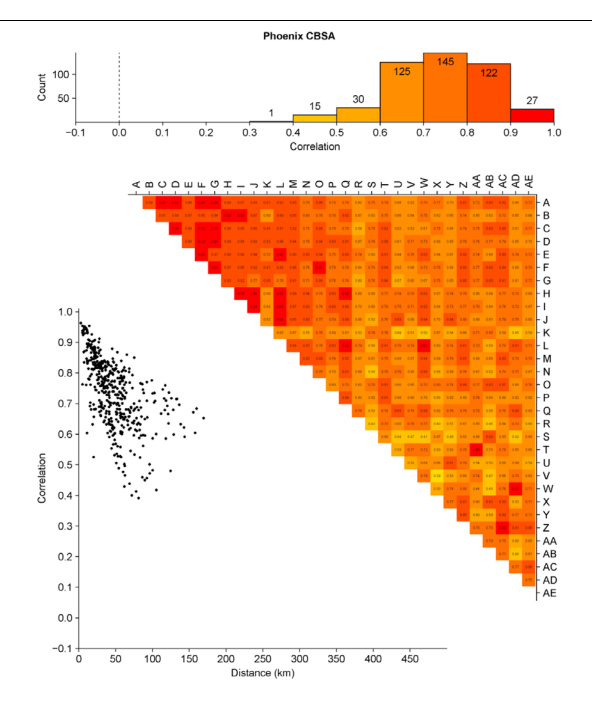
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-127 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the New York CSA.



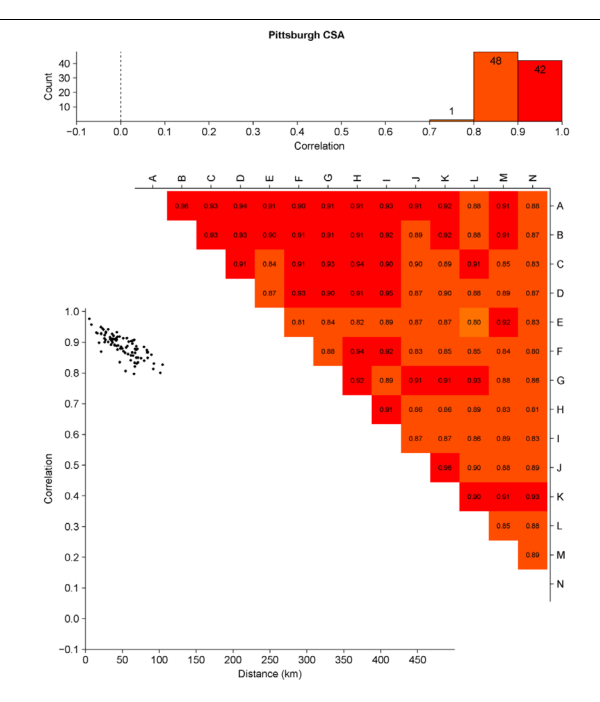
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-128 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Philadelphia CSA.



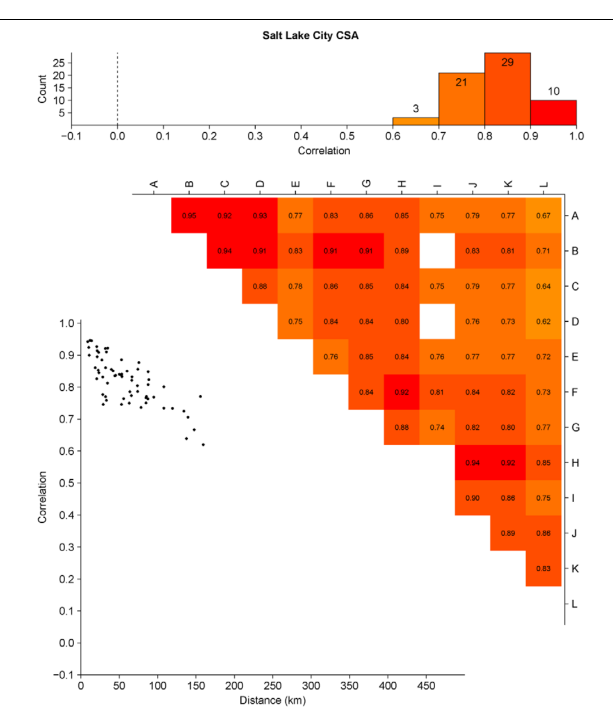
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-129 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Phoenix CBSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-130 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Pittsburgh CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-131 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Salt Lake City CSA.

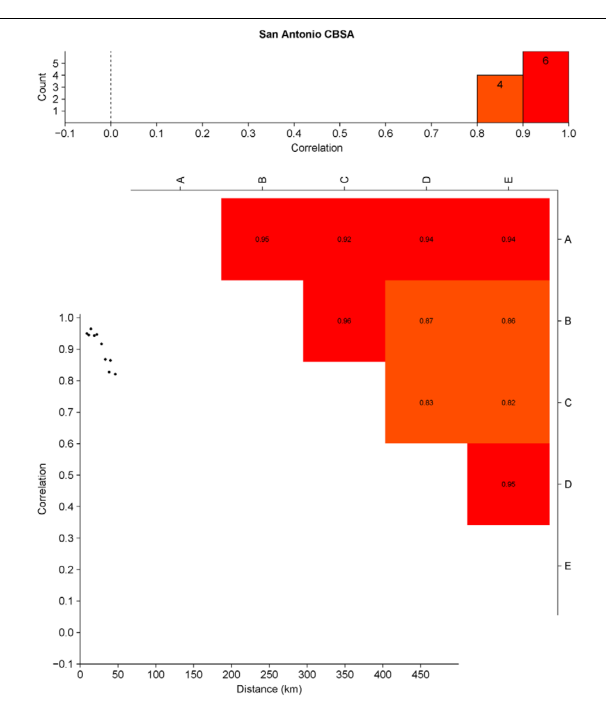
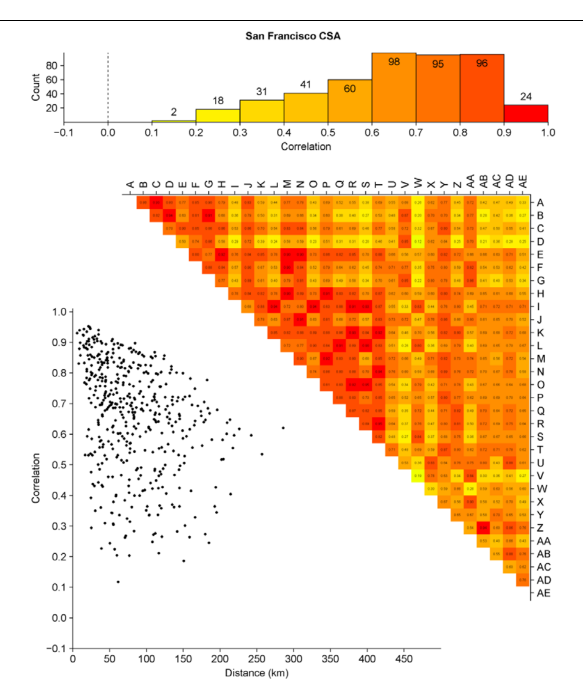


Figure 3-132 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Antonio CBSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-133 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Francisco CSA.

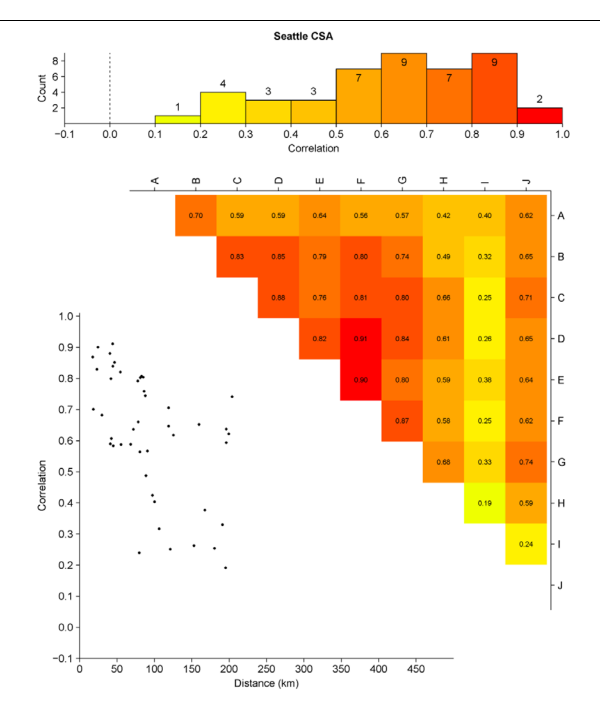


Figure 3-134 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Seattle CSA.

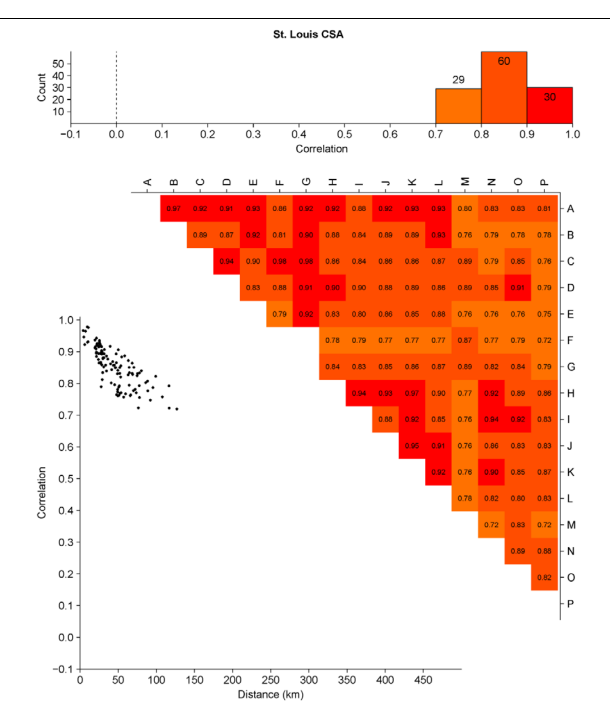


Figure 3-135 Pair-wise monitor correlation coefficients (R) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the St. Louis CSA.

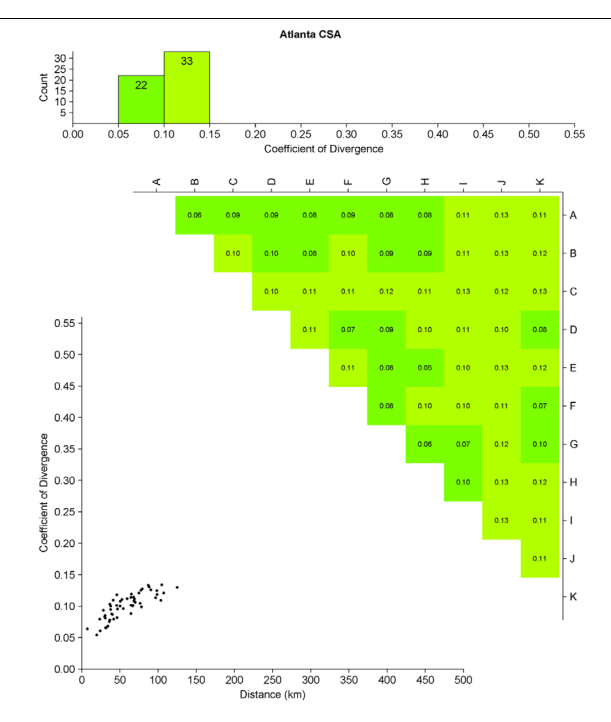
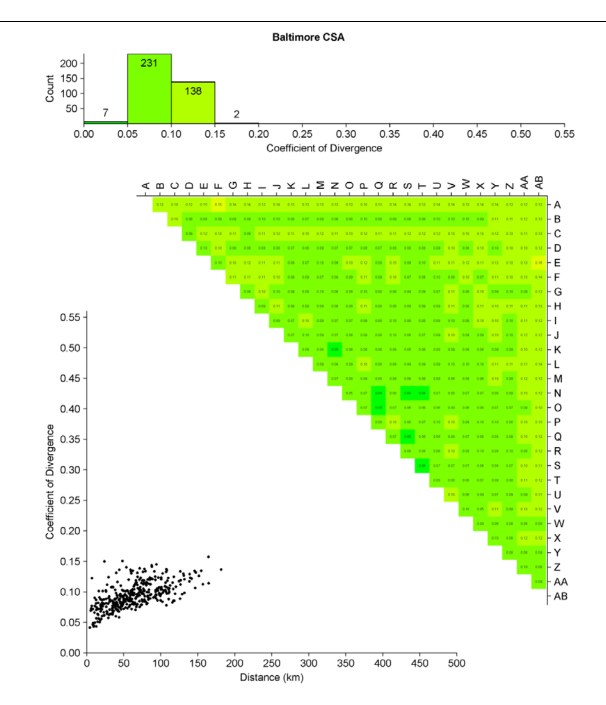
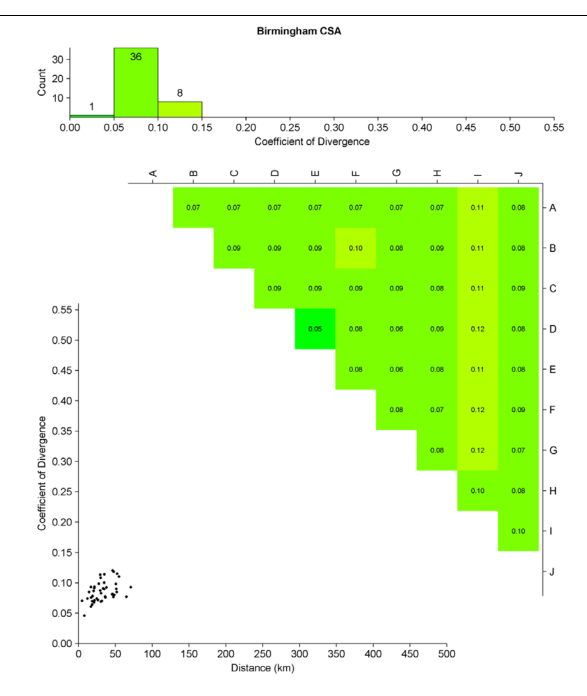


Figure 3-136 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Atlanta CSA.



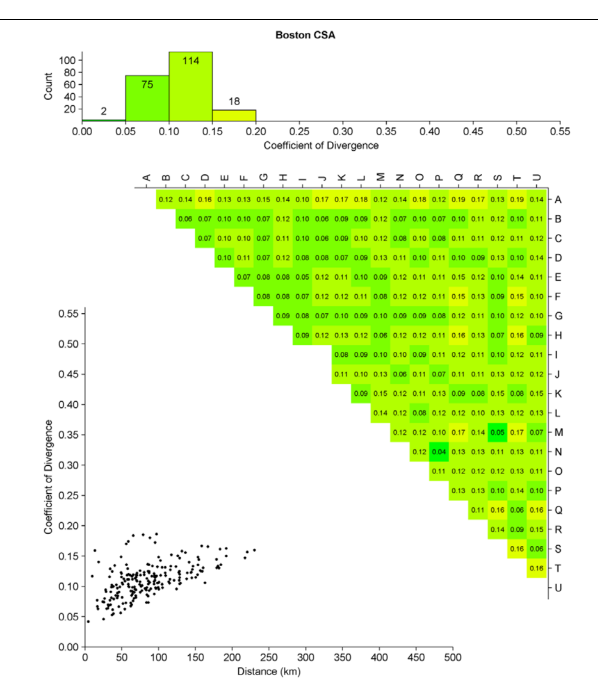
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-137 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Baltimore CSA.



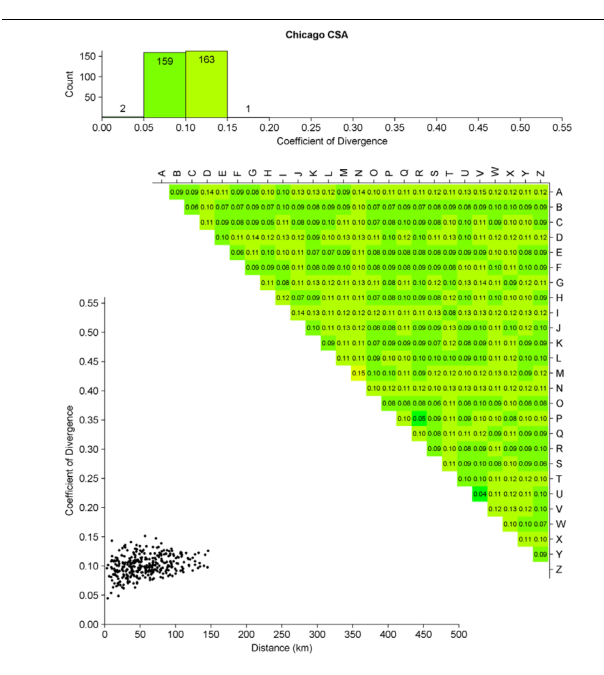
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-138 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Birmingham CSA.



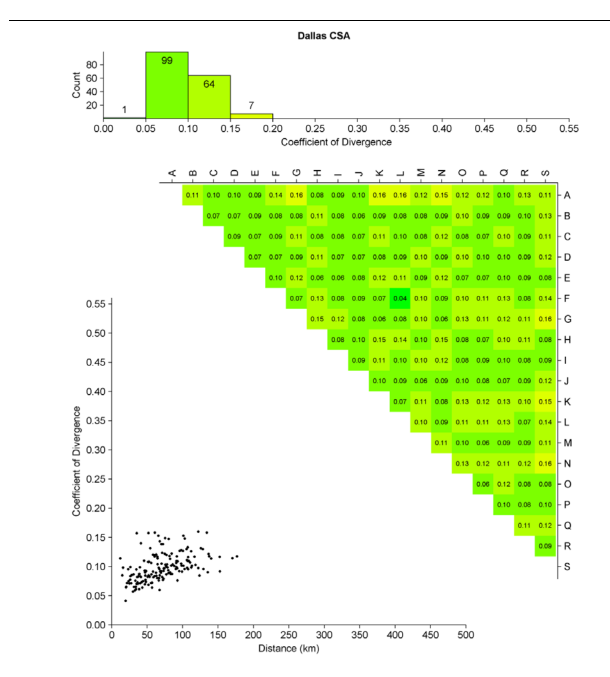
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-139 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Boston CSA.



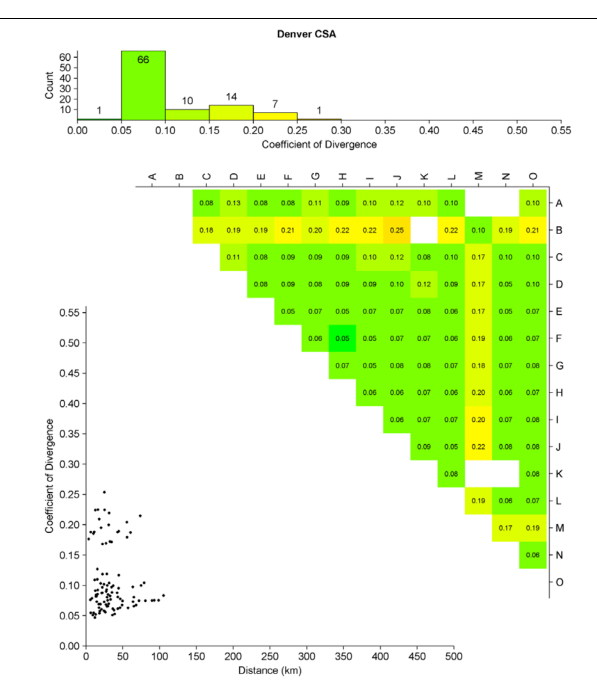
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-140 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Chicago CSA.



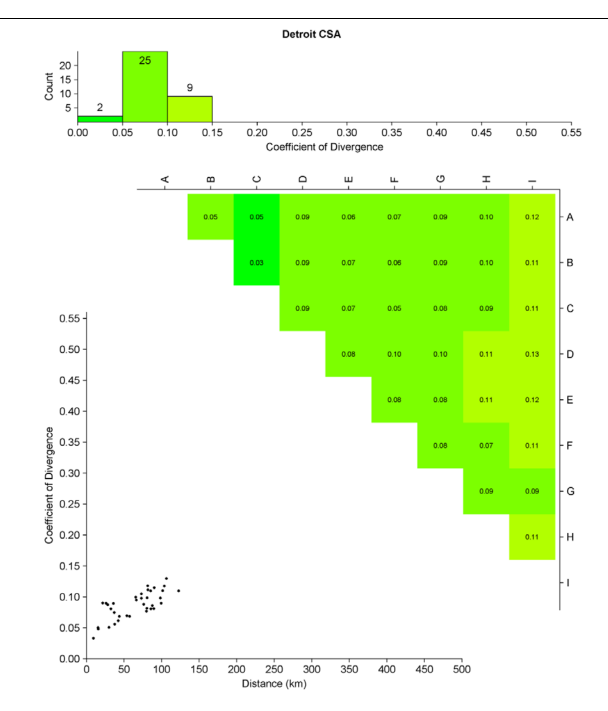
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-141 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Dallas CSA.



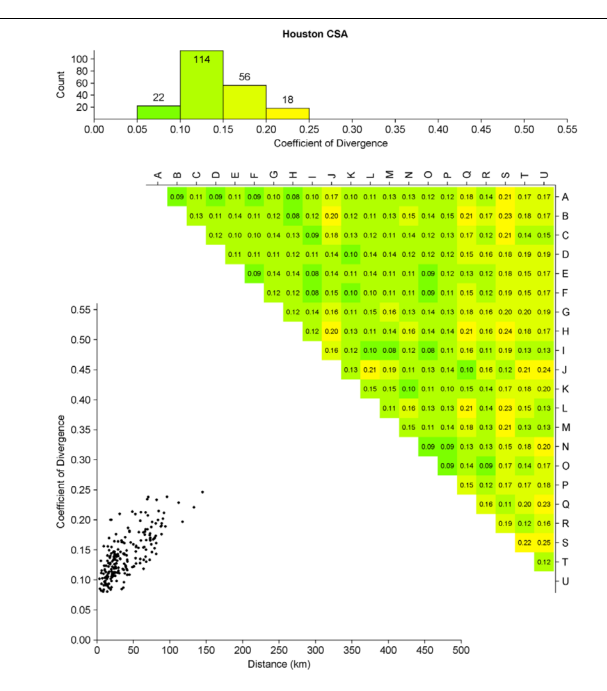
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-142 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Denver CSA.



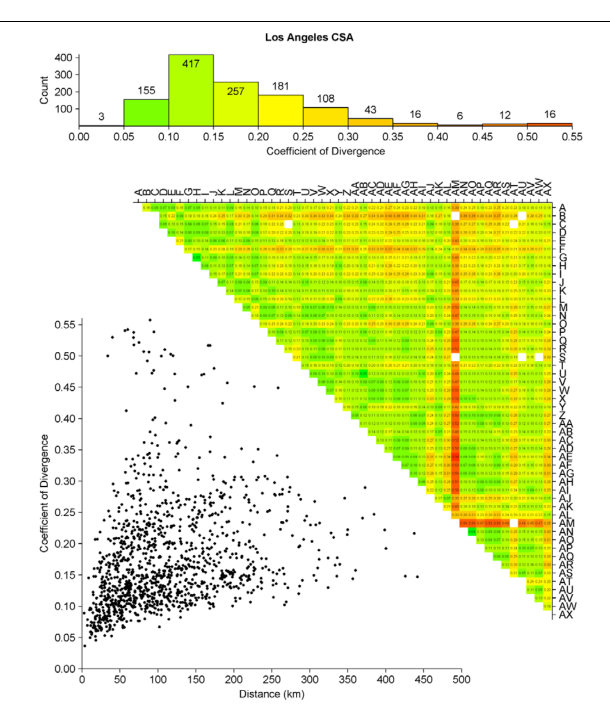
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-143 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Detroit CSA.



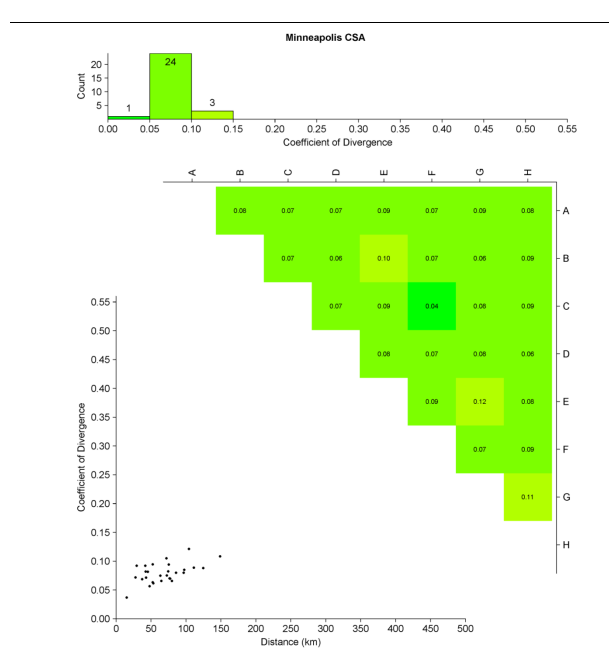
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-144 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Houston CSA.



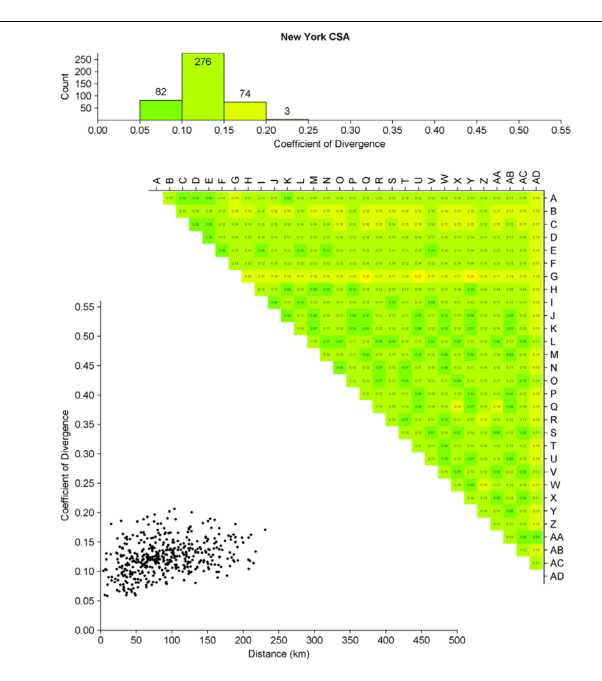
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-145 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Los Angeles CSA.



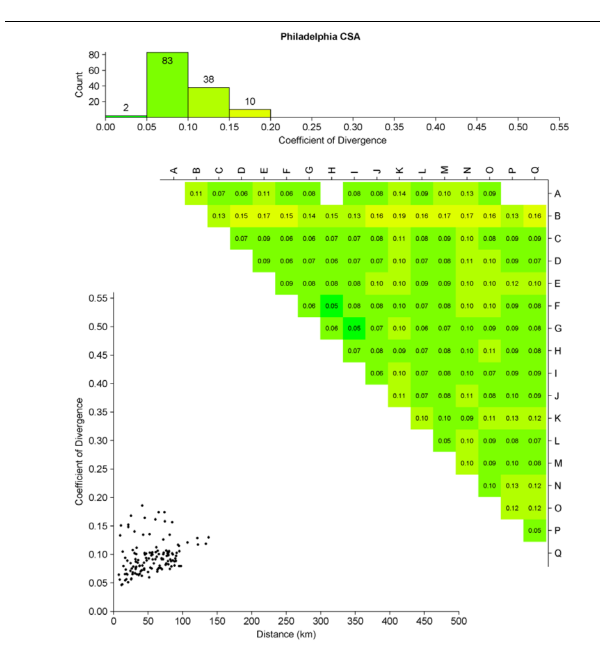
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-146 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Minneapolis CSA.



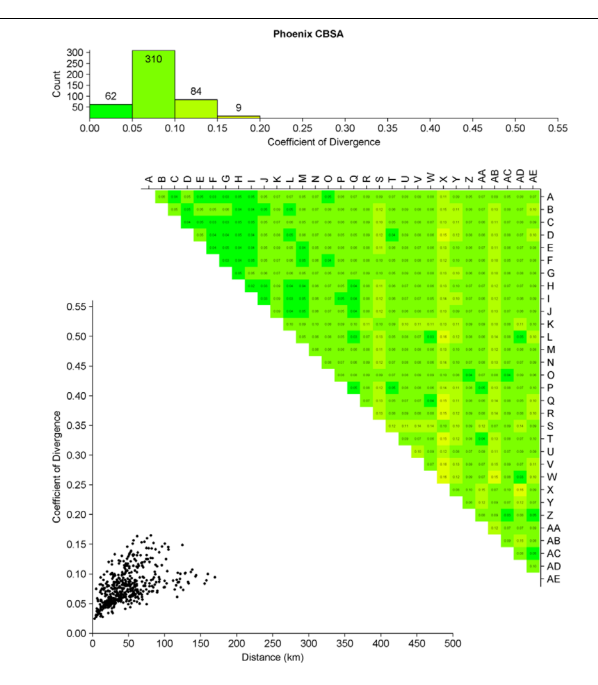
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-147 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the New York CSA.



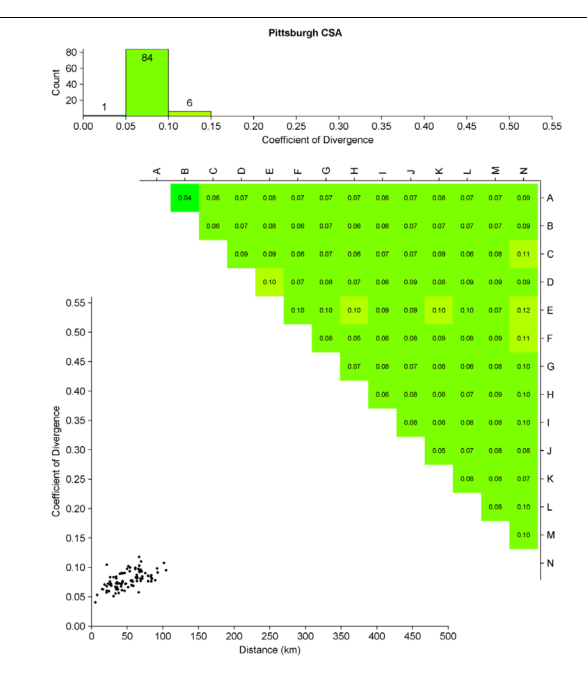
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-148 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Philadelphia CSA.



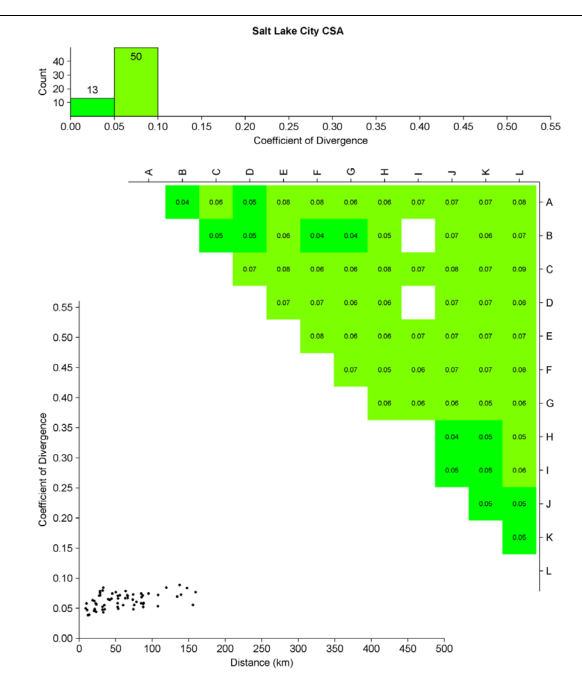
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-149 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Phoenix CBSA.



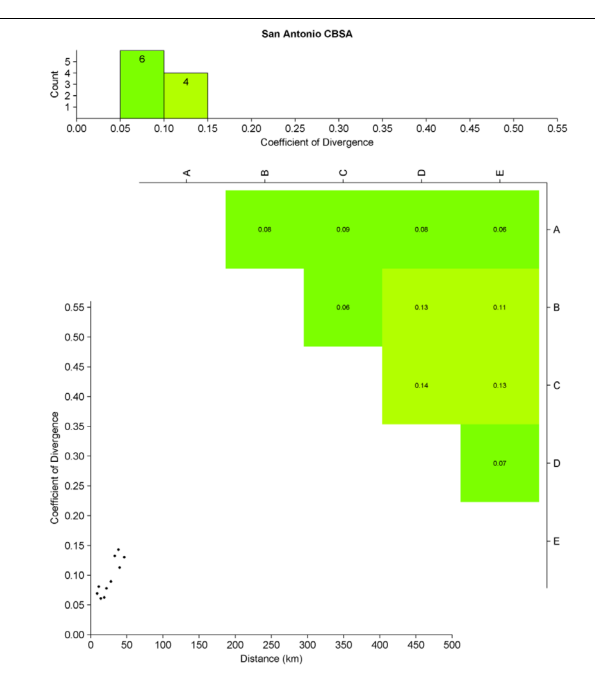
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-150 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Pittsburgh CSA.



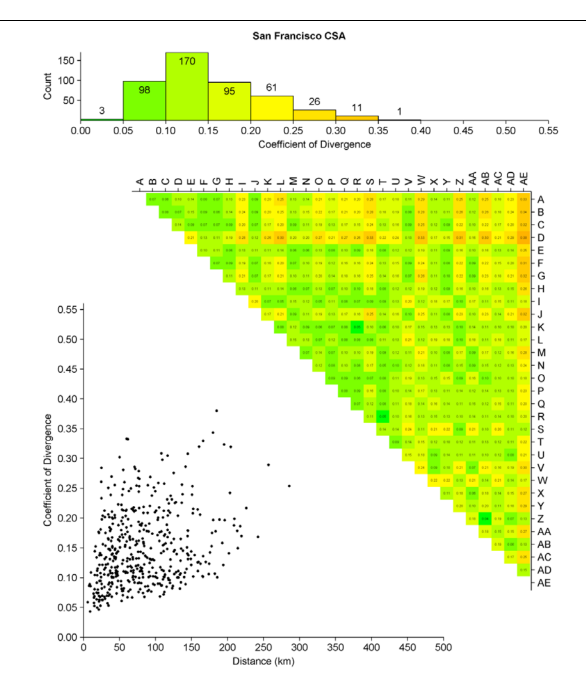
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-151 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Salt Lake City CSA.



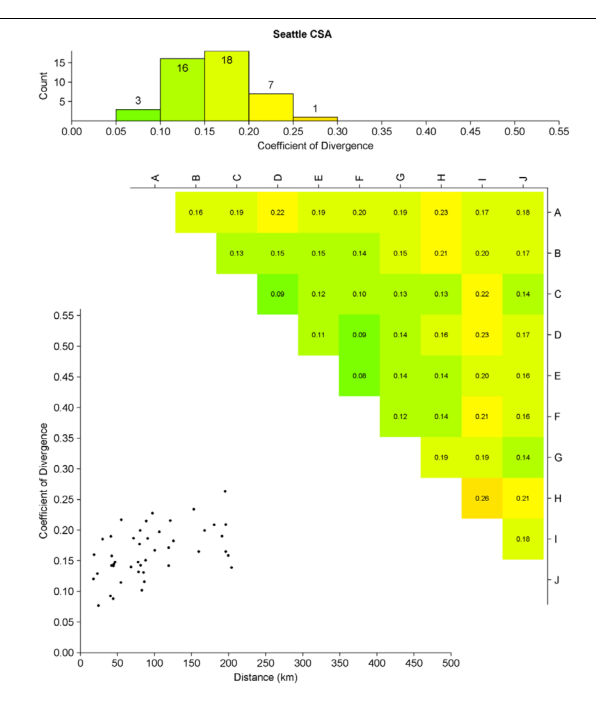
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-152 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Antonio CBSA.



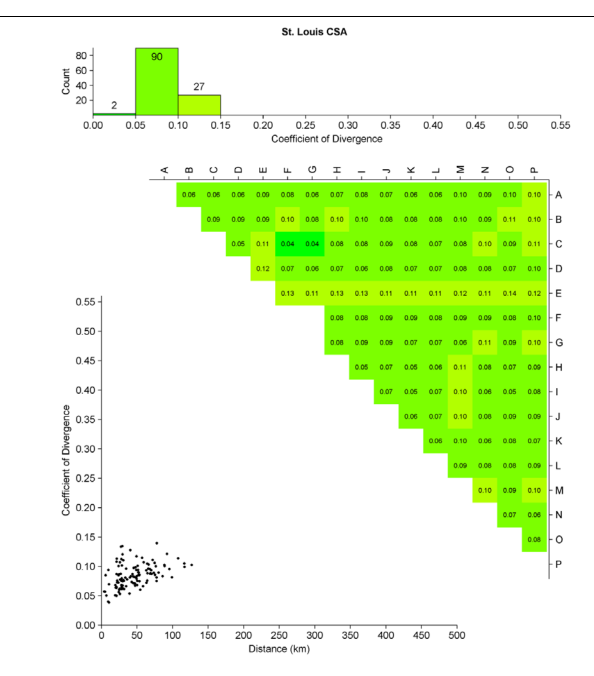
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-153 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the San Francisco CSA.



Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-154 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the Seattle CSA.



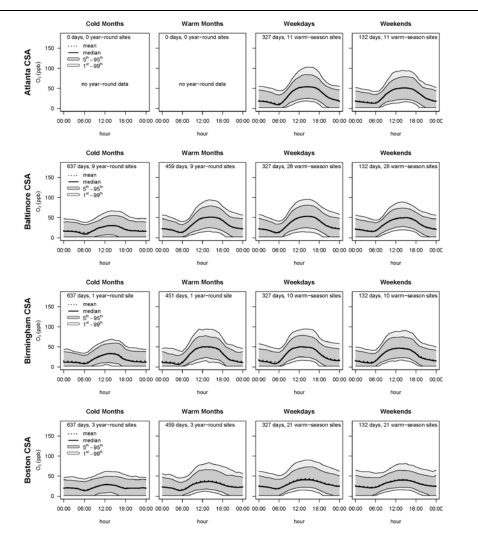
Note: The colors in the histogram bins correspond to the levels of the contour matrix. The histogram includes the number of monitor pairs per bin and the contour matrix includes the numeric values of R.

Figure 3-155 Pair-wise monitor coefficient of divergence (COD) expressed as a histogram (top), contour matrix (middle) and scatter plot versus distance between monitors (bottom) for the St. Louis CSA.

3.9.4 Hourly Variations in Ozone for the Urban Focus Cities

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- 4 5

This section contains diel plots of 1-h avg O_3 data to supplement the discussion on hourly variations in O_3 concentrations from Section <u>3.6.3.2</u> using data from the 20 urban focus cities first introduced in Section <u>3.6.2.1</u>. Comparisons are made between cold months (October-April) and warm months (May-September), using the year-round data set, and between weekdays (Mon-Fri) and weekends (Sat-Sun) using the warm-season data set.



Note: No year-round monitors were available for the cold month/warm month comparison in the Atlanta CSA.

Figure 3-156 Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).

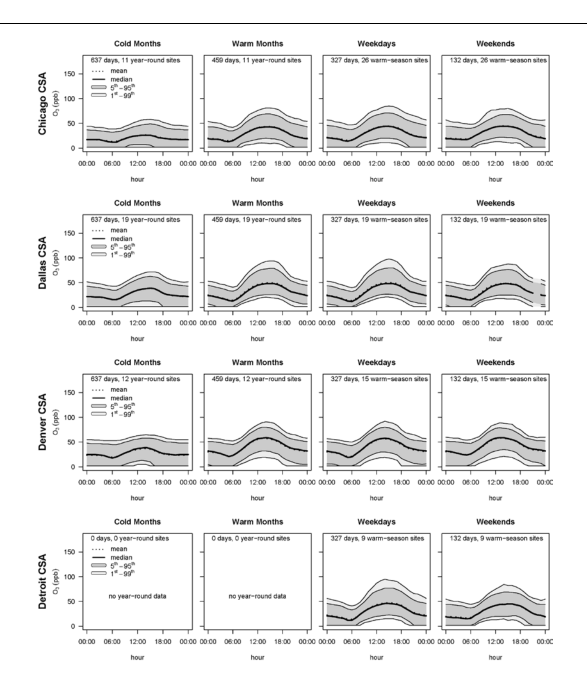
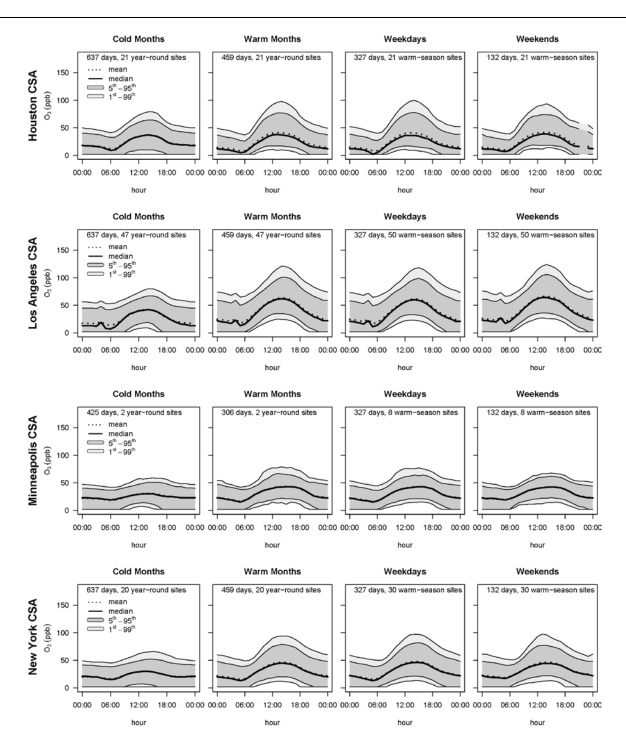


Figure 3-157 Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).



Note: No year-round monitors were available for the cold month/warm month comparison in the Detroit CSA.

Figure 3-158 Diel patterns in 1-h avg ozone for select CSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).

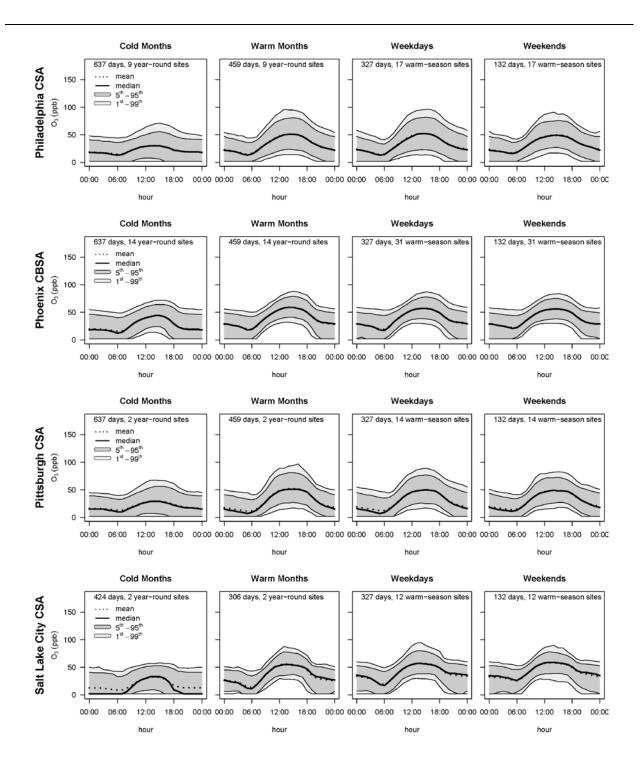


Figure 3-159 Diel patterns in 1-h avg ozone for select CSAs/CBSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).

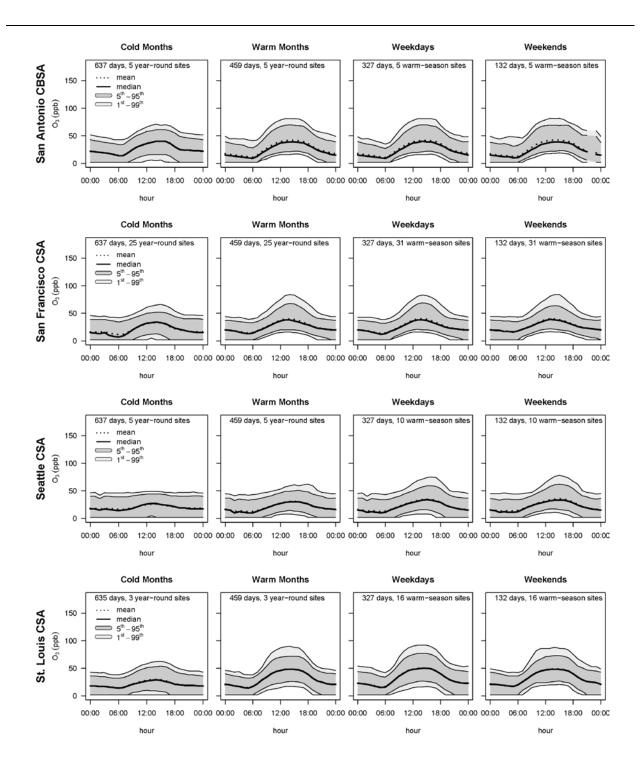


Figure 3-160 Diel patterns in 1-h avg ozone for select CSAs/CBSAs between 2007 and 2009 using the year-round data set for the cold month/warm month comparison (left half) and the warm-season data set for the weekday/weekend comparison (right half).

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4 EXPOSURE TO AMBIENT OZONE

4.1 Introduction

1	The 2006 O ₃ AQCD evaluated O ₃ concentrations and exposures in multiple
2	microenvironments, discussed methods for estimating personal and population exposure
3	via monitoring and modeling, analyzed relationships between personal exposure and
4	ambient concentrations, and discussed the implications of using ambient O ₃
5	concentrations as an estimate of exposure in epidemiologic studies. This chapter presents
6	new information regarding exposure to ambient O ₃ which builds upon the body of
7	evidence presented in the 2006 O_3 AQCD. A brief summary of findings from the 2006 O_3
8	AQCD is presented at the beginning of each section as appropriate.
9	Section 4.2 presents general exposure concepts describing the relationship between
10	ambient pollutant concentrations and personal exposure. Section 4.3 describes exposure
11	measurement techniques and studies that measured personal, ambient, indoor, and
12	outdoor concentrations of O_3 and related pollutants. Section <u>4.4</u> presents material on
13	parameters relevant to exposure estimation, including activity patterns, averting behavior,
14	and population proximity to ambient monitors. Section 4.5 describes techniques for
15	modeling local O3 concentrations, air exchange rates, microenvironmental concentrations,
16	and personal and population exposure. Section 4.6 discusses the implications of using
17	ambient O3 concentrations to estimate exposure in epidemiologic studies, including
18	several factors that contribute to exposure error.

4.2 General Exposure Concepts

19A theoretical model of personal exposure is presented to highlight measurable quantities20and the uncertainties that exist in this framework. An individual's time-integrated total21exposure to O3 can be described based on a compartmentalization of the person's22activities throughout a given time period:

$$E_T = \int C_j \, dt$$

Equation 4-1

23 where E_T = total exposure over a time-period of interest, C_j = airborne O₃ concentration at 24 microenvironment *j*, and *dt* = portion of the time-period spent in microenvironment *j*. 1Equation 4-1can be decomposed into a model that accounts for exposure to O_3 , of2ambient (E_a) and nonambient (E_{na}) origin of the form:

$$E_T = E_a + E_{na}$$

Equation 4-2

3Ambient O_3 is formed through photochemical reactions involving NO_X , VOCs, and other4compounds, as described in Chapter 3. Although nonambient sources of O_3 exist, such as5 O_3 generators and laser printers, these sources are specific to individuals and may not be6important sources of population exposure. Ozone concentrations generated by ambient7and nonambient sources are subject to spatial and temporal variability that can affect8estimates of exposure and influence epidemiologic effect estimates. Exposure parameters9affecting interpretation of epidemiologic studies are discussed in Section 4.5.

10This assessment focuses on the ambient component of exposure because this is more11relevant to the NAAQS review. Assuming steady-state outdoor conditions, E_a can be12expressed in terms of the fraction of time spent in various outdoor and indoor13microenvironments (Wallace et al., 2006; Wilson et al., 2000):

$$E_a = \sum f_o C_o + \sum f_i \mathcal{F}_{inf_i} C_{o,i}$$

Equation 4-3

14	where $f =$ fraction of the relevant time period (equivalent to dt in Equation 4-1), subscript
15	o = index of outdoor microenvironments, subscript $i =$ index of indoor
16	microenvironments, subscript $o, i =$ index of outdoor microenvironments adjacent to a
17	given indoor microenvironment <i>i</i> , and $F_{inf,i}$ = infiltration factor for indoor
18	microenvironment <i>i</i> . Equation 4-3 is subject to the constraint $\Sigma f_0 + \Sigma f_i = 1$ to reflect the
19	total exposure over a specified time period, and each term on the right hand side of the
20	equation has a summation because it reflects various microenvironmental exposures.
21	Here, "indoors" refers to being inside any aspect of the built environment, e.g., home,
22	office buildings, enclosed vehicles (automobiles, trains, buses), and/or recreational
23	facilities (movie theaters, restaurants, bars). "Outdoor" exposure can occur in parks or
24	yards, on sidewalks, and on bicycles or motorcycles. Assuming steady state ventilation
25	conditions, the infiltration factor is a function of the penetration (P) of O_3 into the
26	microenvironment, the air exchange rate (a) of the microenvironment, and the rate of O_3
27	loss (k) in the microenvironment; $F_{inf} = Pa/(a + k)$.

1In epidemiologic studies, the central-site ambient concentration, C_a , is often used in lieu2of outdoor microenvironmental data to represent these exposures based on the availability3of data. Thus it is often assumed that $C_o = C_a$ and that the fraction of time spent outdoors4can be expressed cumulatively as f_o ; the indoor terms still retain a summation because5infiltration differs among different microenvironments. If an epidemiologic study6employs only C_a , then the assumed model of an individual's exposure to ambient O_3 , first7given in Equation 4-3, is re-expressed solely as a function of C_a :

$$E_a = (f_o + \sum f_i \mathcal{F}_{inf_i}) C_a$$

Equation 4-4

8 The spatial variability of outdoor O₃ concentrations due to meteorology, topography, 9 varying precursor emissions and O₃ formation rates; the design of the epidemiologic 10 study; and other factors determine whether or not Equation 4-4 is a reasonable 11 approximation for Equation 4-3. These equations also assume steady-state 12 microenvironmental concentrations. Errors and uncertainties inherent in use of Equation 4-4 in lieu of Equation 4-3 are described in Section 4.6 with respect to implications for 13 14 interpreting epidemiologic studies. Epidemiologic studies often use concentration 15 measured at a central site monitor to represent ambient concentration; thus α , the ratio 16 between personal exposure to ambient O₃ and the ambient concentration of O₃, is defined 17 as:

$$\alpha = \frac{E_a}{C_a}$$

Equation 4-5

$$\alpha = f_o + \sum f_i \mathcal{F}_{inf_i}$$

Equation 4-6

19	where α varies between 0 and 1. If a person's exposure occurs in a single
20	microenvironment, the ambient component of a microenvironmental O ₃ concentration
21	can be represented as the product of the ambient concentration and F_{inf} . Wallace et al.
22	(2006) note that time-activity data and corresponding estimates of F_{inf} for each
23	microenvironmental exposure are needed to compute an individual's α with accuracy. In

1epidemiologic studies, α is assumed to be constant in lieu of time-activity data and2estimates of F_{inf} , which can vary with building and meteorology-related air exchange3characteristics. If important local outdoor sources and sinks exist that are not captured by4central site monitors, then the ambient component of the local outdoor concentration may5be estimated using dispersion models, land use regression models, receptor models, fine6scale CTMs or some combination of these techniques. These techniques are described in7Section $\underline{4.5}$.

4.3 Exposure Measurement

8	This section describes techniques that have been used to measure microenvironmental
9	concentrations of O3 and personal O3 exposures as well as results of studies using those
10	techniques. Previous studies from the 2006 O3 AQCD are described along with newer
11	studies that evaluate indoor-outdoor concentration relationships, associations between
12	personal exposure and ambient monitor concentration, and multipollutant exposure to
13	other pollutants in conjunction with O ₃ . Tables are provided to summarize important
14	study results.

4.3.1 Personal Monitoring Techniques

15	As described in the 2006 O ₃ AQCD, passive samplers have been developed and deployed
16	to measure personal exposure to O ₃ (Grosjean and Hisham, 1992; Kanno and
17	Yanagisawa, 1992). Widely used versions of these samplers utilize a filter coated with
18	nitrite, which is converted to nitrate by O_3 and then quantified by a technique such as ion
19	chromatography (Koutrakis et al., 1993). This method has been licensed and marketed by
20	Ogawa, Inc., Japan (Ogawa & Co, 2007). The cumulative sampling and the detection
21	limit of the passive badges makes them mainly suitable for monitoring periods of 24
22	hours or greater, which limits their ability to measure short-term daily fluctuations in
23	personal O ₃ exposure. Longer sampling periods give lower detection limits; use of the
24	badges for a 6-day sampling period yields a detection limit of 1 ppb, while a 24-hour
25	sampling period gives a detection limit of approximately 5-10 ppb. This can result in a
26	substantial fraction of daily samples being below the detection limit (Sarnat et al., 2006a;
27	Sarnat et al., 2005), which is a limitation of past and current exposure studies.
28	Development of improved passive samplers capable of shorter-duration monitoring with
29	lower detection limits would enable more precise characterization of personal exposure in
30	multiple microenvironments with relatively low participant burden.

1	The nitrite-nitrate conversion reaction has also been used as the basis for an active
2	sampler consisting of a nitrite-coated glass tube through which air is drawn by a pump
3	operating at 65 mL/min (Geyh et al., 1999; Geyh et al., 1997). The reported detection
4	limit is 10 ppb-h, enabling the quantification of O ₃ concentrations measured over a few
5	hours rather than a full day (Geyh et al., 1999).
6	A portable active O ₃ monitor based on the UV photometric technique used for stationary
7	monitors (Chapter $\underline{3}$) has recently been approved as a FEM (75 FR 22126). This monitor
8	includes a Nafion tube in the inlet line to equilibrate humidity, reducing the effect of
9	humidity changes in different microenvironments (Wilson and Birks, 2006). Its size and
10	weight (approximately $10 \times 20 \times 30$ cm; 2 kg) make it suitable for use in a backpack
11	configuration. The monitors are currently used by the U.S. National Park service as
12	stationary monitors to measure O_3 in several national parks (Chapter <u>3</u>). Future
13	improvements and continued miniaturization of real-time O3 monitors can yield highly
14	time-resolved personal measurements to further evaluate O3 exposures in specific
15	situations, such as near roadways or while in transit.

4.3.2 Indoor-Outdoor Concentration Relationships

- 16 Several studies summarized in the 2006 O₃ AQCD, along with some newer studies, have 17 evaluated the relationship between indoor O₃ concentration and the O₃ concentration 18 immediately outside the indoor microenvironment. These studies show that the indoor 19 concentration is often substantially lower than the outdoor concentration unless indoor 20 sources are present. Low indoor O_3 concentrations can be explained by reactions of O_3 21 with surfaces and airborne constituents. Studies have shown that O₃ is deposited onto 22 indoor surfaces where reactions produce secondary pollutants such as formaldehyde 23 (Reiss et al., 1995b; Reiss et al., 1995a). However, the indoor-outdoor relationship is 24 greatly affected by the air exchange rate; under conditions of high air exchange rate, such 25 as open windows, the indoor O_3 concentration may approach the outdoor concentration. 26 Thus, in rooms with open windows, the indoor-outdoor (I/O) ratio may approach 1.0. 27 Table 4-1 summarizes I/O ratios and correlations reported by older and more recent 28 studies, with discussion of individual studies in the subsequent text. In general, I/O ratios 29 range from about 0.1 to 0.4, with some evidence for higher ratios during the O_3 season 30 when concentrations are higher. 31 O₃ concentrations near and below the monitor detection limit cause uncertainty in I/O
- 32ratios, because small changes in low concentration values cause substantial variation in33resulting ratios. This problem is particularly acute in the non-ozone season when ambient34O3 concentrations are low. Further improvements in characterization of

1	microenvironmental O ₃ concentrations and I/O ratios will rely on improved monitoring.
2	Until new monitoring techniques are available and can be used in the field, past studies
3	summarized in the 2006 O3 AQCD remain relevant to consider along with more recent
4	studies in evaluating the relationship between indoor and outdoor O_3 concentrations.

Study	Location	Years/Season	Population	Sample duration	Ratio ^a	Correlation	Micro-environment	Comment
<u>Geyh</u> <u>et al.</u> (2000)	Upland, Southern California	June - September 1995 and May 1996	Children	6 days	0.24	NR Ho	Home	Air- conditioned Ratio: Indoor mean/outdoor
		October 1995- April 1996	_		0.15			mean
	Mountain Communities, Southern California	June - September 1995 and May 1996	_		0.36	-		Opening windows Ratio: indoor mean/outdoor
		October 1995- April 1996	_		0.08	-		mean
<u>Avol et</u> <u>al.</u> (1998a)	Southern California	February- December, 1994	NR	24 h	0.37 SD: 0.25	0.58	Home	Ratio: each pair of measurements
		Summer	_		0.43 SD: 0.29	NR		
		Non-summer	_		0.32 SD: 0.21	NR		
<u>Romieu</u> <u>et al.</u> (1998a)	Mexico City, Mexico	September 1993 - July 1994	Children	7 or 14 days	0.20 SD: 0.18 0.15 ^b Range: 0.01- 1.00	NR	Home	Ratio: each pair of measurements
<u>Lee et</u> <u>al.</u> (2004a)	Nashville, TN	Summer 1994	Children	1 week	0.1 SD: 0.18	NR	Home	Ratio: Indoor mean/outdoor mean

Study	Location	Years/Season	Population	Sample duration	Ratio ^a	Correlation	Micro-environment	Comment
<u>Héroux</u> <u>et al.</u> (2010)	Regina, Saskatchewa, Canada	Summer 2007	All age groups	5 days	0.13	NR	Home	Ratio: Indoor mean/outdoor mean
<u>Liu et al.</u> (1995)	Toronto, Canada	Winter, 1992	All age groups	1 week	0.07 SD: 0.10	NR	Home	Ratio: each pair of measurements
		Summer, 1992			0.40 SD: 0.29	-		
		Summer, 1992		12 h	0.30 SD: 0.32	-		Daytime Ratio: each pair of measurements
		Summer, 1992			0.43 SD: 0.54			Nighttime Ratio: each pair of measurements
<u>Romieu</u> <u>et al.</u> (1998a)	Mexico City, Mexico		Children	24 h/day, 14 days	0.15	NR	School	Ratio: each pair of measurements
			Children (during school hours)	5 h/day, 5 days, 10 days	0.30- 0.40	-		Immediately outside the schools
<u>Blondeau</u> <u>et al.</u> (2005)	La Rochelle, France	Spring, 2000	Children	2 weeks	Range: 0.00- 0.45	NR	School	No air conditioning Ratio: Indoor mean/outdoor mean
<u>López-</u> Aparicio	Prague, Czech Republic		0.10	NR	Historic	No heating or air		
<u>et al.</u> (2011)		Dec 2009	. <u>9</u> .00p0		0.30	- Library	conditioning Ratio: Indoor mean/outdoor mean	
<u>Riediker</u> <u>et al.</u> (2003)	North Carolina	August - October 2001	Adults	9 h	0.51 p- value: 0.000	NR	Vehicle	Ratio: Indoor mean/outdoor mean

^aMean value unless otherwise indicated

⁵Median

NR = not reported

SD = standard deviation.

1Geyh et al. (2000) measured 6-day indoor and outdoor concentrations at 116 homes in2southern California, approximately equally divided between the community of Upland3and several mountain communities. The extended sampling period resulted in a relatively4low detection limit (1 ppb) for the passive samplers used. The Upland homes were nearly5all air-conditioned, while the mountain community homes were ventilated by opening6windows. During the O3 season, the indoor O3 concentration averaged over all homes was

1	approximately 24% of the overall mean outdoor concentration in Upland (11.8 versus
2	48.2 ppb), while in the mountain communities, the indoor concentration was 36% of the
3	outdoor concentration (21.4 versus 60.1 ppb). This is consistent with the increased air
4	exchange rate expected in homes using window ventilation. In the non-ozone season,
5	when homes are likely to be more tightly closed to conserve heat, the ratios of indoor to
6	outdoor concentration were 0.15 (3.2 versus 21.1 ppb) and 0.08 (2.8 versus 35.7 ppb) in
7	Upland and the mountain communities, respectively. Avol et al. (1998a) observed a mean
8	I/O ratio of 0.37 for 239 matched 24-h samples collected between February and
9	December at homes in the Los Angeles area. The I/O ratio during summer was somewhat
10	higher than the non-summer I/O ratio (0.43 versus 0.32). The authors also reported a
11	correlation of 0.58 between the 24-h avg indoor concentration and the outdoor
12	concentration, which was only slightly higher than the correlation between the indoor
13	concentration and the concentration at the neighborhood fixed-site monitor (0.49).
14	Substantially higher summer I/O ratios were reported in a study in Toronto (Liu et al.,
15	1995), which found summer I/O ratios of 0.30-0.43, in comparison with a winter I/O ratio
16	of 0.07. Romieu et al. (1998a) reported a mean I/O ratio of 0.20 in 145 homes in
17	Mexico City for 7- or 14-day cumulative samples, with the highest ratios observed in
18	homes where windows were usually open during the day and where there was no
19	carpeting or air filters. Studies conducted in Nashville, TN and Regina, Saskatchewan
20	reported mean residential I/O ratios of approximately 0.1 (Héroux et al., 2010; Lee et al.,
21	<u>2004a</u>).

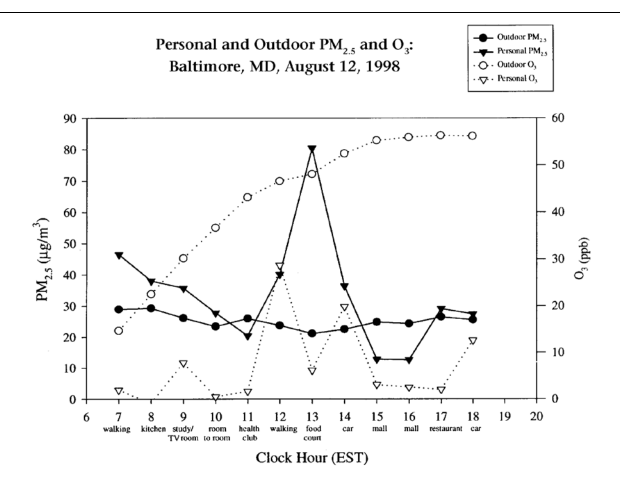
22 Investigators have also measured I/O ratios for non-residential microenvironments, 23 including schools and vehicles. Romieu et al. (1998a) reported that O₃ concentrations 24 measured during school hours (10-day cumulative sample, 5 h/day) were 30-40% of 25 concentrations immediately outside the schools, while overall I/O ratios (14-day 26 cumulative sample, 24 h/day) were approximately 15%. The authors attribute this 27 discrepancy to increased air exchange during the school day due to opening doors and 28 windows. Air exchange was also identified as an important factor in the I/O ratios 29 measured at eight French schools (Blondeau et al., 2005). In this study, the I/O ratios 30 based on simultaneous continuous measurements ranged from 0-0.45, increasing with 31 decreasing building tightness. A historical library building in Prague, Czech Republic 32 with no heating or air conditioning (i.e., natural ventilation) was observed to have ratios 33 of one-month indoor and outdoor concentrations ranging from 0.10-0.30 during a nine-34 month sampling campaign, with the highest ratios reported in Nov-Dec 2009 and the 35 lowest ratios during Jul-Aug 2009 (López-Aparicio et al., 2011). Indoor concentrations were relatively constant (approximately 3-7 μ g/m³ or 2-3 ppb), while outdoor 36 concentrations were lower in the winter (9-10 μ g/m³ or about 5 ppb) than in the summer 37 $(35-45 \mu g/m^3 \text{ or about } 20 \text{ ppb})$. This seasonal variation in outdoor concentrations 38 39 coupled with homogeneous indoor concentrations, together with increased wintertime air

1	exchange rate due to higher indoor-outdoor temperature differences, is likely responsible
2	for the observed seasonal pattern in I/O ratios.
3	Exposures in near-road, on-road and in-vehicle microenvironments are likely to be more
4	variable and lower in magnitude than those in other microenvironments due to reaction of
5	O_3 with NO and other combustion emissions. Depending on wind direction, O_3
6	concentrations near the roadway have been found to be 20-80% of ambient
7	concentrations at sites 400 meters or more distant from roads (Section 3.6.2.1). A study
8	on patrol cars during trooper work shifts reported in-vehicle 9-h concentrations that were
9	approximately 51% of simultaneously measured roadside concentrations (mean of 11.7
10	versus 22.4 ppb) (<u>Riediker et al., 2003</u>).

4.3.3 Personal-Ambient Concentration Relationships

11	Several factors influence the relationship between personal O ₃ exposure and ambient
12	concentration. Due to the lack of indoor O_3 sources, along with reduction of ambient O_3
13	that penetrates into enclosed microenvironments, indoor and in-vehicle O ₃ concentrations
14	are highly dependent on air exchange rate and therefore vary widely in different
15	microenvironments. Ambient O3 varies spatially due to reactions with other atmospheric
16	species, especially near busy roadways where O_3 concentrations are decreased by
17	reaction with NO (Section $3.6.2.1$). This is in contrast with pollutants such as CO and
18	NO_X , which show appreciably higher concentrations near the roadway than several
19	hundred meters away (Karner et al., 2010). O_3 also varies temporally over multiple
20	scales, with generally increasing concentrations during the daytime hours, and higher O_3
21	concentrations during summer than in winter. An example of this variability is shown in
22	Figure 4-1, taken from a personal exposure study conducted by Chang et al. (2000).
23	In this figure, hourly personal exposures are seen to vary from a few ppb in some indoor
24	microenvironments to tens of ppb in vehicle and outdoor microenvironments. The
25	increase in ambient O_3 concentration during the day is apparent from the outdoor
26	monitoring data. In comparison, ambient PM _{2.5} exhibits less temporal variability over the
27	day than O ₃ , although personal exposure to PM _{2.5} also varies by microenvironment. This
28	combined spatial and temporal variability for O3 results in varying relationships between
29	personal exposure and ambient concentration.
20	
30	Correlations between personal exposure to O_3 and corresponding ambient concentrations,
31	summarized in <u>Table 4-2</u> , exhibit a wide range (generally 0.3-0.8, although both higher
32	and lower values have been reported), with higher correlations generally observed in
33	outdoor microenvironments, high building ventilation conditions, and during the summer
34	season. Low O ₃ concentrations indoors and during the winter lead to a high proportion of

1 personal exposures below the sampler detection limit, which may partially explain the 2 low correlations observed in some studies under those conditions. Studies report varying 3 correlations over a range of averaging times, with no clear trend. Ratios of personal 4 exposure to ambient concentration, summarized in Table 4-3, are generally lower in 5 magnitude (typically 0.1-0.3), and are also variable, with increasing time spent outdoors 6 associated with higher ratios. The next two subsections describe studies that have 7 reported personal-ambient correlations and slopes for a variety of seasons, locations, and 8 populations.



Note: the notation below each clock hour shows the location or activity during that hour. Source: Reprinted with permission of Air and Waste Management Association (<u>Chang et al., 2000</u>).

Figure 4-1 Variation in hourly personal and ambient concentrations of ozone and PM_{2.5} in various microenvironments during daytime hours.

1 Ozone concentrations near and below the passive sampler detection limit lead to 2 uncertainty in personal-ambient correlations and ratios. Correlations are reduced in 3 magnitude by values below the detection limit because noise obscures the underlying 4 signal in the data, while ratios tend to fluctuate widely at low concentration since small 5 changes in measured values cause large relative changes in resulting ratios. As with I/O 6 ratios, this problem is particularly acute in the non-ozone season when ambient O_3 7 concentrations are low. Improved characterization of the relationship between personal 8 exposure and ambient concentration will depend on the use of recent improved 9 monitoring techniques to accurately capture low O_3 concentrations, preferably at high 10 time resolution to facilitate evaluation of the effect of activity pattern on exposure 11 (Section 4.3.1). While data from studies using new monitoring techniques become 12 available, past studies summarized in the 2006 O₃ AQCD remain relevant to consider 13 along with more recent studies in evaluating personal-ambient concentration 14 relationships.

- 15 Personal-Ambient Correlations. Correlations between personal exposure and 16 ambient O₃ concentrations have been evaluated in several research studies, many of 17 which were conducted prior to 2005 and are discussed in the 2006 O₃ AQCD. Some 18 studies evaluated subject-specific, or longitudinal correlations, which describe multiple 19 daily measurements for a single individual. These studies indicate the inter-individual 20 variability of personal-ambient correlations. Another type of correlation is a pooled 21 correlation, which combines data from multiple individuals over multiple monitoring 22 periods (e.g., days), providing an overall indicator of the personal-ambient relationship 23 for all study subjects. A third type of correlation is a community-average correlation, 24 which correlates average exposure across all study subjects with fixed-site monitor 25 concentrations. Community-average correlations are particularly informative for 26 interpreting time-series epidemiologic studies, in which ambient concentrations are used 27 as a surrogate for community-average exposure. However, few studies report this metric; 28 this represents another opportunity for improvement of future personal exposure studies. 29 Table 4-2 summarizes studies reporting personal-ambient correlations, and the studies in 30 the table are discussed in the subsequent text.
- 31 The results of these studies generally indicate that personal exposures are moderately 32 well correlated with ambient concentrations, and that the ratio of personal exposure to 33 ambient concentration is higher in outdoor microenvironments and during the summer 34 season. In some situations, a low correlation was observed, and this may be due in part to 35 a high proportion of personal measurements below the detection limit of the personal 36 sampler. The effect of season is unclear, with mixed evidence on whether higher 37 correlations are observed during the O_3 season. Chang et al. (2000) measured hourly 38 personal exposures in multiple microenvironments and found that the pooled correlation

1	between personal exposure and ambient concentration was highest for outdoor
2	microenvironments ($r = 0.68-0.91$). In-vehicle microenvironments showed moderate to
3	high correlations (0.57-0.72). Correlations in residential indoor microenvironments were
4	very low (r = $0.05-0.09$), with moderate correlations ($0.34-0.46$) in other indoor
5	microenvironments such as restaurants and shopping malls. Liard et al. (1999) evaluated
6	community-average correlations based on 4-day mean personal O ₃ exposure
7	measurements for adults and children and found a relatively high correlation ($r = 0.83$)
8	with ambient concentrations, even though 31-82% of the personal measurements were
9	below the detection limit. Sarnat et al. (2000) studied a population of older adults in
10	Baltimore and found that longitudinal correlations between 24-h personal exposure and
11	ambient concentration varied by subject and season, with somewhat higher correlations
12	observed in this study during summer (mean = 0.20) than in winter (mean = 0.06). Some
13	evidence was presented that subjects living in well-ventilated indoor environments have
14	higher correlations than those living in poorly ventilated indoor environments, although
15	exceptions to this were also observed. Ramírez-Aguilar et al. (2008) measured 48- to
16	72-h personal exposures of four groups of asthmatic children aged 6-14 in Mexico City
17	during 1998-2000. A moderate pooled correlation ($r = 0.35$) was observed between these
18	exposures and corresponding ambient concentrations.

Study	Location	Years/Season	Population	Sample duration	Correlation	Study Type	Comment
Chang et al.	Baltimore,	Summer 1998	Older adults	1 h	0.91	Pooled	Outdoor
<u>(2000</u>)	MD	Winter 1999	_		0.77		near roadway
		Summer 1998	-		0.68	-	Outdoor
		Winter 1999	_		0.86		away from road
		Summer 1998	_		0.72		In vehicle
		Winter 1999	_		0.57		
		Summer 1998	_		0.09		Indoors-
		Winter 1999	_		0.05		residence
		Summer 1998	_		0.34		Indoors-
		Winter 1999	_		0.46		other
<u>Liard et al.</u> (1999)	Paris, France	Summer 1996	All age groups	4 day	0.83	Community- averaged	
<u>Sarnat et al.</u> (2000)	Baltimore, MD	Summer	Older adults	24 h	0.20 SD: 0.28 95% Cl: 0.06, 0.34	Longitudinal	
		Winter	_		0.06 SD: 0.34 95% Cl: -0.88, 0.24		
<u>Linn et al.</u> (1996)	Southern California	All seasons from 1992 to 1993	Children	24 h	0.61	Community- averaged	
<u>Brauer and</u> <u>Brook</u> (1997)	Vancouver, Canada	Summers 1992 and 1993	Health clinic workers	24 h	0.60	Pooled	0-25% of time outdoors
			Camp counselors	24 h	0.42	Pooled	7.5-45% of time outdoors
			Farm workers	24 h	0.64	Pooled	100% of time outdoors
<u>Ramírez-</u> Aguilar et al. (2008)	Mexico City, Mexico	December 1998- April 2000	Asthmatic children	48 h to 72 h	0.35	Pooled	
<u>Delfino et</u> al. (1996).	San Diego, California	September and October 1993	Asthmatic children	12-h	0.45 Range: 0.35-0.69	Pooled	

Table 4-2 Correlations between personal and ambient ozone concentration.

NR = not reported

1Consistent with hourly microenvironment-specific results from the Chang et al. (2000)2study described above, studies have found moderate to high personal-ambient3correlations for individuals spending time outdoors. A moderate pooled correlation of40.61 was reported between 24-h avg personal and central-site measurements by Linn et5al. (1996) for a population of southern California schoolchildren who spent an average of

1 101-136 minutes per day outdoors. The authors also report a correlation of 0.70 between 2 central-site measurements and concentrations outside the children's schools. Although 3 the average school outdoor concentration (34 ppb) was higher than the average central-4 site concentration (23 ppb) and the average personal exposure concentration was lower 5 (5 ppb) than the central-site value, the similarity between the correlations in this study 6 indicate that central-site monitor concentrations can represent personal exposures in 7 addition to representing local outdoor concentrations. A study in Vancouver, BC 8 provided another illustration of the effect of outdoor microenvironments on personal-9 ambient relationships by comparing three groups spending different amounts of time 10 outdoors: health clinic workers (0-25% of sampling period outdoors), camp counselors (7.5-45% of sampling period outdoors), and farm workers (100% of sampling period 11 12 outdoors) (Brauer and Brook, 1997). Health clinic workers and camp counselors were 13 monitored 24 h/day, while farm workers were monitored during their work shift 14 (6-14 hours). In this study, the pooled correlations between personal exposure and fixed-15 site concentration were not substantially different among the groups (r = 0.60, 0.42, and 16 0.64, respectively). The ratios of personal exposure to fixed-site monitor concentration 17 increased among the groups with increasing amount of time spent outdoors (0.35, 0.53, 18 and 0.96, respectively). This indicates that temporal variations in personal exposure to O_3 19 are driven by variations in ambient concentration, even for individuals that spend little 20 time outdoors.

Personal-Ambient Ratios. Studies indicate that the ratio between personal O₃
 exposure and ambient concentration varies widely, depending on activity patterns,
 housing characteristics, and season. Higher personal-ambient ratios are generally
 observed with increasing time spent outside, higher air exchange rate, and in seasons
 other than winter. <u>Table 4-3</u> summarizes the results of several such studies discussed in
 the 2006 O₃ AQCD together with newer studies showing the same pattern of results.

27 O'Neill et al. (2003) studied a population of shoe cleaners working outdoors in 28 Mexico City and presented a regression model indicating a 0.56 ppb increase in 6-h 29 personal exposure for each 1 ppb increase in ambient concentration. Regression analyses 30 by (2005; 2001) for 24-h data from mixed populations of children and older adults in 31 Baltimore (Sarnat et al., 2001) and Boston (Sarnat et al., 2005) found differing results 32 between the two cities, with Baltimore subjects showing a near-zero slope (0.01) during 33 the summertime while Boston subjects showed a positive slope of 0.27 ppb personal 34 exposure per 1 ppb ambient concentration. In both cities, the winter slope was near zero. 35 The low slope observed in Baltimore may have been due to differences in time spent 36 outdoors, residential ventilation conditions, or other factors. Xue et al. (2005) measured 37 6-day personal exposure of children in southern California and found that the average 38 ratio of personal exposure to ambient concentration was relatively stable throughout the

1	year at 0.3. These authors also regressed personal exposures on ambient concentration
2	after adjusting for time-activity patterns and housing characteristics and found a slope of
3	0.54 ppb/ppb, with the regression R^2 value of 0.58. Unadjusted regression slopes were
4	not presented. It should also be noted that the ratio and slope would not be expected to be
5	identified unless the intercept and other regression parameters were effectively zero.

Table 4-3Ratios of personal to ambient ozone concentration.

Study	Location	Years/Season	Population	Sample duration	Ratio ^a	Slope	Inter- cept	Study Type	Comment
Sarnat et al. (2001)	Baltimore	Summer 1998	Older adults and children	24 h	NR	0.01	1.84	Longitudinal	t-value: 1.21
		Winter 1999	Older adults, children, and individuals with COPD	-	NR	0.00	0.46	-	t-value: 0.03
Brauer and Brook (1997)	Vancouver, Canada	Summers 1992 and 1993	Health clinic workers	24 h	0.35	NR	NR	Pooled	0-25% of time outdoors
			Camp counselors		0.53	NR	NR	Pooled	7.5-45% of time outdoors
			Farm workers		0.96	NR	NR	Pooled	100% of time outdoors
<u>O'Neill et al.</u> (2003)	Mexico City, Mexico	April - July 1996	Shoe cleaners	6 h	0.40 0.37 ^b SD: 0.22	0.56 95% CI: 0.43-0.69	NR	Longitudinal	
<u>Sarnat et al.</u> (2005)	Boston	Summer	Older adults and children	24 h	NR	0.27 95% CI: 0.18-0.37	NR	Longitudinal	
		Winter	-		NR	0.04 95% CI: 0.00-0.07	NR	-	

Study	Location	Years/Season	Population	Sample duration	Ratio ^a	Slope	Inter- cept	Study Type	Comment
<u>Xue et al.</u> (2005)	Southern California	June 1995 - May 1996	Children	6 day	0.3 SD: 0.13	NR	NR	Longitudinal	
<u>Sarnat et al.</u> Steubenville, (2006a) OH	Steubenville, OH	Summer	Older adults	24 h	NR	0.15 SE: 0.02 t-value: 7.21 R ² : 0.24	NR	Longitudinal	All individuals
				NR	0.18 SE: 0.03 t-value: 7.34 R ² : 0.27	NR	-	High- ventilation	
					NR	0.08 SE: 0.04 t-value: 1.89 R ² : 0.19	NR	-	Low- ventilation
		Fall			NR	0.27 SE: 0.03 t-value: 8.64 R ² : 0.25	NR	-	All individuals
					NR	0.27 SE: 0.04 t-value: 7.38 R ² : 0.33	NR	-	High- ventilation
					NR	0.20 SE: 0.05 t-value: 3.90 R ² : 0.12	NR	-	Low- ventilation
<u>Ramírez-</u> Aguilar et al. (2008)	Mexico City, Mexico	Dec.1998- Apr. 2000	Asthmatic children	48 h to 72 h	0.23	0.17 SE: 0.02 95% CI : 0.13-0.21 p-value: 0.00		Pooled	

^a Mean value unless otherwise indicated ^b Median NR = not reported SD = standard deviation

1	A few additional studies have been published since the 2006 O ₃ AQCD comparing
2	personal exposures with ambient concentrations, and these findings generally confirm the
3	conclusions of the 2006 O ₃ AQCD that ventilation conditions, activity pattern, and season
4	may impact personal-ambient ratios. Sarnat et al. (2006a) measured 24-h personal
5	exposures for a panel of older adults in Steubenville, OH during summer and fall 2000.

1	Subjects were classified as high-ventilation or low-ventilation based on whether they
2	spent time in indoor environments with open windows. Regression of personal exposures
3	on ambient concentration found a higher slope for high-ventilation subjects compared
4	with low-ventilation subjects in both summer (0.18 versus 0.08) and fall (0.27 versus
5	0.20). Suh and Zanobetti (2010) reported an average 24-h personal exposure of 2.5 ppb as
6	compared to 24-h ambient concentration of 29 ppb for a group of individuals with either
7	recent MI or diagnosed COPD in Atlanta. A similar result was observed in Detroit, where
8	the mean 24-h personal exposure across 137 participants in summer and winter was
9	2.1 ppb, while the mean ambient concentration on sampling days was 25 ppb (Williams
10	et al., 2009b). Although no personal exposures were measured, Mcconnell et al. (2006)
11	found that average 24-h home outdoor O_3 concentrations were within 6 ppb of O_3
12	concentrations measured at central-site monitors in each of three southern California
13	communities, with a combined average home outdoor concentration of 33 ppb compared
14	to the central-site average of 36 ppb. In Mexico City, Ramírez-Aguilar et al. (2008)
15	regressed 48- to 72-h personal exposures of four groups of asthmatic children aged 6-14
16	with ambient concentrations and found slope of 0.17 ppb/ppb after adjustment for
17	distance to the fixed-site monitor, time spent outdoors, an interaction term combining
18	these two variables, and an interaction term representing neighborhood and study group.

4.3.4 Co-exposure to Other Pollutants and Environmental Stressors

19	Exposure to ambient O_3 occurs in conjunction with exposure to a complex mixture of
20	ambient pollutants that varies over space and time. Multipollutant exposure is an
21	important consideration in evaluating health effects of O ₃ since these other pollutants
22	have either known or potential health effects that may impact health outcomes due to O_3 .
23	The co-occurrence of high O_3 concentrations with high heat and humidity may also
24	contribute to health effects. This section presents data on relationships between overall
25	personal O ₃ exposure and exposure to other ambient pollutants, as well as co-exposure
26	relationships for near-road O ₃ exposure.

4.3.4.1 Personal Exposure to Ozone and Copollutants

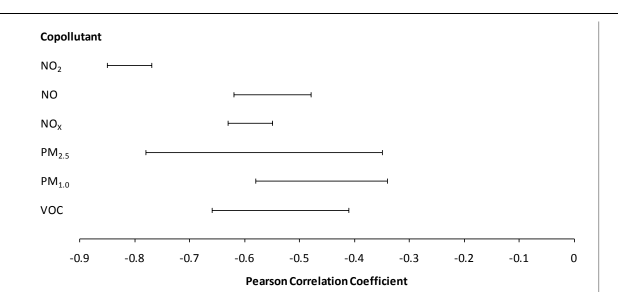
27	Personal exposure to O_3 shows variable correlation with personal exposure to other
28	pollutants, with differences in correlation depending on factors such as instrument
29	detection limit, season, city-specific characteristics, time scale, and spatial variability of
30	the copollutant. Suh and Zanobetti (2010) reported Spearman rank correlation
31	coefficients during spring and fall between 24-h avg O3 measurements and co-pollutants
32	of 0.14, 0.00, and -0.03 for $PM_{2.5}$, EC, and NO ₂ , respectively. Titration of O ₃ near

- 1 roadways is likely to contribute to the low or slightly negative correlations with the 2 traffic-related pollutants EC and NO₂. The somewhat higher correlation with PM_{2.5} may 3 reflect the influence of air exchange rate and time spent outdoors on co-exposures to 4 ambient $PM_{2.5}$ and O_3 . Overall, the copollutant correlations are quite small, which may be 5 due to the very low personal exposures observed in this study (2-3 ppb), likely to be near 6 or below the detection limit of the passive sampler over a 24-h period. Chang et al. 7 (2000) measured hourly personal exposures to PM_{2.5} and O₃ in summer and winter in 8 Baltimore, Maryland. Correlations between PM2.5 and O3 were 0.05 and -0.28 in summer 9 and winter, respectively. Results indicate personal O_3 exposures were not significantly 10 associated with personal PM2.5 exposures in either summer or winter. These non-11 significant correlations may be attributed in part to the relatively low personal O₃ 12 exposures observed in this study; in both summer and winter, the mean personal O_3 13 exposure was below the calculated limit of detection.
- 14 Studies conducted in Baltimore (Sarnat et al., 2001) and Boston (Sarnat et al., 2005) 15 found differing results for the correlation between 24-h avg personal O_3 and personal 16 $PM_{2.5}$ exposures, particularly during the winter season. Sarnat et al. (2001) found a 17 positive slope when regressing personal exposures of both total $PM_{2.5}$ (0.21) and $PM_{2.5}$ of 18 ambient origin (0.22) against personal O₃ exposures during the summer season, but 19 negative slopes (-0.05 and -0.18, respectively) during the winter season. The summertime 20 slope for personal PM_{2.5} exposure versus personal O₃ exposure was much higher for 21 children (0.37) than for adults (0.07), which may be the result of different activity 22 patterns. This team of researchers also found a positive, although higher, summer slope 23 between 24-h avg personal O_3 and personal $PM_{2.5}$ in Boston (0.72) (Sarnat et al., 2005). 24 However, the winter slope was positive (1.25) rather than negative, as in Baltimore. In 25 both cities during both seasons, there was a wide range of subject-specific correlations 26 between personal O₃ and personal PM_{2.5} exposures, with some subjects showing 27 relatively strong positive correlations (>0.75) and others showing strong negative 28 correlations (<-0.50). The median correlation in both cities was slightly positive in the 29 summer and near zero (Boston) or slightly negative (Baltimore) in the winter. These 30 results indicate the potential effects of city-specific characteristics, such as housing stock 31 and building ventilation patterns, on relationships between O₃ and copollutants.
- 32The lack of long-term exposure studies limits evaluation of long-term correlations33between O3 exposure and copollutant exposure. However, some insight may be provided34by an analysis of correlations between O3 and other criteria pollutants, such as is35provided in Section 3.6.4. Warm-season 8-h daily max O3 concentrations are generally36positively correlated with co-located 24-h avg measurements of other criteria pollutants37(Figure 3-57). Median correlations range from approximately 0.15 to 0.55 for CO, SO2,38NO2, PM10, and PM2.5, in that order. In contrast, year-round 8-h daily max O3 data show

1	negative median correlations with CO and NO ₂ , positive correlations with PM_{10} and
2	PM _{2.5} , and essentially zero correlation with SO ₂ . This reflects mostly negative
3	correlations between O_3 and all pollutants during wintertime, as shown in Figure 3-56.
4	Titration of O ₃ near roadways also likely contributes to overall negative correlations with
5	NO_2 and CO. Positive correlations between O_3 and $PM_{2.5}$ during the summertime can be
6	partly explained by meteorological conditions favoring increased formation of both
7	secondary PM and O ₃ . Strong positive correlations can influence the interpretation of
8	epidemiologic results, potentially complicating the ability to identify the independent
9	effect of a pollutant.

4.3.4.2 Near-Road Exposure to Ozone and Copollutants

10	Beckerman et al. (2008) measured both 1-week and continuous concentrations of O ₃ , NO,
11	NO ₂ , NO _X , PM _{2.5} , PM _{1.0} , and several VOCs (the BTEX compounds, MTBE, hexane, and
12	THC) in the vicinity of heavily traveled (annual average daily traffic [AADT] >340,000)
13	roadways in Toronto, Canada. Passive samplers were deployed for one week in August
14	2004. Ozone concentrations were negatively correlated with all pollutants, with the
15	exception of VOCs at one of the monitoring sites which were suspected of being
16	influenced by small area sources. Site specific correlations are given in Figure 4-2.
17	Correlations were -0.77 to -0.85 for NO ₂ , -0.48 to -0.62 for NO, and -0.55 to -0.63 for
18	NO _X . Pooled correlations using data from both sites were somewhat lower in magnitude.
19	$PM_{2.5}$ and $PM_{1.0}$ correlations were -0.35 to -0.78 and -0.34 to -0.58, respectively. At the
20	monitoring site not influenced by small area sources, O3-VOC correlations ranged from -
21	0.41 to -0.66.
22	Beckerman et al. (2008) also made on-road measurements of multiple pollutants with a
23	instrumented vehicle. Concentrations were not reported, but correlations between O3 and
24	other pollutants were negative and somewhat greater in magnitude (i.e., more negative)
25	than the near-road correlations. SO_2 , CO, and BC were measured in the mobile
26	laboratory, although not at the roadside, and they all showed negative correlations with
27	O_3 when the data were controlled for site. Correlations for continuous concentrations
28	between O ₃ and co-pollutants were somewhat lower than the 1-week correlations, except
29	for O_3 -PM _{2.5} correlations. Correlations were -0.90, -0.66, -0.77, and -0.89 for NO ₂ , NO,
30	NO_X , and $PM_{1.0}$ respectively. The continuous O_3 - $PM_{2.5}$ correlation was -0.62, which is in
31	the range of the 1-week correlation.



Source: Data from: Beckerman et al. (2008)

Figure 4-2 Correlations between 1-week concentrations of ozone and copollutants measured near roadways.

4.3.4.3 Indoor Exposure to Ozone and Copollutants

1	Ambient O ₃ that infiltrates indoors reacts with organic compounds and other chemicals to
2	form oxidized products, as described in Section $3.2.3$ as well as the 2006 O ₃ AQCD. It is
3	anticipated that individuals are exposed to these reaction products, although no evidence
4	was identified regarding personal exposures. The reactions are similar to those occurring
5	in the ambient air, as summarized in Chapter $\underline{3}$. For example, O_3 can react with terpenes
6	and other compounds from cleaning products, air fresheners, and wood products both in
7	the gas phase and on surfaces to form particulate and gaseous species, such as
8	formaldehyde (Chen et al., 2011; Shu and Morrison, 2011; Aoki and Tanabe, 2007; Reiss
9	et al., 1995b). Ozone has also been shown to react with material trapped on HVAC filters
10	and generate airborne products (Bekö et al., 2007; Hyttinen et al., 2006). Potential
11	oxygenated reaction products have been found to act as irritants (Anderson et al., 2007),
12	indicating that these reaction products may have health effects separate from those of O ₃
13	itself (Weschler and Shields, 1997). Ozone may also react to form other oxidants, which
14	then go on to participate in additional reactions. White et al. (2010) found evidence that
15	HONO or other oxidants may have been present during experiments to estimate indoor
16	OH concentrations, indicating complex indoor oxidant chemistry. Rates of these reactions
17	are dependent on indoor O_3 concentration, temperature, and air exchange rate, making
18	estimation of exposures to reaction products difficult.

4.4 **Exposure-Related Metrics**

In this section, parameters are discussed that are relevant to the estimation of exposure, but are not themselves direct measures of exposure. Time-location-activity patterns, including behavioral changes to avoid exposure, have a substantial influence on exposure and dose. Proximity of populations to ambient monitors may influence how well their exposure is represented by measurements at the monitors, although factors other than distance play an important role as well.

4.4.1 **Activity Patterns**

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7	The activity pattern of individuals is an important determinant of their exposure.
8	Variation in O3 concentrations among various microenvironments means that the amount
9	of time spent in each location, as well as the level of activity, will influence an
10	individual's exposure to ambient O ₃ . The effect of activity pattern on exposure is
11	explicitly accounted for in Equation $4-3$ by the fraction of time spent in different
12	microenvironments.

- 13 Activity patterns vary both among and within individuals, resulting in corresponding 14 variations in exposure across a population and over time. Large-scale human activity 15 databases, such as those developed for the National Human Activity Pattern Survey 16 (NHAPS) (Klepeis et al., 2001) or the Consolidated Human Activity Database (CHAD) 17 (McCurdy et al., 2000), which includes NHAPS data together with other activity study 18 results, have been designed to characterize exposure patterns among much larger 19 population subsets than can be examined during individual panel studies. The complex 20 human activity patterns across the population (all ages) are illustrated in Figure 4-3 21 (Klepeis et al., 2001), which is presented to illustrate the diversity of daily activities 22 among the entire population as well as the proportion of time spent in each 23 microenvironment. For example, about 25% of the individuals reported being outdoors or 24 in a vehicle between 2:00 and 3:00 p.m., when daily O₃ levels are peaking, although 25 about half of this time was spent in or near a vehicle, where O_3 concentrations are likely 26 to be lower than ambient concentrations.
- 27 Time spent in different locations has also been found to vary by age. Table 4-4 28 summarizes NHAPS data reported for four age groups, termed Very Young (0-4 years), 29 School Age (5-17 years), Working (18-64 years), and Retired (65+ years) (Klepeis et al., 30 1996). The working population spent the least time outdoors, while the school age 31 population spent the most time outdoors. NHAPS respondents aged 65 and over spent 32 somewhat more time outdoors than adults aged 18-64, with a greater fraction of time

1	spent outdoors at a residence. Children aged 0-4 also spent most of their outdoor time in a
2	residential outdoor location. On average, the fraction of time spent outdoors by school
3	age respondents was 2.62 percentage points higher than working respondents,
4	corresponding to approximately 38 minutes more time outdoors per day. Although not
5	accounting for activity level, this increased time spent outdoors is consistent with
6	evidence in Chapter <u>8</u> suggesting that younger and older age groups are more at risk for
7	O ₃ -related health effects.

Table 4-4Mean fraction of time spent in outdoor locations by various age
groups in the NHAPS study

Age Group	Residential-Outdoor	Other Outdoor	Total Outdoors	
0-4 yr	5.38%	0.96%	6.34%	
5-17 yr	5.05%	2.83%	7.88%	
18-64 yr	2.93%	2.33%	5.26%	
65+ yr	4.48%	1.27%	5.75%	

Source: Data from Klepeis et al. (1996).

8	Together with location, exertion level is an important determinant of exposure. Table 4-5
9	summarizes ventilation rates for different age groups at several levels of activity as
10	presented in Table 6-2 of the EPA's Exposure Factors Handbook (U.S. EPA, 2011b).
11	Most of the age-related variability is seen for moderate and high intensity activities,
12	except for individuals under 1 year. For moderate intensity, ventilation rate increases with
13	age through childhood and adulthood until age 61, after which a moderate decrease is
14	observed. Ventilation rate is most variable for high intensity activities. Children aged 1 to
15	<11 years have ventilation rates of approximately 40 L/min, while children aged 11+ and
16	adults have ventilation rates of approximately 50 L/min. The peak is observed for the 51
17	to <61 age group, at 53 L/min, with lower ventilation rates for older adults.

Age Group	Sleep or Nap	Sedentary/Passive	Light Intensity	Moderate Intensity	High Intensity
Birth to <1 yr	3.0	3.1	7.6	14	26
1 to <2 yr	4.5	4.7	12	21	38
2 to <3 yr	4.6	4.8	12	21	39
3 to <6 yr	4.3	4.5	11	21	37
6 to <11 yr	4.5	4.8	11	22	42
11 to <16 yr	5.0	5.4	13	25	49
16 to <21 yr	4.9	5.3	12	26	49
21 to <31 yr	4.3	4.2	12	26	50
31 to <41 yr	4.6	4.3	12	27	49
41 to <51 yr	5.0	4.8	13	28	52
51 to <61 yr	5.2	5.0	13	29	53
61 to <71 yr	5.2	4.9	12	26	47
71 to <81 yr	5.3	5.0	12	25	47
81+ yr	5.2	4.9	12	25	48

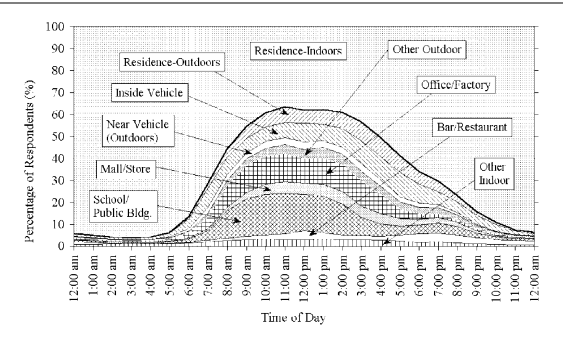
Table 4-5Mean ventilation rates (L/min) at different activity levels for
different age groups.

Source: Data from *Exposure Factors Handbook* (U.S. EPA, 2011b).

1A dramatic increase in ventilation rate occurs as exercise intensity increases. For children2and adults <31 years, high intensity activities result in nearly double the ventilation rate</td>3for moderate activity, which itself is nearly double the rate for light activity. Children4have other important differences in ventilation compared to adults. As discussed in5Chapter 5, children tend to have a greater oral breathing contribution than adults, and6they breathe at higher minute ventilations relative to their lung volumes. Both of these7factors tend to increase dose normalized to lung surface area.

8 Longitudinal activity pattern information is also an important determinant of exposure, as 9 different people may exhibit different patterns of time spent outdoors over time due to 10 age, gender, employment, and lifestyle-dependent factors. These differences may 11 manifest as higher mean exposures or more frequent high-exposure episodes for some 12 individuals. The extent to which longitudinal variability in individuals contributes to the 13 population variability in activity and location can be quantified by the ratio of between-14 person variance to total variance in time spent in different locations and activities (the 15 intraclass correlation coefficient, ICC). Xue et al. (2004) quantified ICC values in time-16 activity data collected by Harvard University for 160 children aged 7–12 years in 17 Southern California (Geyh et al., 2000). For time spent outdoors, the ICC was 18 approximately 0.15, indicating that 15% of the variance in outdoor time was due to

between-person differences. The ICC value might be different for other population groups.



Source: Reprinted with permission of Nature Publishing Group (Klepeis et al., 2001).

Figure 4-3 Distribution of time that NHAPS respondents spent in ten microenvironments based on smoothed 1-min diary data.

3 The EPA's National Exposure Research Laboratory (NERL) ha	as consolidated the most
4 important human activity databases into one comprehensive da	tabase called the
5 Consolidated Human Activity Database (CHAD). The current v	version of CHAD contains
6 data from nineteen human activity pattern studies (including N	HAPS), which were
7 evaluated to obtain over 33,000 person-days of 24-h human act	ivities in CHAD
8 (<u>McCurdy et al., 2000</u>). The surveys include probability-based	recall studies conducted
9 by EPA and the California Air Resources Board, as well as real	-time diary studies
10 conducted in individual U.S. metropolitan areas using both prob	bability-based and
11 volunteer subject panels. All ages of both genders are represent	ed in CHAD. The data for
12 each subject consist of one or more days of sequential activities	s, in which each activity is
13 defined by start time, duration, activity type, and microenviron	ment classification
14 (i.e., location). Activities vary from one minute to one hour in c	luration, with longer
15 activities being subdivided into clock-hour durations to facilitat	te exposure modeling.
16 CHAD also provides information on the level of exertion assoc	iated with each activity,

1 2 which can be used by exposure models, including the APEX model (Section 4.5.3), to estimate ventilation rate and pollutant dose.

4.4.2 Ozone Averting Behavior

3 Individuals can reduce their exposure to O_3 by altering their behaviors, such as by staying 4 indoors, being active outdoors when air quality is better, and by reducing activity levels 5 or time spent being active outdoors on high- O_3 days. To assist the public in avoiding 6 exposure to air pollution on days with high pollutant concentrations, EPA has developed 7 an information tool known as the Air Quality Index (AQI) to provide information to the 8 public on ambient levels of pollutants and the potential for individuals to experience 9 health effects (U.S. EPA, 2011a). The AQI describes the potential for health effects from 10 O₃ (and other individual pollutants) in six color-coded categories of air-quality, ranging 11 from good (green), moderate (yellow), unhealthy for sensitive groups (orange), unhealthy 12 (red), very unhealthy (purple), and hazardous (maroon). The levels are associated with 13 descriptors of the likelihood of health effects and the populations most likely to be 14 affected. For example, the orange level indicates that the general population is not likely 15 to be at risk, but susceptible groups may experience health effects. These advisories 16 explicitly state that children, older adults, people with lung disease, and those who are 17 active outdoors may be at greater risk from exposure to air pollution. Forecasted and 18 actual conditions typically are reported to the public during high-O₃ months through local 19 media outlets, using various versions of this air-quality categorization scheme. People are 20 advised to change their behavior to reduce exposure depending on predicted O_3 21 concentrations and the likelihood of risk. Behavioral recommendations include being 22 active outdoors when air quality is better, and reducing activity levels or the time spent 23 being active outdoors on high- O_3 days. Staying indoors to reduce exposure is only 24 recommended when the AQI is at or above the very unhealthy range.

25 Evidence of individual averting behaviors in response to advisories has been found in 26 several studies, especially for potentially susceptible populations, such as children, older 27 adults, and asthmatics. Reduced time spent outdoors was reported in an activity diary 28 study in 35 U.S. cities (Mansfield et al., 2006), which found that asthmatic children who 29 spent at least some time outdoors reduced their total time spent outdoors by an average of 30 30 min on a code red O_3 day relative to a code green, yellow, or orange day; however, the 31 authors noted that there was appreciable variation in both the overall amount of time 32 spent outdoors and the reduction in outdoor time on high ozone days among asthmatic 33 children. Bresnahan et al. (1997) examined survey data collected during 1985-86 from a 34 panel of adults in the Los Angeles area, many of whom had compromised respiratory 35 function, by an averting behavior model. A regression analysis indicated that individuals

1	with smog-related symptoms spent about 12 minutes less time outdoors over a two-day
2	period for each 10 ppb increase in O_3 concentration above 120 ppb. Considering that the
3	average daily maximum O_3 concentration at the time was approximately 180 ppb on days
4	when the then-current standard (1-h max of 120 ppb) was exceeded, this implies that
5	those individuals spent about 40 minutes less time outside per day on a typical high O_3
6	day compared to days with O_3 concentrations below the standard. However, the behavior
7	was not specifically linked to exceedances or air quality alerts.
8	The fraction of individuals who reduce time spent outdoors, or restrict their children's
9	outdoor activity, has been found to vary based on health status. In the <u>Bresnahan et al.</u>
10	(1997) study, 40 percent of respondents reported staying indoors on days when air quality
10	was poor. Individuals who reported experiencing smog-related symptoms were more
12	likely to take the averting actions, although the presence of asthma or other chronic
13	respiratory conditions did not have a statistically significant effect on behavior. A study
13	of parents of asthmatic children (<u>McDermott et al., 2006</u>) suggests that parents are aware
15	of the hazard of outdoor air pollution and the official alerts designed to protect them and
16	their children. It also suggests that a majority of parents (55%) comply with
17	recommendations of the alerts to restrict children's outdoor activity, with more parents of
18	asthmatics reporting awareness and responsiveness to alerts. However, only 7% of all
19	parents complied with more than one-third of the advisories issued (McDermott et al.,
20	2006). Wen et al. (2009) analyzed data from the 2005 Behavioral Risk Factor
20	Surveillance System (BRFSS) and indicated that people with asthma are about twice as
21 22	likely as people without asthma to reduce their outdoor activities based on either media
22	alerts of poor air quality (31% vs. 16%) or individual perception of air quality (26% vs.
23	12%). Respondents who had received advice from a health professional to reduce outdoor
25	activity when air quality is poor were more likely to report a reduction based on media
26	alerts, both for those with and without asthma. In a study of randomly selected
20 27	individuals in Houston, TX and Portland, OR, <u>Semenza et al. (2008)</u> found that a
28	relatively small fraction of survey respondents (9.7% in Houston, 10.5% in Portland)
28	changed their behaviors during poor air quality episodes. This fraction is appreciably
30	lower than the fraction reported for people with asthma in the <u>Wen et al. (2009)</u> study,
30	although it is similar to the fraction reported in that study for those without asthma. Most
31	of the people in the <u>Semenza et al. (2008</u>) study reported that their behavioral changes
33	were motivated by self-perception of poor air quality rather than an air quality advisory.
33 34	It should be noted that the <u>McDermott et al. (2006)</u> , <u>Wen et al. (2009</u>), and <u>Semenza et al.</u>
35	(2008) studies evaluated air quality in general and therefore are not necessarily specific to
35 36	(2008) studies evaluated air quanty in general and therefore are not necessarily specific to O_3 .
30	03.
37	Commuting behavior does not seem to change based on air quality alerts. A study in the
38	Atlanta area showed that advisories can raise awareness among commuters but do not

1	necessarily result in a change in an individual's travel behavior (Henry and Gordon,
2	2003). This finding is consistent with a survey for 1000 commuters in Denver, Colorado,
3	which showed that the majority (76%) of commuters heard and understood the air quality
4	advisories, but did not alter their commuting behavior (Blanken et al., 2001).

- 5 Some evidence is available for other behavioral changes in response to air quality alerts. 6 Approximately 40 percent of the respondents in the Los Angeles study by Bresnahan et 7 al. (1997) limited or rearranged leisure activities, and 20 percent increased use of air 8 conditioners. As with changes in time spent outdoors, individuals who reported 9 experiencing smog-related symptoms, but not those with asthma or chronic respiratory 10 conditions, were more likely to take the averting actions. Other factors influencing 11 behavioral changes, such as increased likelihood of averting behavior among high school 12 graduates, are also reported in the study. In a separate Southern California study, 13 attendance at two outdoor facilities (i.e., a zoo and an observatory) was reduced by 14 6-13% on days when smog alerts were announced, with greater decreases observed 15 among children and older adults (Neidell, 2010, 2009).
- 16The studies discussed in this section indicate that averting behavior is dependent on17several factors, including health status and lifestage. People with asthma and those18experiencing smog-related symptoms reduce their time spent outdoors and are more19likely to change their behavior than those without respiratory conditions. Children and20older adults appear more likely to change their behavior than the general population.21Commuters, even when aware of air quality advisories, tend not to change their22commuting behavior.

4.4.3 Population Proximity to Fixed-Site Ozone Monitors

23	The distribution of O_3 monitors across urban areas varies between cities (Section <u>3.6.2.1</u>),
24	and the population living near each monitor varies as well. Monitoring sites in rural areas
25	are generally located in national or state parks and forests, and these monitors may be
26	relevant for exposures of exercising visitors as well as those who live in similar locations.
27	They also serve as an important source of data for evaluating ecological effects of O_3
28	(Chapter 9). Rural monitors tend to be less affected than urban monitors by strong and
29	highly variable anthropogenic sources of species participating in the formation and
30	destruction of O ₃ (e.g., onroad mobile sources) and more highly influenced by regional
31	transport of O_3 or O_3 precursors (Section <u>3.6.2.2</u>). This may contribute to less diel
32	variability in O_3 concentration than is observed in urban areas.
33	A variety of factors determine the siting of the O ₃ monitors that are part of the SLAMS
34	network reporting to AQS. As discussed in Section 3.5.6, the number and location of

1	required O ₃ monitors in an urban area depend on O ₃ concentration and population, among
2	other factors. Areas classified as serious, severe, or extreme nonattainment have
3	additional monitoring requirements. Generally, high-O3 urban areas with a population of
4	50,000 or greater are required to have at least one monitor; in low- or moderate-
5	concentration areas, the minimum population for a required monitor is 350,000. Most
6	large U.S. cities have several monitors, as shown in Figure 3-76 through Figure 3-95.
7	As an illustration of the location of O_3 monitors and their concentrations with respect to
8	population density, Figure 4-4 through Figure 4-6 present this information for Atlanta,
9	Boston, and Los Angeles, the three cities selected for detailed analysis in Chapter 3. They
10	represent a cross-section with respect to geographic distribution, O ₃ concentration,
11	layout, geographic features, and other factors. The maps show the location of O_3
12	monitors, identified by the same letters as in Chapter $\underline{3}$ to facilitate intercomparisons,
13	along with the 2007-09 mean 8-h daily max O_3 concentration for perspective on the
14	variation in O_3 concentration across the urban area. Population density at the census
15	block group level is also presented on the maps.

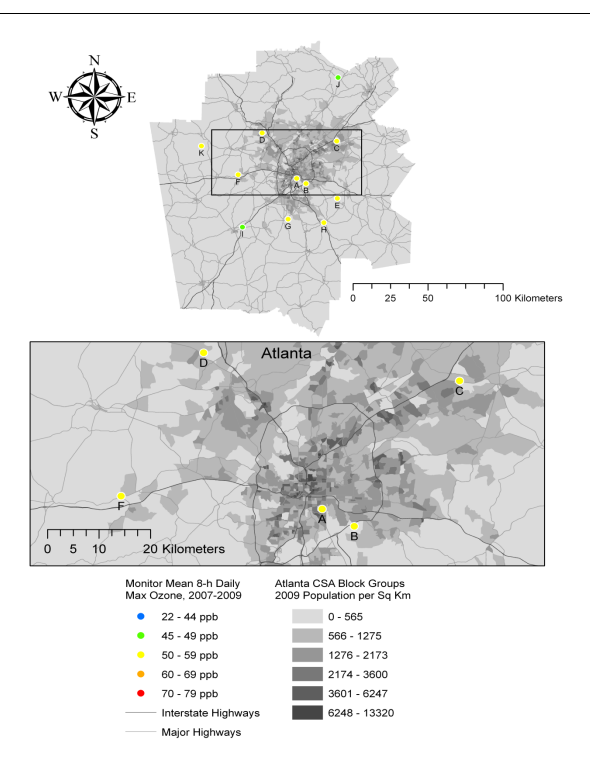


Figure 4-4 Map of the Atlanta CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.

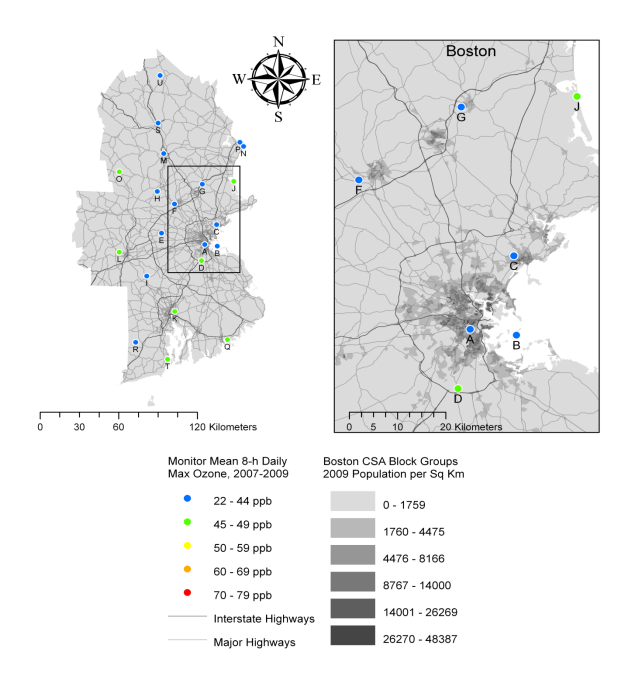


Figure 4-5 Map of the Boston CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.

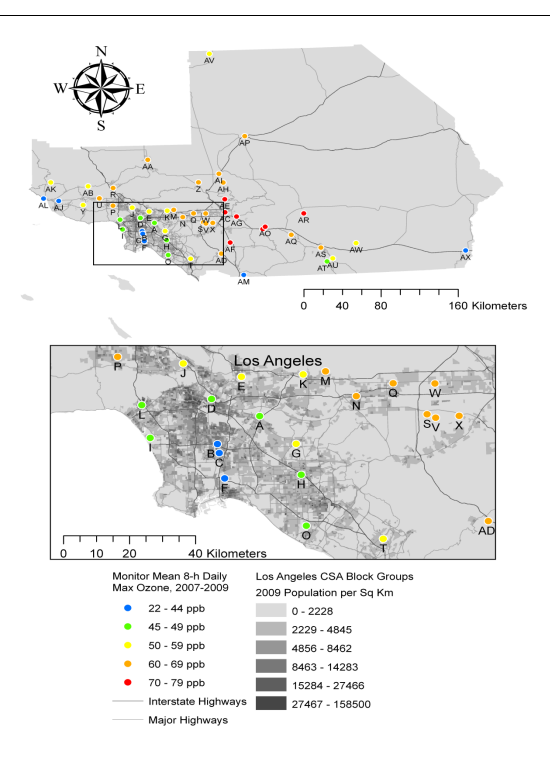


Figure 4-6 Map of the Los Angeles CSA including ozone monitor locations and major roadways with respect to census block group population density estimates for 2009.

1 Similarities and differences are apparent among the cities. The spatial distribution of 2 monitor locations in Atlanta and Boston is similar, with one site (site A) near the high 3 population density area and other monitors in surrounding areas of lower population 4 density. In Atlanta, the monitors near the city all have similar concentrations, while 5 somewhat lower concentrations are observed at sites I and J, which are located >50 km 6 from the city center. Boston shows a different spatial concentration pattern, with some 7 low and some high concentrations in urban and less-populated areas. The differences in 8 spatial concentration profiles between the two cities may be due to more consistent 9 terrain in Atlanta compared with Boston, which has a coastline, along with the downwind 10 influence of New York and other northeastern cities contributing to concentration 11 variability.

- 12 Los Angeles has a much more complex spatial pattern of monitors, population, and 13 geography. There are a large number of monitors located in multiple levels of population 14 density across the entire CSA, which includes substantial rural areas. Most monitors are 15 near at least moderate population density areas, but there are some high-density areas 16 without a monitor. Concentrations increase in a somewhat radial or west-east pattern 17 from the city, with lower concentrations near the port of Long Beach (monitors B, C, and 18 F). The highest concentrations are located near the San Bernadino forest (e.g., monitors 19 AG, AO, and AR), which have lower population density, but more potential for ecological 20 impacts. Low concentrations in highly populated areas near the coast likely reflect 21 titration by NO_X and other atmospheric constituents, while high downwind 22 concentrations reflect lack of local sources and increased photochemical processing time.
- 23 The location of these monitors relative to the location of dense population centers varies 24 among urban areas. NCore sites, a subset of the overall monitoring network, are designed 25 with population exposure as a monitoring objective, and the monitoring requirements in 26 40 CFR Part 58, Appendix D include population density as one of several factors that 27 would be considered in designing the O_3 monitoring program for an area. At least one site 28 for each MSA is designed to be a maximum concentration site, which could presumably 29 represent the location with the maximum exposure potential in the city. Sites may also be 30 required upwind and downwind of high-concentration urban areas.
- 31All three cities have some high population density areas without an O3 monitor. The32siting considerations for NCore monitors generally target the neighborhood (0.5-4 km) or33urban (4-50 km) scale to provide representative concentrations throughout the34metropolitan area; however, a middle-scale (0.1-0.5 km) site may be acceptable in cases35where the site can represent many such locations throughout a metropolitan area. In other36words, a monitor could potentially represent exposures in other similar areas of the city if37land use and atmospheric chemistry conditions are similar. This is supported by the

correlation analyses in Chapter <u>3</u>. For example, in Los Angeles, monitors H and L are
 located in medium-density areas and show moderately high correlation (0.78), although
 they are some 50 km apart.

4 Although proximity to a monitor does not determine the degree to which that monitor 5 represents an individual's ambient exposure, it is one indicator. One way to calculate 6 monitor representativeness is to calculate the fraction of the urban population living 7 within a certain radius of a monitor. Table 4-6 presents the fraction of the population in 8 selected cities living within 1, 5, 10, and 20 km of an O₃ monitor. Values are presented 9 for both total population and for those under 18 years of age, a potentially susceptible 10 population to the effects of O_3 . The data indicate that relatively few people live within 1 km of an O₃ monitor, while nearly all of the population in most cities lives within 11 12 20 km of a monitor. Looking at the results for a 5-km radius, corresponding roughly to 13 the neighborhood scale (Section 3.5.6.1), generally 20-30% of the population lives within 14 this distance from an O_3 monitor. Some cities have a greater population in this buffer, 15 such as Salt Lake City, while others have a lower percentage, such as Minneapolis and 16 Seattle. Percentages for children are generally similar to the total population, with no 17 clear trend.

18 Another approach is to divide the metropolitan area into sectors surrounding each 19 monitor such that every person in the sector lives closer to that monitor than any other. 20 This facilitates calculation of the fraction of the city's population represented (according 21 to proximity) by each monitor. In Atlanta, for example, the population fraction 22 represented by each of the 11 monitors in the city ranged from 2.9-22%. The two 23 monitors closest to the city center (sites A and B on Figure 4-4) accounted for 16% and 24 8% of the population, respectively. Site B has two listed monitoring objectives, highest 25 concentration and population exposure. The other monitor in Atlanta with a listed 26 objective of highest concentration is Site C, which represents the largest fraction of the 27 population (22%). The eight monitors with a primary monitoring objective of population 28 exposure account for 2.9-17% of the population per monitor.

Table 4-6	Fraction of the 2009 population living within a specified distance of
	an ozone monitor in selected U.S. cities.

City	Population		Within 1 km		Within 5 km		Within 10 km		Within 20 km	
	Total	<18 yr	Total	<18 yr	Total	<18 yr	Total	<18 yr	Total	<18 yr
Atlanta CSA	5,901,670	1,210,932	0.3%	0.3%	8%	9%	28%	29%	75%	77%
Baltimore CSA	8,421,016	1,916,106	1.3%	1.1%	25%	24%	57%	55%	89%	89%
Birmingham CSA	1,204,399	281,983	1.4%	1.6%	22%	24%	56%	59%	73%	74%
Boston CSA	7,540,533	1,748,918	0.9%	0.9%	17%	16%	49%	47%	85%	85%
Chicago CSA	9,980,113	2,502,454	1.5%	1.5%	28%	29%	63%	65%	89%	91%
Dallas CSA	6,791,942	1,530,877	0.4%	0.4%	13%	13%	45%	44%	87%	87%
Denver CSA	3,103,801	675,380	1.7%	1.6%	35%	36%	66%	68%	92%	93%
Detroit CSA	5,445,448	1,411,875	0.8%	0.9%	15%	17%	42%	44%	77%	78%
Houston CSA	5,993,633	1,387,851	1.5%	1.8%	26%	28%	54%	57%	83%	84%
Los Angeles CSA	18,419,720	4,668,441	1.6%	1.7%	28%	29%	77%	79%	98%	98%
Minneapolis CSA	3,652,490	872,497	0.3%	0.3%	5%	4%	16%	16%	57%	56%
New York CSA	22,223,406	5,284,875	1.5%	1.7%	23%	23%	51%	50%	91%	91%
Philadelphia CSA	6,442,836	1,568,878	0.9%	1.0%	22%	24%	55%	56%	89%	89%
Phoenix CBSA	4,393,462	873,084	2.0%	2.4%	35%	41%	74%	79%	96%	97%
Pittsburgh CSA	2,471,403	563,309	1.5%	1.4%	22%	21%	52%	50%	88%	88%
Salt Lake City CSA	1,717,045	460,747	3.0%	3.0%	41%	38%	79%	79%	95%	95%
San Antonio CBSA	2,061,147	484,473	0.5%	0.5%	12%	12%	42%	43%	78%	80%
San Francisco CSA	7,497,443	1,675,711	2.6%	2.9%	41%	40%	81%	81%	98%	98%
Seattle CSA	4,181,278	918,309	0.3%	0.3%	5%	5%	18%	16%	43%	39%
St. Louis CSA	2,914,754	720,746	1.3%	1.5%	17%	18%	52%	53%	80%	82%

1 Atlanta population fractions for children (<18 years of age) are similar to those for the 2 general population, but other populations show a different pattern of monitor 3 representativeness. Older adults (age 65 and up) were somewhat differently distributed 4 with respect to the monitors, with most monitors showing a difference of more than a 5 percentage point compared to the general population. Based on 2000 population data, the 6 fraction of older adults closest to the two city center monitors (A and B) was 4% higher 7 and 2% lower, respectively, than the fraction for the population as a whole. Site C 8 showed the highest differential, with 21% of the total population but only 15% of the 9 older adult population. This indicates the potential for monitors to differentially represent 10 potentially susceptible populations.

4.5 Exposure Modeling

1	In the absence of personal exposure measurements, modeling techniques are used to
2	estimate exposures, particularly for large populations for which individual-level
3	measurements would be impractical. Model estimates may be used as inputs to
4	epidemiologic studies or as stand-alone assessments of the level of exposure likely to be
5	experienced by a population under certain air quality conditions. This section describes
6	approaches used to improve exposure estimates, including concentration surface
7	modeling, which calculates local outdoor concentrations over a geographic area; air
8	exchange rate modeling, which estimates building ventilation based on housing
9	characteristics and meteorological parameters; and microenvironment-based exposure
10	modeling, which combines air quality data with demographic information and activity
11	pattern simulations to estimate time-weighted exposures based on concentrations in
12	multiple microenvironments. These models each have strengths and limitations, as
13	summarized in <u>Table 4-7</u> . The remainder of this section provides more detail on specific
14	modeling approaches, as well as results of applying the models.

Model Type	Model	Description	Strengths	Limitations
Concentration Surface	Spatial Interpolation (e.g., Inverse Distance Weighting, Kriging)	Measured concentrations are interpolated across an area to yield local outdoor concentration estimates	High concentration resolution; uses available data; requires low to moderate resources for implementation	Spatial heterogeneity not fully captured; a single high-concentration monitor can skew results; no location-activity information
	Chemistry- transport (e.g., CMAQ)	Grid-based O ₃ concentrations are calculated from precursor emissions, meteorology, and atmospheric chemistry and physics	First-principles characterization of physical and chemical processes influencing O ₃ formation	Grid cell resolution; resource-intensive; no location-activity information
	Land-use regression (LUR)	Merges concentration data with local-scale variables such as land use factors to yield local concentration surface	High concentration resolution	Reactivity and small-scale spatial variability of O_3 ; location-specific, limiting generalizability; no location-activity information
Air Exchange Rate	Mechanistic (LBL, LBLX)	Uses database on building leakage tests to predict AER based on building characteristics and meteorological variables (including natural ventilation in LBLX)	Physical characterization of driving forces for air exchange	Moderate resource requirement; no location- activity information
	Empirical	Predicts AER based on factors such as building age and floor area	Low input data requirements	Cannot account for meteorology; no location- activity information
Integrated Microenvironmental Exposure and Dose	Population (APEX, SHEDS)	Stochastic treatment of air quality data, demographic variables, and activity pattern to generate estimates of microenvironmental concentrations, exposures, and doses	Probabilistic estimates of exposure and dose distributions for specific populations; consideration of nonambient sources; small to moderate uncertainty for exercising asthmatic children (APEX)	Resource-intensive; evaluation with measured exposures; underestimation of multiple high-exposure events in an individual (APEX)

Table 4-7Characteristics of exposure modeling approaches.

4.5.1 Concentration Surface Modeling

1	One approach to improve exposure estimates in urban areas involves construction of a
2	concentration surface over a geographic area, with the concentration at locations between
3	monitors estimated using a model to compensate for missing data. The calculated O_3
4	concentration surface can then be used to estimate exposures outside residences, schools,
5	workplaces, roadways, or other locations of interest. This technique does not estimate
6	exposure directly because it does not account for activity patterns or concentrations in
7	different microenvironments. This is an important consideration in the utility of these
8	methods for exposure assessment; while improved local-scale estimates of outdoor
9	concentrations may contribute to better assignment of exposures, information on activity

- 1 patterns is needed to produce estimates of personal exposure. There are three main types 2 of approaches: spatial interpolation of measured concentrations; statistical models using 3 meteorological variables, pollutant concentrations, and other predictors to estimate 4 concentrations at receptors in the domain; and rigorous first-principle models, such as 5 chemistry-transport models or dispersion models incorporating O₃ chemistry. Some 6 researchers have developed models that combine these techniques. The models may be 7 applied over urban, regional, or national spatial scales, and can be used to estimate daily 8 concentrations or longer-term averages. This discussion will focus on short-term 9 concentrations estimated across urban areas.
- 10The 2006 O3 AQCD discussed concentration surface models, focusing on chemistry-11transport models as well as geospatial and spatiotemporal interpolation techniques (e.g.,12Christakos and Vyas, 1998a, b; Georgopoulos et al., 1997). Recent research has13continued to refine and extend the modeling approaches. A few recent papers have14compared different approaches for the same urban area.
- 15 Marshall et al. (2008) compared four spatial interpolation techniques for estimation of O₃ 16 concentrations in Vancouver, BC. The investigators assigned a daily average O₃ 17 concentration to each of the 51,560 postal-code centroids using one of the following 18 techniques: (1) the concentration from the nearest monitor within 10 km; (2) the average 19 of all monitors within 10 km; (3) the inverse-distance-weighted (IDW) average of all 20 monitors in the area; and (4) the IDW average of the 3 closest monitors within 50 km. 21 Method 1 (the nearest-monitor approach) and Method 4 (IDW-50 km) had similar mean 22 and median estimated annual- and monthly-average concentrations, although the 10th-23 90th percentile range was smaller for IDW-50. This is consistent with the averaging of 24 extreme values inherent in IDW methods. The Pearson correlation coefficient between 25 the two methods was 0.93 for monthly-average concentrations and 0.78 for annual-26 average concentrations. Methods 2 and 3 were considered sub-optimal and were excluded 27 from further analysis. In the case of Method 2, a single downtown high-concentration 28 monitor skewed the results in the vicinity, partially as a result of the asymmetric layout of 29 the coastal city of Vancouver. Method 3 was too spatially homogenous because it 30 assigned most locations a concentration near the regional average, except for locations 31 immediately adjacent to a monitoring site. CMAQ concentration estimates using a 32 4 km×4 km grid were also compared to the interpolation techniques in this study. Mean 33 and median concentrations from CMAQ were approximately 50% higher than Method 1 34 and Method 4 estimates for both annual and monthly average concentrations. This may 35 be due in part to the CMAQ grid size, which was too coarse to reveal near-roadway 36 decrements in O₃ concentration due to titration by NO. The IQR for the annual average 37 was similar between CMAQ and the interpolation techniques, but the monthly average 38 CMAQ IQR was approximately twice as large, indicating a seasonal effect.

1	Bell (2006) compared CMAQ estimates for northern Georgia with nearest-monitor and
2	spatial interpolation techniques, including IDW and kriging. The area-weighted
3	concentration estimates from CMAQ indicated areas of spatial heterogeneity that were
4	not captured by approaches based on the monitoring network. The author concluded that
5	some techniques, such as spatial interpolation, were not suitable for estimation of
6	exposure in certain situations, such as for rural areas. Using the concentration from the
7	nearest monitor resulted in an overestimation of exposure relative to model estimates.
8	Land use regression (LUR) models have been developed to estimate levels of air
9	pollutants, predominantly NO ₂ , as a function of several land use factors, such as land use
10	designation, traffic counts, home heating usage, point source strength, and population
11	density (Ryan and LeMasters, 2007; Gilliland et al., 2005; Briggs et al., 1997). LUR,
12	initially termed regression mapping (Briggs et al., 1997), is a regression derived from
13	monitored concentrations as a function of data from a combination of the land use
14	factors. The regression is then used for predicting concentrations at multiple locations
15	based on the independent variables at those particular locations without monitors. Hoek
16	et al. (2008) warn of several limitations of LUR, including distinguishing real
17	associations between pollutants and covariates from those of correlated copollutants,
18	limitations in spatial resolution from monitor data, applicability of the LUR model under
19	changing temporal conditions, and introduction of confounding factors when LUR is used
20	in epidemiologic studies. These limitations may partially explain the lack of LUR models
21	that have been developed for O_3 at the urban scale. Brauer et al. (2008) evaluated the use
22	of LUR and IDW-based spatial-interpolation models in epidemiologic analyses for
23	several different pollutants in Vancouver, BC and suggested that LUR is appropriate for
24	directly-emitted pollutants with high spatial variability, such as NO and BC, while IDW
25	is appropriate for secondary pollutants such as NO ₂ and PM _{2.5} with less spatial variability.
26	Although O ₃ is also a secondary pollutant, its reactivity and high small-scale spatial
27	variability near high-traffic roadways indicates this conclusion may not apply for O_3 .
28	At a much larger spatial scale, EU-wide, Beelen et al. (2009) compared a LUR model for
29	O ₃ with ordinary kriging and universal kriging, which incorporated meteorological,
30	topographical, and land use variables to characterize the underlying trend. The LUR
31	model performed reasonably well at rural locations (5-km resolution), explaining a higher
32	percentage of the variability ($R^2 = 0.62$) than for other pollutants. However, at the urban
33	scale (1-km resolution), only one variable was selected into the O ₃ LUR model
34	(high-density residential land use), and the R^2 value was very low (0.06). Universal
35	kriging was the best method for the large-scale composite EU concentration map, for O ₃
36	as well as for NO ₂ and PM ₁₀ , with an R^2 value for O ₃ of 0.70. The authors noted that
37	these methods were not designed to capture spatial variation in concentrations that are

known to occur within tens of meters of roadways (Section 3.6.2.1), which could partially explain poor model performance at the urban scale.

- 3 Titration of O_3 with NO emitted by motor vehicles tends to reduce O_3 concentrations near 4 roadways. Mcconnell et al. (2006) developed a regression model to predict residential O_3 5 concentrations in southern California using estimates of residential NO_x calculated from 6 traffic data with the CALINE4 line source dispersion model. The annual average model results were well-correlated ($R^2 = 0.97$) with multi-year average monitoring data. The 7 authors estimated that local traffic contributes 18% of NO_x concentrations measured in 8 9 the study communities, with the remainder coming from regional background. Their 10 regression model indicates that residential NO_X reduces residential O_3 concentrations by 0.51 ppb (SE 0.11 ppb) O₃ per 1 ppb NO_X, and that a 10th-90th percentile increase in 11 local NO_X results in a 7.5 ppb decrease in local O₃ concentrations. This intra-urban 12 13 traffic-related variability in O₃ concentrations suggests that traffic patterns are an 14 important factor in the relationship between central site monitor and residential O_3 , and 15 that differences in traffic density between the central site monitor and individual homes 16 could result in either an overestimate or underestimate of residential O₃.
- 17 A substantial number of researchers have used geostatistical methods and chemistry-18 transport models to estimate O_3 concentrations at urban, regional, national, and 19 continental scales, both in the U.S. and in other countries (Section 3.3). In addition to 20 short-term exposure assessment for epidemiologic studies, such models may also be used 21 for long-term exposure assessment, O_3 forecasts, or evaluating emission control 22 strategies. However, as discussed at the beginning of this section, caveats regarding the 23 importance of activity pattern information in estimating personal and population exposure 24 should be kept in mind.

4.5.2 Residential Air Exchange Rate Modeling

25	The residential air exchange rate (AER), which is the airflow into and out of a home, is
26	an important mechanism for entry of ambient O_3 . As described in Section <u>4.3.2</u> , the
27	indoor-outdoor relationship is greatly affected by the AER. Since studies show that
28	people spend approximately 66% of their time indoors at home (Leech et al., 2002;
29	Klepeis et al., 2001), the residential AER is a critical parameter for exposure models,
30	such as APEX, SHEDS, and EMI (discussed in Section 4.5.3) (U.S. EPA, 2011c, 2009b;
31	Burke et al., 2001). Since the appropriate AER measurements may not be available for
32	exposure models, mechanistic and empirical (i.e., regression-based) AER models can be
33	used for exposure assessments. The input data for the AER models can include building
34	characteristics (e.g., age, number of stories, wind sheltering), occupant behavior

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1	(e.g., window opening), climatic region, and meteorology (e.g., local temperature and
2	wind speed). Mechanistic AER models use these meteorological parameters to account
3	for the physical driving forces of the airflows due to pressure differences across the
4	building envelope from wind and indoor-outdoor temperature differences (ASHRAE,
5	2009). Empirical AER models do not consider the driving forces from the wind and
6	indoor-outdoor temperature differences. Instead, a scaling constant can be used based on
7	factors such as building age and floor area (Chan et al., 2005b).
8	Single-zone mechanistic models represent a whole-building as a single, well-mixed
9	compartment. These AER models, such as the Lawrence Berkeley Laboratory (LBL)
10	model, can predict residential AER using input data from whole-building pressurization
11	tests (Sherman and Grimsrud, 1980), or leakage area models (Breen et al., 2010; Sherman
12	and McWilliams, 2007). Recently, the LBL air infiltration model was linked with a
13	leakage area model using population-level census and residential survey data (Sherman
14	and McWilliams, 2007) and individual-level questionnaire data (Breen et al., 2010). The
15	LBL model, which predicts the AER from air infiltration (i.e., small uncontrollable
16	openings in the building envelope) was also extended to include airflow from natural
17	ventilation (LBLX), and evaluated using window opening data (Breen et al., 2010). The
18	AER predictions from the LBL and LBLX models were compared to daily AER
19	measurements on seven consecutive days during each season from detached homes in
20	central North Carolina (Breen et al., 2010). For the individual model-predicted and
21	measured AER, the median absolute difference was 43% (0.17 h^{-1}) and 40% (0.17 h^{-1}) for
22	the LBL and LBLX models, respectively. Given the uncertainty of the AER
23	measurements (accuracy of 20-25% for occupied homes), these results demonstrate the
24	feasibility of using these AER models for both air infiltration (e.g., uncontrollable
25	openings) and natural ventilation (e.g., window opening) to help reduce the AER
26	uncertainty in exposure models. The capability of AER models could help support the
27	exposure modeling needs, as described in Section $4.5.3$, which includes the ability to
28	predict indoor concentrations of ambient O ₃ that may be substantial for conditions of high
29	AER such as open windows.

4.5.3 Microenvironment-Based Models

Population-based methods, such as the Air Pollution Exposure (APEX) and Stochastic
Human Exposure and Dose Simulation (SHEDS) integrated microenvironmental
exposure and dose models, involve stochastic treatment of the model inputs (U.S. EPA,
2009b; Burke et al., 2001). These are described in detail in the 2008 NO_X ISA (U.S. EPA,
2008b), in AX3.6.1. Stochastic models utilize distributions of pollutant-related and
individual-level variables, such as ambient and local O₃ concentration contributions and

1 breathing rate respectively, to compute the distribution of individual exposures across the 2 modeled population. The models also have the capability to estimate received dose 3 through a dosimetry model. Using distributions of input parameters in the model 4 framework rather than point estimates allows the models to incorporate uncertainty and 5 variability explicitly into exposure estimates (Zidek et al., 2007). These models estimate 6 time-weighted exposure for modeled individuals by summing exposure in each 7 microenvironment visited during the exposure period. 8 The initial set of input data for population exposure models is ambient air quality data, 9 which may come from a monitoring network or model estimates. Estimates of 10 concentrations in a set of microenvironments are generated either by mass balance methods, which can incorporate AER models (Section 4.5.3), or microenvironmental 11 12 factors. Microenvironments modeled include indoor residences; other indoor locations, 13 such as schools, offices, and public buildings; and vehicles. The sequence of 14 microenvironments and exertion levels during the exposure period is determined from 15 characteristics of each modeled individual. The APEX model does this by generating a 16 profile for each simulated individual by sampling from distributions of demographic 17 variables such as age, gender, and employment; physiological variables such as height 18 and weight; and situational variables such as living in a house with a gas stove or air 19 conditioning. Activity and location (microenvironmental) patterns from a database such 20 as CHAD are assigned to the simulated individual in a longitudinal manner, using age, 21 gender, and biometric characteristics (U.S. EPA, 2009a; Glen et al., 2008). Breathing 22 rates for each individual are calculated for each activity based on predicted energy 23 expenditures, and the corresponding dose may then be computed. APEX has an algorithm 24 to estimate O₃ dose and changes in FEV₁ resulting from O₃ exposure. Summaries of 25 individual- and population-level metrics are produced, such as maximum exposure or 26 dose, number of individuals exceeding a specified exposure/dose, and number of 27 person-days at or above benchmark exposure levels. The models also consider the 28 nonambient contribution to total exposure. Nonambient source terms are added to the 29 infiltration of ambient pollutants to calculate the total concentration in the 30 microenvironment. Output from model runs with and without nonambient sources can be 31 compared to estimate the ambient contribution to total exposure and dose. 32 Georgopoulos et al. (2005) used a version of the SHEDS model as the exposure 33 component of a modeling framework known as MENTOR (Modeling Environment for 34 Total Risk Studies) in a simulation of O_3 exposure in Philadelphia over a 2-week period 35 in July 1999. 500 individuals were sampled from CHAD in each of 482 census tracts to 36 match local demographic characteristics from U.S. Census data. Outdoor concentrations 37 over the modeling domain were calculated from interpolation of photochemical modeling 38 results and fixed-site monitor concentrations. These concentrations were then used as

1	input data for SHEDS. Median microenvironmental concentrations predicted by SHEDS
2	for nine simulated microenvironments were strongly correlated with outdoor
3	concentrations, a result consistent with the lack of indoor O ₃ sources in the model. A
4	regression of median microenvironmental concentrations against outdoor concentrations
5	indicated that the microenvironmental concentrations were appreciably lower than
6	outdoor concentrations (regression slope = 0.26). 95th percentile microenvironmental
7	concentrations were also well correlated with outdoor concentrations and showed a
8	regression slope of 1.02, although some microenvironmental concentrations were well
9	below the outdoor values. This suggests that in most cases the high-end concentrations
10	were associated with outdoor microenvironments. Although the authors did not report
11	exposure statistics for the population, their dose calculations also indicated that O ₃ dose
12	due to time spent outdoors dominated the upper percentiles of the population dose
13	distribution. They found that both the 50th and 95th percentile O_3 concentrations were
14	correlated with census-tract level outdoor concentrations estimated by photochemical
15	modeling combined with spatiotemporal interpolation, and attributed this correlation to
16	the lack of indoor sources of O ₃ . Relationships between exposure and concentrations at
17	fixed-site monitors were not reported.
18	An analysis has been conducted for the APEX model to evaluate the contribution of
	-

19 uncertainty in input parameters and databases to the uncertainty in model outputs 20 (Langstaff, 2007). The Monte Carlo analysis indicates that the uncertainty in model 21 exposure estimates for asthmatic children during moderate exercise is small to moderate, 22 with 95% confidence intervals of at most ± 6 percentage points at exposures above 60, 23 70, and 80 ppb (8-h avg) However, APEX appears to substantially underestimate the 24 frequency of multiple high-exposure events for a single individual. The two main sources 25 of uncertainty identified were related to the activity pattern database and the spatial 26 interpolation of fixed-site monitor concentrations to other locations. Additional areas 27 identified in the uncertainty analysis for potential improvement include: further 28 information on children's activities, including longitudinal patterns in the activity pattern 29 database; improved information on spatial variation of O₃ concentrations, including in 30 near-roadway and indoor microenvironments; and data from personal exposure monitors 31 with shorter averaging times to capture peak exposures and lower detection limits to 32 capture low indoor concentrations. A similar modeling approach has been developed for 33 panel epidemiologic studies or for controlled human exposure studies, in which activity 34 pattern data specific to the individuals in the study can be collected. Time-activity data is 35 combined with questionnaire data on housing characteristics, presence of indoor or 36 personal sources, and other information to develop a personalized set of model input 37 parameters for each individual. This model, the Exposure Model for Individuals, has been 38 developed by EPA's National Exposure Research Laboratory (U.S. EPA, 2011c; 39 Zartarian and Schultz, 2010).

4.6 Implications for Epidemiologic Studies

1	Exposure measurement error, which refers to the uncertainty associated with using
2	exposure metrics to represent the actual exposure of an individual or population, can be
3	an important contributor to variability in epidemiologic study results. Time-series studies
4	assess the daily health status of a population of thousands or millions of people over the
5	course of multiple years (i.e., thousands of days) across an urban area by estimating their
6	daily exposure using a short monitoring interval (hours to days). In these studies, the
7	community-averaged concentration of an air pollutant measured at central-site monitors
8	is typically used as a surrogate for individual or population ambient exposure. In
9	addition, panel studies, which consist of a relatively small sample (typically tens) of
10	study participants followed over a period of days to months, have been used to examine
11	the health effects associated with short-term exposure to ambient concentrations of air
12	pollutants (Delfino et al., 1996). Panel studies may also apply a microenvironmental
13	model to represent exposure to an air pollutant. A longitudinal cohort epidemiologic
14	study, such as the ACS cohort study, typically involves hundreds or thousands of subjects
15	followed over several years or decades (Jerrett et al., 2009). Concentrations are generally
16	aggregated over time and by community to estimate exposures.
17	Exposure error can under- or over-estimate epidemiologic associations between ambient
17 18	Exposure error can under- or over-estimate epidemiologic associations between ambient pollutant concentrations and health outcomes by biasing effect estimates toward or away
18	pollutant concentrations and health outcomes by biasing effect estimates toward or away
18 19	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (<u>Sheppard</u>
18 19 20	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (<u>Sheppard</u> et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the
18 19 20 21	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (<u>Sheppard</u> et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study
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 18 19 20 21 22 23 24 	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (Sheppard et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study of nondifferential exposure misclassification (Brauer et al., 2002). The importance of exposure misclassification varies with study design and is dependent on the spatial and temporal aspects of the design. For example, the use of a community-averaged O_3
 18 19 20 21 22 23 24 25 	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (Sheppard et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study of nondifferential exposure misclassification (Brauer et al., 2002). The importance of exposure misclassification varies with study design and is dependent on the spatial and temporal aspects of the design. For example, the use of a community-averaged O ₃ concentration in a time-series epidemiologic study may be adequate to represent the day-
 18 19 20 21 22 23 24 25 26 	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (Sheppard et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study of nondifferential exposure misclassification (Brauer et al., 2002). The importance of exposure misclassification varies with study design and is dependent on the spatial and temporal aspects of the design. For example, the use of a community-averaged O_3 concentration in a time-series epidemiologic study may be adequate to represent the day- to-day temporal concentration variability used to evaluate health effects, but may not
 18 19 20 21 22 23 24 25 26 27 	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (Sheppard et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study of nondifferential exposure misclassification (Brauer et al., 2002). The importance of exposure misclassification varies with study design and is dependent on the spatial and temporal aspects of the design. For example, the use of a community-averaged O ₃ concentration in a time-series epidemiologic study may be adequate to represent the day- to-day temporal concentration variability used to evaluate health effects, but may not capture differences in the magnitude of exposure due to spatial variability. Other factors
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 18 19 20 21 22 23 24 25 26 27 28 29 30 	pollutant concentrations and health outcomes by biasing effect estimates toward or away from the null, and tends to widen confidence intervals around those estimates (Sheppard et al., 2005; Zeger et al., 2000). Exposure misclassification can also tend to obscure the presence of potential thresholds for health effects, as demonstrated by a simulation study of nondifferential exposure misclassification (Brauer et al., 2002). The importance of exposure misclassification varies with study design and is dependent on the spatial and temporal aspects of the design. For example, the use of a community-averaged O_3 concentration in a time-series epidemiologic study may be adequate to represent the day- to-day temporal concentration variability used to evaluate health effects, but may not capture differences in the magnitude of exposure due to spatial variability. Other factors that could influence exposure estimates include nonambient exposure, topography of the natural and built environment, meteorology, measurement errors, use of ambient O_3 concentration as a surrogate for ambient O_3 exposure, and the presence of O_3 in a mixture

4.6.1 Non-Ambient Ozone Exposure

1	For other criteria pollutants, nonambient sources can be an important contributor to total
2	personal exposure. There are relatively few indoor sources of O_3 ; as a result, personal O_3
3	exposure is expected to be dominated by ambient O ₃ in outdoor microenvironments and
4	in indoor microenvironments with high air exchange rates (e.g., with open windows).
5	Even in microenvironments where nonambient exposure is substantial, such as in a room
6	with an O ₃ generator, this nonambient exposure is unlikely to be temporally correlated
7	with ambient O_3 exposure (Wilson and Suh, 1997), and therefore would not affect
8	epidemiologic associations between O_3 and a health effect (<u>Sheppard et al., 2005</u>). In
9	simulations of a nonreactive pollutant, <u>Sheppard et al. (2005</u>) concluded that nonambient
10	exposure does not influence the health outcome effect estimate if ambient and
11	nonambient concentrations are independent. Since personal exposure to ambient O_3 is
12	some fraction of the ambient concentration, it should be noted that effect estimates
13	calculated based on personal exposure rather than ambient concentration will be
14	increased in proportion to the ratio of ambient concentration to ambient exposure, and
15	daily fluctuations in this ratio can widen the confidence intervals in the ambient
16	concentration effect estimate, but uncorrelated nonambient exposure will not bias the
17	effect estimate (Sheppard et al., 2005; Wilson and Suh, 1997).

4.6.2 Spatial and Temporal Variability

18 Spatial and temporal variability in O₃ concentrations can contribute to exposure error in 19 epidemiologic studies, whether they rely on central-site monitor data or concentration 20 modeling for exposure assessment. Spatial variability in the magnitude of concentrations 21 may affect cross-sectional and large-scale cohort studies by undermining the assumption 22 that intra-urban concentration and exposure differences are less important than inter-23 urban differences. This issue may be less important for time-series studies, which rely on 24 day-to-day temporal variability in concentrations to evaluate health effects. Low inter-25 monitor correlations contribute to exposure error in time-series studies, including bias 26 toward the null and increased confidence intervals.

4.6.2.1 Spatial Variability

27	Spatial variability of O ₃ concentrations is highly dependent on spatial scale; in effect, O ₃
28	is a regional pollutant subject to varying degrees of local variability. In the immediate
29	vicinity of roadways, O ₃ concentrations are reduced due to reaction with NO and other
30	species (Section $4.3.4.2$); over spatial scales of a few kilometers, O ₃ may be more

- 1 homogeneous due to its formation as a secondary pollutant; over scales of tens of 2 kilometers, atmospheric processing can result in higher concentrations downwind of an 3 urban area than in the urban core. Local-scale variations have a large impact on the 4 relative magnitude of concentrations among urban monitors, while conditions favoring 5 high or low rates of O_3 formation (e.g., temperature) vary over large spatial scales. This 6 suggests that neighborhood monitors are likely to track one another temporally, but miss 7 small-scale spatial variability in magnitude. This is supported by an analysis in Atlanta 8 that found correlations greater than 0.8 for daily O₃ concentration metrics (1-h max, 9 8-h max, and 24-h avg) measured at monitors 10-60 km apart (Darrow et al., 2011a). In 10 rural areas, a lower degree of fluctuation in O_3 precursors such as NO and VOCs is likely 11 to make the diel concentration profile less variable than in urban areas, resulting in more 12 sustained ambient levels. Spatial variability contributes to exposure error if the ambient 13 O₃ concentration measured at the central site monitor is used as an ambient exposure 14 surrogate and differs from the actual ambient O₃ concentration outside a subject's 15 residence and/or worksite (in the absence of indoor O_3 sources). Averaging data from a 16 large number of samplers will dampen intersampler variability, and use of multiple 17 monitors over smaller land areas may allow for more variability to be incorporated into 18 an epidemiologic analysis.
- 19 Community exposure may not be well represented when monitors cover large areas with 20 several subcommunities having different sources and topographies, such as the 21 Los Angeles CSA (Section 3.6.2.1 and Section 4.4.3). Ozone monitors in Los Angeles 22 had a much wider range of intermonitor correlations (-0.06 to 0.97) than Atlanta (0.61 to 0.97)23 0.96) or Boston (0.56 to 0.97) using 2007-2009 data. Although the negative and near-zero 24 correlations in Los Angeles were observed for monitors located some distance apart 25 (>150 km), some closer monitor pairs had low positive correlations, likely due to changes 26 in land use, topography, and airflow patterns over short distances. Lower COD values, 27 which indicate less variability among monitors in the magnitude of O_3 concentrations, 28 were observed in Atlanta (0.05-0.13) and Boston (0.05-0.19) than Los Angeles 29 (0.05-0.56), although a single monitor (AM) was responsible for all Los Angeles COD 30 values above 0.40. The spatial and temporal variability in O₃ concentration in 24 MSAs 31 across the U.S. was also examined in the 2006 O₃ AQCD by using Pearson correlation 32 coefficients, values of the 90th percentile of the absolute difference in O_3 concentrations, 33 and CODs. No clear discernible regional differences across the U.S. were found in the 34 ranges of parameters analyzed.
- An analysis of the impact of exposure error due to spatial variability and instrument imprecision on time-series epidemiologic study results indicated that O₃ has relatively low exposure error compared to other routinely monitored pollutants, and that the simulated impact on effect estimates is minor. Goldman et al. (2011) computed

1population-weighted scaled semivariances and Pearson correlation coefficients for daily2concentration metrics of twelve pollutants measured at multiple central-site monitors in3Atlanta. 8-h daily max O3 exhibited the lowest semivariance and highest correlation of4any of the pollutants. Although this indicates some degree of urban-scale homogeneity5for O3, the analysis did not account for near-road effects on O3 concentrations.

6 Studies evaluating the influence of monitor selection on epidemiologic study results have 7 found that O₃ effect estimates are similar across different spatial averaging scales and 8 monitoring sites. A study in Italy compared approaches for using fixed-site monitoring 9 data in a case-crossover epidemiologic study of daily O₃ and mortality (Zauli Sajani et 10 al., 2011). O_3 effect estimates were found to be similar whether the nearest monitor was used, or whether single-city, three-city, or six-city regional averages were used for 11 12 exposure assessment. In contrast, effect estimates for PM₁₀ and NO₂ increased with 13 increasing scale of spatial averaging. Confidence intervals increased with increasing 14 spatial scale for all pollutants. The authors attributed the consistency of O_3 effect 15 estimates to the relative spatial homogeneity of O_3 over multi-km spatial scales, and 16 pointed to the high (0.85-0.95) inter-monitor correlations to support this. The use of 17 background monitors rather than monitors influenced by local sources in this study 18 suggests that local-scale spatial variation in O₃, such as that due to titration by traffic 19 emissions, was not captured in the analyses. A multi-city U.S. study of asthmatic children 20 found comparable respiratory effect estimates when restricting the analysis to the 21 monitors closest to the child's zip code centroid as when using the average of all 22 monitors in the urban area (Mortimer et al., 2002), suggesting little impact of monitor 23 selection. Sarnat et al. (2010) studied the spatial variability of O_3 , along with $PM_{2.5}$, NO_2 , 24 and CO, in the Atlanta, GA, metropolitan area and evaluated how spatial variability 25 affects interpretation of epidemiologic results, using time-series data for circulatory 26 disease ED visits. The authors found that associations with ambient 8-h daily maximum 27 O₃ concentration were similar among all sites tested, including multiple urban sites and a 28 rural site some 38 miles from the city center. This result was also observed for 24-h PM_{2.5} 29 concentrations. In contrast, hourly CO and NO₂ showed different associations for the 30 rural site than the urban sites, although the urban site associations were similar to one 31 another for CO. This suggests that the choice of monitor may have little impact on the 32 results of O₃ time-series studies, consistent with the moderate to high inter-monitor 33 correlations observed in Atlanta (Chapter 3).

34One potential explanation for this finding from the study by Sarnat et al. (2010) is that35although spatial variability at different scales contributes to a complicated pattern of36variations in the magnitude of O3 concentrations between near-road, urban core, and37urban downwind sites, day-to-day fluctuations in concentrations may be reflected across38multiple urban microenvironments. In addition, time-averaging of O3 and PM2.5

1	concentrations may smooth out some of the intra-day spatial variability observed with the
2	hourly CO and NO ₂ concentrations. However, some uncertainty in observed effect
3	estimates due to spatial variability and associated exposure error is expected to remain,
4	including a potential bias towards the null.

4.6.2.2 Seasonality

- 5 The relationship between personal exposure and ambient concentration has been found to 6 vary by season, with at least three factors potentially contributing to this variation: 7 differences in building ventilation (e.g., air conditioning or heater use versus open 8 window ventilation), higher O_3 concentrations during the O_3 season contributing to 9 increased exposure and improved detection by personal monitors; and changes in activity 10 pattern resulting in more time spent outside. Evidence has been presented in studies 11 conducted in several cities regarding the effect of ventilation on personal-ambient and 12 indoor-outdoor O_3 relationships (see Section 4.3.2 and Section 4.3.3). More limited 13 evidence is available regarding the specific effects of O_3 detection limits and activity 14 pattern changes on O₃ relationships.
- 15 Several studies have found increased summertime correlations or ratios between personal 16 exposure and ambient concentration (Sarnat et al., 2005; Sarnat et al., 2000) or between 17 indoor and outdoor O₃ concentrations (Geyh et al., 2000; Avol et al., 1998a). However, 18 others have found higher ratios in fall than in summer (Sarnat et al., 2006a) or equivalent, 19 near-zero ratios in winter and summer (Sarnat et al., 2001), possibly because summertime 20 use of air conditioners decreases building air exchange rates. It should be noted that O_3 21 concentrations during winter are generally much lower than summertime concentrations, 22 possibly obscuring wintertime relationships due to detection limit issues. Studies 23 specifically evaluating the effect of ventilation conditions on O_3 relationships have found 24 increased correlations or ratios for individuals or buildings experiencing higher air 25 exchange rates (Sarnat et al., 2006a; Geyh et al., 2000; Sarnat et al., 2000; Romieu et al., 26 1998a).
- 27Increased correlations or ratios between personal exposure and ambient concentration, or28between indoor and outdoor concentration, are likely to reduce error in exposure29estimates used in epidemiologic studies. This suggests that studies conducted during the30O3 season or in periods when communities are likely to have high air exchange rates31(e.g., during mild weather) may be less prone to exposure error than studies conducted32only during winter. Year-round studies that include both the O3 and non-O3 seasons may33have an intermediate level of exposure error.

4.6.3 Exposure Duration

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Epidemiologic studies of health effects associated with short-term and long-term exposures use different air pollution metrics and thus have different sources of exposure error. The following subsections discuss the impact of using different short-term and long-term exposure metrics on epidemiologic results.

4.6.3.1 Short-Term Exposure

5 The averaging time of the daily exposure metrics used to evaluate daily aggregated health 6 data (e.g., 1-h or 8-h daily maximum vs. 24-h avg concentration) may also impact 7 epidemiologic results, since different studies report different daily metrics. Correlations 8 between 1-h daily max, 8-h daily max, and 24-h avg concentrations for U.S. monitoring 9 sites are presented in Section 3.6.1 (Figure 3-23 and accompanying text). The two daily 10 peak values (1-h max and 8-h max) are well correlated, with a median (IQR) correlation 11 of 0.97 (0.96-0.98). The correlation between the 8-h max and 24-h avg are somewhat less 12 well correlated with a median (IQR) correlation of 0.89 (0.86-0.92). While this may 13 complicate quantitative comparisons between epidemiologic studies using different daily 14 metrics, as well as the interpretation of studies using metrics other than the current 8-h 15 standard, the high inter-metric correlations suggest it is a relatively small source of 16 uncertainty in comparing the results of studies using different metrics. This is supported 17 by a study comparing each of these metrics in a time-series study of respiratory ED visits 18 (Darrow et al., 2011a), which found positive associations for all metrics, with the 19 strongest association for the 8-h daily max exposure metric (Section 6.2.7.3). 20 The ratios of 1-h daily max, 8-h daily max, and 24-h avg concentrations to one another 21 have been found to differ across communities and across time within individual 22 communities (Anderson and Bell, 2010). For example, 8:24 hour ratios ranged from 23 1.23-1.83, with a median of 1.53. Lower ratios were generally observed in the spring and 24 summer compared to fall and winter. O₃ concentration was identified as the most 25 important predictor of ozone metric ratios, with higher overall O_3 concentrations 26 associated with lower ratios. In communities with higher long-term ozone concentrations, 27 the lower 8:24 hour ratio is attributed to high baseline O_3 , which results in elevated 24-h 28 average values. Differences in the representativeness of O_3 metrics introduces uncertainty 29 into the interpretation of epidemiologic results and complicates comparison of studies 30 using different metrics. Preferably, studies will report results using multiple metrics. In 31 cases where this does not occur, the results of the study by Anderson and Bell (2010) can 32 inform the uncertainty associated with using a standard increment to adjust effect 33 estimates based on different metrics so that they are comparable (Chapter 6).

1 A study compared measures of spatial and temporal variability for 1-h daily max and 2 24-h daily avg O₃ concentrations in Brazil (Bravo and Bell, 2011). The 1-h daily max 3 value was found to have higher correlation between monitors (i.e., lower temporal 4 variability) and lower COD (a measure of spatiotemporal variability which incorporates 5 differences in concentration magnitude, with lower values indicating lower variability; 6 see Chapter 3) than the 24-h avg value. The range of correlation coefficients and COD 7 values was similar between the two metrics, although the variation was lower for the 1-h 8 daily max, as indicated by the R^2 value for the regression of correlation coefficient on 9 inter-monitor distance.

4.6.3.2 Long-Term Exposure

10 Long-term O_3 exposure studies are not available that permit evaluation of the relationship 11 between long-term O₃ concentrations and personal or population exposure. The value of 12 short-term exposure data for evaluating long-term concentration-exposure relationships is 13 uncertain. If the longer averaging time (annual vs. daily or hourly) smooths out short-14 term fluctuations, long-term concentrations may be well-correlated with long-term 15 exposures. However, lower correlation between long-term exposures and ambient 16 concentration could occur if important exposure determinants change over a period of 17 several years, including activity pattern and residential air exchange rate.

- 18 A study in Canada suggests that an exposure metric based on a single year can represent 19 exposure over a multi-decade period. The authors compared exposure assessment 20 methods for long-term O_3 exposure and found that the annual average concentration in 21 the census tract of a subject's residence during 1980 and 1994 was well-correlated (0.76 22 and 0.82, respectively) with a concentration metric accounting for movement among 23 census subdivisions during 1980-2002 (Guay et al., 2011). This may have been due in 24 part to a relatively low rate of movement, with subjects residing on average for 71% of 25 the 22-year period in the same census subdivision they were in during 1980.
- 26 Analysis of the exposure assessment methodology in a recent study of mortality 27 associated with long-term O_3 exposure (Jerrett et al., 2009) is illustrative. In this study, 28 the authors computed quarterly averages of the daily 1-h max O₃ concentration, averaged 29 the two summer quarters together to produce an annual value, then calculated a 23-year 30 average value for each city in the study. Producing a single value for each city enables a 31 comparison of relatively cleaner cities with relatively more polluted cities. In this case, 32 the average was calculated using the 1-h daily max value; if the 24-h avg value had been 33 used, concentrations would have been lower and potentially more variable, based on 34 analyses in Chapter 3. According to

1	T.11. 2.7. (h. 2007.2000. 2
1	Table 3-7, the 2007-2009, 3-year average 1-h daily max value during the warm season
2	was approximately 50% higher than the corresponding 24-avg value on a nationwide
3	basis. Correlation between the two metrics varies by site, indicating the differential
4	influence of the overnight period on 24-h avg concentrations. The median correlation
5	between 1-h daily max and 24-h avg is 0.83, with an IQR of 0.78-0.88. It is not clear,
6	however, that a different exposure assignment method would yield different results.
7	Long-term O_3 trends, as discussed in Chapter <u>3</u> , show gradually decreasing
8	concentrations. Figure 3-48 shows that concentrations have decreased most for the 90th
9	percentile, with relatively little change among the 10th percentile monitors. The decrease
10	has been greater in the eastern U.S. than in the western part of the country (excluding
11	California). For the most part, the rank order of regions in terms of O_3 concentration has
12	remained the same, as shown in Figure 3-50, with the Northeast, Southeast, and
13	California exhibiting the highest concentrations. The decreasing trend is consistent across
14	nearly all monitors in the U.S., with only 1-2% of monitors reporting an increase of more
15	than 5 ppb between the 2001-2003 and 2008-2010 time periods (Figure 3-52 and
16	Figure 3-53). This provides some evidence that epidemiologic studies of long-term
17	exposure are not affected by drastic changes in O_3 concentration, such as a relatively
18	clean city becoming highly polluted or the reverse.
19	A few epidemiologic studies have evaluated the impact of distance to monitor on
20	associations between long-term O ₃ concentration and reproductive outcomes, as
21	discussed in Chapter $\frac{7}{2}$. It is not clear from this evidence whether using a local monitor
22	for these multi-month concentration averages improves exposure assessment. Jalaludin et
23	al. (2007) found somewhat higher effect estimates for women living within 5 km of a
24	fixed-site O ₃ monitor than for all women in the Sydney metropolitan area, suggesting that
25	increased monitor proximity reduced exposure misclassification. In contrast, <u>Darrow et</u>
26	al. (2011b) found no substantial difference between effect estimates for those living
27	within 4 mi of a fixed-site monitor and those living in the five-county area around
28	Atlanta. This result could be due to spatial variability over smaller scales than the 4-mi
29	radius evaluated, time spent away from the residence impacting O_3 exposure, or
30	similarity in monitor location and representativeness across the urban area (see
31	Figure 4-4). At this time, the effect of exposure error on long-term exposure
32	
61	epidemiologic studies is unclear.

4.6.4 Exposure to Copollutants and Ozone Reaction Products

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34			

Although indoor O_3 concentrations are usually well below ambient concentrations, the same reactions that reduce O_3 indoors form particulate and gaseous species, including

1 other oxidants, as summarized in Section 4.3.4.3. Exposures to these reaction products 2 would therefore be expected to be correlated with ambient O_3 concentrations, although no 3 evidence was identified regarding personal exposures. Such exposure could potentially 4 contribute to health effects observed in epidemiologic studies.

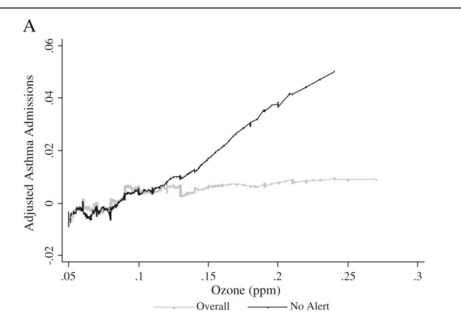
4.6.5 **Averting Behavior**

7

5 As described in Section 4.4.2, several recent studies indicate that some populations alter 6 their behavior on high ozone days to avoid exposure. Such behavioral responses to information about forecasted air quality may introduce systematic measurement error in 8 air pollution exposure, leading to biased estimates of the impact of air pollution on health. 9 For example, studies have hypothesized that variation in time spent outdoors may be a 10 driving factor behind the considerable heterogeneity in ozone mortality impacts across 11 communities (Bell et al., 2004). If averting behavior reduces outdoor O_3 exposure, then 12 studies that do not account for averting behavior may produce effect estimates that are 13 biased towards the null (Section 6.2.7.2).

14 This is supported by an epidemiologic study that examined the association between 15 exposure to ambient ozone concentrations and asthma hospitalizations in Southern 16 California during 1989-1997, which indicates that controlling for avoidance behavior 17 increases the effect estimate for both children and older adults, but not for adults aged 18 20-64 (Neidell and Kinney, 2010; Neidell, 2009). Figure 4-7 and Figure 4-8, reproduced 19 from Neidell (2009), show covariate-adjusted asthma hospital admissions as a function of 20 daily maximum 1-h O_3 concentration for all days (gray line) and days when no O_3 alert 21 was issued (black line). Stage 1 smog alerts were issued by the State of California for 22 days when ambient O_3 concentrations were forecast to be above 0.20 ppm; however, the 23 concentration-response functions are based on measured O₃ concentrations. For children 24 aged 5-19 (Figure 4-7), hospital admissions were higher on high- O_3 days when no alert 25 was issued, especially on days with O_3 concentrations above 0.15 ppm (150 ppb). The 26 concentration-response curves for all days and days with no alert diverge at measured O_3 27 concentrations between 0.10 and 0.15 ppm because smog alerts begin to be issued more 28 frequently in this range. This suggests that in the absence of information that would 29 enable averting behavior, children experience higher ozone exposure and subsequently a 30 greater number of asthma hospital admissions than on alert days with similar O_3 31 concentrations. The lower rate of admissions observed when alert days were included in 32 the analysis suggests that averting behavior reduced O₃ exposure and asthma hospital 33 admissions. In both cases, O_3 was found to be associated with asthma hospital 34 admissions, although the strength of the association is underestimated when not 35 accounting for averting behavior. A different result was observed when examining

associations for adults aged 20-64 (Figure 4-8), who had similar rates of hospital admissions on non-alert days as on all days. The lack of change for adults aged 20-64, which is primary employment age, may reflect lower response to air quality alerts due to the increased opportunity cost of behavior change. The finding that air quality information reduces the daily asthma hospitalization rate in these populations provides additional support for a link between ozone and health effects.



Source: Reprinted with permission of the Board of Regents of the University of Wisconsin System, University of Wisconsin Press (Neidell, 2009).

Figure 4-7 Adjusted asthma hospital admissions by age on lagged ozone by alert status, ages 5-19.

1

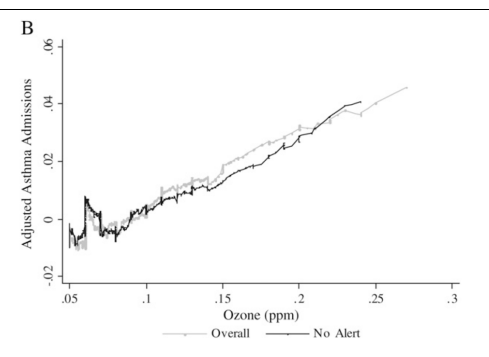
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Source: Reprinted with permission of the Board of Regents of the University of Wisconsin System, University of Wisconsin Press; <u>Neidell (2009</u>).

Figure 4-8 Adjusted asthma hospital admissions by age on lagged ozone by alert status, ages 20-64.

4.6.6 Exposure Estimation Methods in Epidemiologic Studies

1	Epidemiologic studies use a variety of methods to assign exposure. Study design, data
2	availability, and research objectives are all important factors for epidemiologists when
3	selecting an exposure assessment method. Common methods for assigning exposure
4	using monitoring data include using a single fixed-site monitor to represent population
5	exposure, averaging concentrations from multiple monitors, and selecting the closest
6	monitor. Investigators may also use statistical adjustment methods, such as trimming
7	extreme values, to prepare the concentration data set. Panel or small-scale cohort studies
8	involving dozens of individuals may use more individualized concentration
9	measurements, such as personal exposures, residential indoor or outdoor measurements,
10	or concentration data from local study-specific monitors. For long-term epidemiologic
11	studies, the lack of personal exposure data or dedicated measurements means that
12	investigators must rely on fixed-site monitoring data. These data may be used directly,
13	averaged across counties or other geographic areas, or used to construct geospatial or
14	regression models to assign concentrations to unmonitored locations. Longer-term
15	averages (months to years) are typically used (e.g., in studies discussed in
16	Section <u>7.3.1.1</u>). Chapters 6 and 7 describe the exposure assessment methods used in the

- epidemiologic studies described therein, providing additional detail on studies with
 innovative or expanded techniques designed to improve exposure assessment and reduce
 exposure error.
- 4 The use of O_3 measurements from central ambient monitoring sites is the most common 5 method for assigning exposure in epidemiologic studies. However, fixed-site 6 measurements do not account for the effects of spatial variation in O₃ concentration, 7 ambient and non-ambient concentration differences, and varying activity patterns on 8 personal exposures (Brown et al., 2009; Chang et al., 2000; Zeger et al., 2000). The use 9 of fixed-site concentrations results in minimal exposure error when: (1) O_3 concentrations 10 are uniform across the region; (2) personal activity patterns are similar across the 11 population; and (3) housing characteristics, such as air exchange rate and indoor reaction 12 rate, are constant over the study area. Since these factors vary by location and population, 13 there will be errors in the magnitude of total exposure based solely on ambient 14 monitoring data.
- 15 Modeled concentrations can also be used as exposure surrogates in epidemiologic studies, 16 as discussed in Section 4.5. Geostatistical spatial interpolation techniques, such as IDW 17 and kriging, can provide finer-scale estimates of local concentration over urban areas. A 18 microenvironmental modeling approach simulates exposure using empirical distributions 19 of concentrations in specific microenvironments together with human activity pattern 20 data. The main advantage of the modeling approach is that it can be used to estimate 21 exposures over a wide range of population and scenarios. A main disadvantage of the 22 modeling approach is that the results of modeling exposure assessment must be compared 23 to an independent set of measured exposure levels (Klepeis, 1999). In addition, 24 resource-intensive development of validated and representative model inputs is required, 25 such as human activity patterns, distributions of air exchange rate, and deposition rate. 26 Therefore, modeled exposures are used much less frequently in epidemiologic studies.

4.7 Summary and Conclusions

- This section will briefly summarize and synthesize the main points of the chapter, with particular attention to the relevance of the material for the interpretation of epidemiologic studies.
- 30Passive badge samplers are the most widely used technique for measuring personal O331exposure (Section 4.3.1). The detection limit of the badges for a 24-h sampling period is32approximately 5-10 ppb, with lower detection limits at longer sampling durations. In low-33concentration conditions this may result in an appreciable fraction of 24-h samples being34below the detection limit. The use of more sensitive portable active monitors, including

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- some that have recently become available, may help overcome this issue and improve
 personal monitoring in the future.
- Since there are relatively few indoor sources of O₃, indoor O₃ concentrations are often
 substantially lower than outdoor concentrations due to reactions of O₃ with indoor
 surfaces and airborne constituents (Section 4.3.2). Air exchange rate is a key determinant
- 6 of the I/O ratio, which is generally in the range of 0.1-0.4 (<u>Table 4-1</u>), with some 7 evidence for higher ratios during the O₃ season when concentrations are higher.
- 8 Personal exposure is moderately correlated with ambient O_3 concentration, as indicated 9 by studies reporting correlations generally in the range of 0.3-0.8 (Table 4-2). Hourly 10 concentration correlations are more variable than those averaged over 24 hours or longer, 11 with correlations in outdoor microenvironments (0.7-0.9) much higher than those in 12 residential indoor (0.1) or other indoor (0.3-0.4) microenvironments. Some studies report 13 substantially lower personal-ambient correlations, a result attributable in part to low air 14 exchange rate and O₃ concentrations below the sampler detection limit, conditions often 15 encountered during wintertime. Low correlations may also occur for individuals or 16 populations spending substantial time indoors.
- 17The ratio between personal exposure and ambient concentration varies widely depending18on activity patterns, housing characteristics, and season, with higher personal-ambient19ratios generally observed with increasing time spent outside, higher air exchange rate,20and in seasons other than winter (Table 4-3). Personal-ambient ratios are typically210.1-0.3, although individuals spending substantial time outdoors (e.g., outdoor workers)22may have much higher ratios (0.5-0.9).
- 23 Personal exposure to other pollutants shows variable association with personal exposure 24 to O_3 , with differences in copollutant relationships depending on factors such as season, 25 city-specific characteristics, activity pattern, and spatial variability of the copollutant 26 (Section 4.3.4). In near-road and on-road microenvironments, correlations between O_3 27 and traffic-related pollutants are moderately to strongly negative, with the most strongly 28 negative correlations observed for NO_2 (-0.8 to -0.9). This is consistent with the 29 chemistry of NO oxidation, in which O_3 is consumed to form NO_2 . The more moderate 30 negative correlations observed for PM_{2.5}, PM_{1.0}, and VOC may reflect reduced 31 concentrations of O₃ in polluted environments due to other scavenging reactions. A 32 similar process occurs indoors, where infiltrated O₃ reacts with airborne or surface-33 associated materials to form secondary compounds, such as formaldehyde. Although such 34 reactions decrease indoor O_3 exposure, they result in increasing exposure to other species 35 which may themselves have health effects.

- 1 Variations in ambient O_3 concentrations occur over multiple spatial and temporal scales. 2 Near roadways, O₃ concentrations are reduced due to reaction with NO and other species 3 (Section 4.3.4.2). Over spatial scales of a few kilometers and away from roads, O_3 may 4 be somewhat more homogeneous due to its formation as a secondary pollutant, while 5 over scales of tens of kilometers, additional atmospheric processing can result in higher 6 concentrations downwind of an urban area. Although local-scale variability impacts the 7 magnitude of O_3 concentrations, O_3 formation rates are influenced by factors that vary 8 over larger spatial scales, such as temperature (Section 3.2), suggesting that urban 9 monitors may track one another temporally but miss small-scale variability in magnitude. 10 The resulting uncertainty in exposure contributes to exposure measurement error in 11 epidemiologic studies.
- 12 Another factor that may influence epidemiologic results is the tendency for people to 13 avoid O_3 exposure by altering their behavior (e.g., reducing time spent outdoors) on high-14 O_3 days. Activity pattern has a substantial effect on ambient O_3 exposure, with time spent 15 outdoors contributing to increased exposure (Section 4.4.2). Averting behavior has been 16 predominantly observed among children, older adults, and people with respiratory 17 problems. Such effects are less pronounced in the general population, possibly due to the 18 opportunity cost of behavior modification. Evidence from one recent epidemiologic study 19 indicates increased asthma hospital admissions among children and older adults when O₃ 20 alert days were excluded from the analysis (presumably thereby eliminating averting 21 behavior based on high O₃ forecasts). The lower rate of admissions observed when alert 22 days were included in the analysis suggests that estimates of health effects based on 23 concentration-response functions which do not account for averting behavior may be 24 biased towards the null.
- 25 The range of personal-ambient correlations reported by most studies (0.3-0.8) is similar 26 to that for NO₂ (U.S. EPA, 2008b) and somewhat lower than that for PM_{2.5} (U.S. EPA, 27 2009d). To the extent that relative changes in central-site monitor concentration are 28 associated with relative changes in exposure concentration, this indicates that ambient 29 monitor concentrations are representative of day-to-day changes in average total personal 30 exposure and in personal exposure to ambient O_3 . The lack of indoor sources of O_3 , in 31 contrast to NO₂ and PM_{2.5}, is partly responsible for low indoor-outdoor ratios (generally 32 0.1-0.4) and low personal-ambient ratios (generally 0.1-0.3), although it contributes to 33 increased personal-ambient correlations. The lack of indoor sources also suggests that 34 fluctuations in ambient O_3 may be primarily responsible for changes in personal 35 exposure, even under low-ventilation, low-concentration conditions. Nevertheless, low 36 personal-ambient correlations are a source of exposure error for epidemiologic studies, 37 tending to obscure the presence of potential thresholds, bias effect estimates toward the 38 null, and widen confidence intervals, and this impact may be more pronounced among

1	populations spending substantial time indoors. The impact of this exposure error may
2	tend more toward widening confidence intervals than biasing effect estimates, since
3	epidemiologic studies evaluating the influence of monitor selection indicate that effect
4	estimates are similar across different spatial averaging scales and monitoring sites.

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5 DOSIMETRY AND MODE OF ACTION

5.1 Introduction

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This chapter has two main purposes. The first is to describe the principles that underlie the dosimetry of O_3 and to discuss factors that influence it. The second is to describe the modes of action leading to the health effects that will be presented in Chapters 6 and 7. This chapter is not intended to be a comprehensive overview, but rather, it updates the basic concepts derived from O_3 literature presented in previous documents (U.S. EPA, 2006b, 1996a) and introduces the recent relevant literature.

- 7In Section 5.2, particular attention is given to dosimetric factors influencing individual8risk of developing effects from O3 exposure. As there have been few O3 dosimetry studies9published since the last AQCD, the reader is referred to previous documents (U.S. EPA,102006b, 1996a) for more detailed discussion of the past literature. Evaluation of the11progress in the interpretation of past dosimetry studies, as well as studies published since122005, in the areas of uptake, reactions, and models for O3 dosimetry, is discussed.
- 13 Section 5.3 highlights findings of studies published since the 2006 O₃ AQCD, which 14 provide insight into the biological pathways by which O₃ exerts its actions. Since 15 common mechanisms lead to health effects from both short- and long-term exposure to 16 O_3 , these pathways are discussed in Chapter 5 rather than in later chapters. The related 17 sections of Chapters 6 and 7 are indicated. Earlier studies that represent the current state 18 of the science are also discussed. Studies conducted at more environmentally-relevant 19 concentrations of O_3 are of greater interest, since mechanisms responsible for effects at 20 low O_3 concentrations may not be identical to those occurring at high O_3 concentrations. 21 Some studies at higher concentrations are included if they were early demonstrations of 22 key mechanisms or if they are recent demonstrations of potentially important new 23 mechanisms. The topics of dosimetry and mode of action are bridged by reactions of O_3 24 with components of the extracellular lining fluid (ELF), which play a role in both O_3 25 uptake and biological responses (Figure 5-1).
- 26In addition, this chapter discusses interindividual variability in responses, and issues27related to species comparison of doses and responses (Section 5.4 and Section 5.5). These28topics are included in this chapter because they are influenced by both dosimetric and29mechanistic considerations.

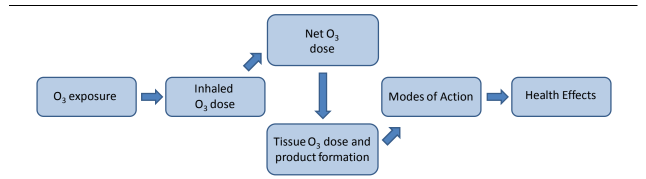


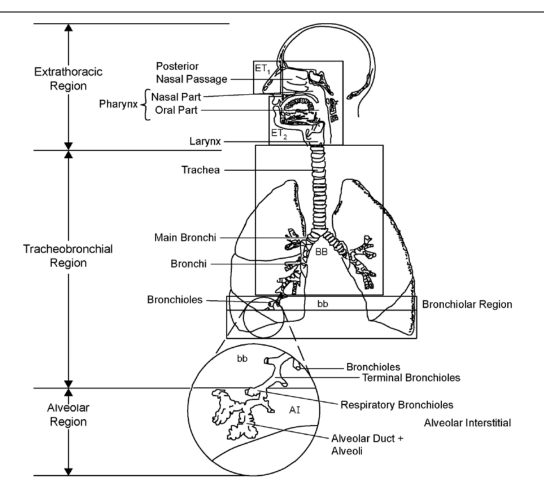
Figure 5-1 Schematic of the ozone exposure and response pathway. Ozone transport follows a path from exposure concentration, to inhaled dose, to net dose, to the local tissue dose. Chapter 5 discusses the concepts of dose and modes of action that result in the health effects discussed in Chapters 6 and 7.

5.2 Human and Animal Ozone Dosimetry

5.2.1 Introduction

1	Dosimetry refers to the measurement or estimation of the quantity of or rate at which a
2	chemical and/or its reaction products are absorbed and retained at target sites. Figure 5-1
3	illustrates the transport of O_3 or its reaction products from exposure to dose to the
4	development of health effects. Ozone exposure has been defined in Section 4.2 and
5	consists of contact between the human or animal and O_3 at a specific concentration for a
6	specified period of time (i.e., exposure = concentration \times time). The amount of O ₃ present
7	in a given volume of air for which animals and individuals are exposed is termed
8	exposure concentration. Ozone exposure will result in some amount (dose) of O_3 crossing
9	an exposure surface to enter a target area. The initial measure of dose after O_3 enters the
10	RT is inhaled dose and is the amount or rate of O_3 that crosses the outer RT surface
11	before crossing the ELF and is effectively C×t× \dot{V}_E . Ozone may then cross from the gas
12	phase across the ELF interface where net dose may be measured. Net dose is the amount
13	or rate of entry of O_3 across the gas/ELF interface. In modeling studies, the dose rate is
14	often expressed as a flux per unit of surface area of a region of respiratory epithelium.
15	Finally, O_3 or its reaction products may reach the tissues and tissue dose of O_3 can be
16	reported. Tissue dose is the amount of O_3 or its reaction products absorbed and available
17	for reacting with tissues and is difficult and rarely measured. In the literature, the
18	exposure concentration and various measures of dose (i.e., net dose and inhaled dose) are
19	often used as surrogates for tissue dose. However, ambient or exposure concentrations are

not a true measure of dose so understanding the relationship between ambient
 concentrations and tissue dose allows for a greater appreciation of the dose-response from
 O₃ exposure.

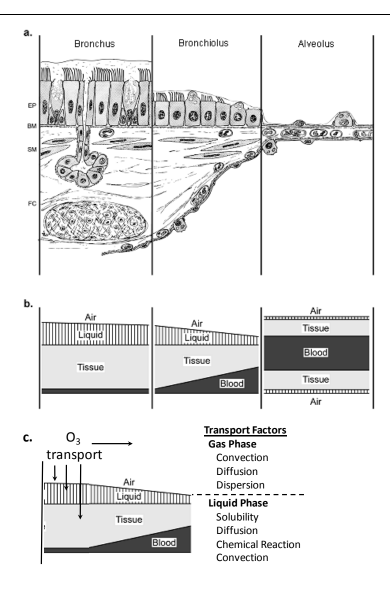


Note: Structures are anterior nasal passages, ET₁; oral airway and posterior nasal passages, ET₂; bronchial airways, BB; bronchioles, bb; and alveolar interstitial, AI. Source: Based on ICRP (1994).

Figure 5-2 Representation of respiratory tract regions in humans.

Ozone is a highly reactive, though poorly water soluble, gas at physiological temperature.
The latter feature is believed to be the reason why it is able to penetrate into targets in the
lower respiratory tract (LRT). Figure 5-2 presents the basic structure of the human
respiratory tract (RT). The lung can be divided into three major regions: the extrathoracic
(ET) region or upper respiratory tract (URT, from the nose/mouth to larynx); the
tracheobronchial (TB) tree (from trachea to the terminal bronchioles); and the alveolar or

- 1 pulmonary region (from the respiratory bronchioles to the terminal alveolar sacs). The 2 latter two regions comprise the LRT. Although the structure varies, the illustrated 3 anatomic regions are common to all mammalian species with the exception of the 4 respiratory bronchioles. Respiratory bronchioles, the transition region between ciliated 5 and fully alveolated airways, are found in humans, dogs, ferrets, cats, and monkeys. 6 Respiratory bronchioles are absent in rats and mice and abbreviated in hamsters, guinea 7 pigs, sheep, and pigs. The branching structure of the ciliated bronchi and bronchioles also 8 differs between species from being a rather symmetric and dichotomous branching 9 network of airways in humans to a more monopodial branching network in other 10 mammals.
- 11 Figure 5-3 illustrates the structure of the LRT with progression from the large airways in 12 the TB region to the alveolus in the alveolar region. The fact that O_3 is so chemically 13 reactive has suggested to some that its tissue dose at the target sites exists in the form of 14 oxidation products such as aldehydes and peroxides (see Section 5.2.3). Reaction 15 products are formed when O₃ interacts with components of the ELF such as lipids and 16 antioxidants. The ELF varies throughout the length of the RT with the bronchial tree 17 lined with a thin film of mucus and the alveolar region lined with a thinner layer of surfactant solution (Figure 5-3b). Ozone dose is directly related to the coupled diffusion 18 19 and chemical reactions occurring in ELF, a process termed "reactive absorption." Thus, 20 the O_3 dose depends on both the concentration of O_3 as well as the availability of 21 substrates within the ELF.
- 22Ozone dose is affected by complex interactions between a number of other major factors23including RT morphology, breathing route, frequency, and volume, physicochemical24properties of the gas, physical processes of gas transport, as well as the physical and25chemical properties of the ELF and tissue layers (Figure 5-3c). The role of these26processes varies throughout the length of the RT and as O3 moves from the gas to liquid27compartments of the RT.



Note: (a) Illustrates basic airway anatomy. Structures are epithelial cells, EP; basement membrane, BM; smooth muscle cells, SM; and fibrocartilaginous coat, FC. (b) Illustrates the relative amounts of liquid, tissue, and blood with distal progression. In the bronchi there is a thick surface lining over a relatively thick layer of tissues. With distal progress, the lining diminishes allowing increased access of compounds crossing the air-liquid interface to the tissues and the blood. (c) Presents the factors acting in the gas and liquid phases of O_3 transport.

Source: Panel (a) reprinted with permission of McGraw-Hill (Weibel, 1980).

Figure 5-3 Structure of lower airways with progression from the large airways to the alveolus.

1	Two types of measurements have been used to arrive at the O_3 dose to target sites during
2	breathing: (1) measurement of removal of O ₃ from the air stream (termed "uptake"); and
3	(2) measurement of chemical reactions in tissues or with biomolecules known to be
4	present in tissues (termed "reactants"). The results of the above measurements have been
5	incorporated into mathematical models for the purpose of explaining, predicting, and

1extrapolating O3 dose in different exposure scenarios. Few new studies have investigated2the uptake of O3 in the RT since the last O3 assessment (U.S. EPA, 2006b). The studies3that have been conducted generally agree with the results presented in the past and do not4change the dosimetry conclusions of the last document.

5.2.2 Ozone Uptake

5	Past AQCDs provide information on the majority of literature relevant to understanding
6	the state of the science in O3 dosimetry. Measurements of O3 dose have been inferred
7	from simultaneous measurements of airflow and O3 concentration at the airway opening
8	of the nose or mouth (Nodelman and Ultman, 1999; Wiester et al., 1996a) as well as at
9	internal sampling catheters (Gerrity et al., 1995; Gerrity et al., 1988). One method of
10	quantifying O_3 dose is to measure the amount of O_3 removed from the air stream during
11	breathing (termed "uptake"). The difference in the amount of O_3 inhaled and exhaled
12	relative to the amount of inhaled O ₃ is termed fractional absorption. Uptake efficiency is
13	also reported and refers to the fraction of O_3 absorbed in a region as a function of the total
14	amount of O_3 entering the given region. Uptake studies have utilized bolus and
15	continuous O ₃ breathing techniques as well as modeling to investigate these measures of
16	uptake and the distribution of O_3 uptake between the upper and lower respiratory tract. A
17	number of the studies that have measured the fractional absorption and uptake efficiency
18	of O_3 in the human RT, URT, and LRT are presented in <u>Table 5-1</u> . For bolus exposure
19	studies that reported fractional absorption, the total RT uptake efficiency was estimated
20	as the sum of the products of the experimental bolus absorption and incremental volume
21	of a bolus into a breath divided by the tidal volume of the breath, or where available, was
22	taken from Table 1A of <u>Schlesinger et al. (1997</u>).

Reference	Nose ^a	Inspiratory	V _T (mL)	f _B (bpm) [♭]		Uptake Efficiency		
		Flow (mL/sec)			URT, complete breath	URT, inspiration	LRT, complete breath	Total RT, tidal breath
Continuous Expos	ure							
Gerrity et al.	OR	509	832	18		0.40	0.91	
<u>(1988</u>)	N	456	754	18		0.36	0.91	
	OR/N	500	800	18		0.43	0.91	
	OR/N	350	832	12		0.41	0.93	
	OR/N	634	778	24		0.38	0.89	
Gerrity et al.	OR	1,360	1,650	25		0.37	0.43	0.81
<u>(1994)</u>	OR	1,360	1,239	35		0.41	0.36	0.78
Gerrity et al. (1995)	OR	330	825	12		0.27	0.95	0.91
Wiester et al.	OR	539	631	16				0.76
<u>(1996a</u>)	N	514	642	16				0.73
<u>Rigas et al. (2000</u>)	Face mask	480	1,100	27.6				0.86
Santiago et al.	Ν	50				0.80 ^c		
<u>(2001</u>)	N	250				0.33		
Bolus Exposure								
<u>Hu et al. (1992</u>)	Mouth- piece	250	500		0.46			0.88
Kabel et al. (1994)	Mouth- piece	250	500		0.50			0.88
	Mouth- piece	250	500		0.53			0.88
	N	250	500		0.78			0.94
<u>Hu et al. (1994</u>)	Mouth- piece	150	500		0.65			0.91
	Mouth- piece	250	500		0.51			0.87
	Mouth- piece	500	500		0.26			0.82
	Mouth- piece	750	500		0.16			0.78
	Mouth- piece	1,000	500		0.11			0.76
<u>Ultman et al.</u> (1994)	Mouth- piece	250	500 ^d	15	0.30			
	Mouth- piece	250	500	15	0.47			

Table 5-1 Human respiratory tract uptake efficiency data

Reference	Mouth/ Nose ^ª	Inspiratory Flow (mL/sec)	V _T (mL)	f _Β (bpm) ^b	Uptake Efficiency			
					URT, complete breath	URT, inspiration	LRT, complete breath	Total RT, tidal breath
Bush et al. (1996)	Mouth- piece	250	500		0.51			0.89
Nodelman and Ultman (1999)	Nasal Cannula	150	500	18	0.90			0.92
	Nasal Cannula	1,000	500	120	0.50			0.84
	Mouth- piece	150	500	18	0.77			0.91
	Mouth- piece	1,000	500	120	0.25			0.75
Ultman et al.	OR	490	450 ^d	32.7				0.87
<u>(2004</u>)	OR	517	574	27				0.91

^aOR = oral exposure during spontaneous breathing; N = nasal exposure during spontaneous breathing; OR/N = pooled data from oral and nasal exposure; mouthpiece = exposure by mouthpiece.

 ${}^{\rm b}f_{\rm B}$ is either measured or is computed from flows and V_r.

^c F_{URT} from <u>Santiago et al. (2001</u>) represents nasal absorption (F_{nose}).

 $^{d}V_{T}$ is computed from flow and f_B.

5.2.2.1 Gas Transport Principles

1	The three-dimensional transport of O_3 in the lumen of an airway is governed by diffusion
2	and bulk flow or convection. When modeled as a one-dimensional process in which the
3	radial profiles of axial velocity and O ₃ concentration profiles are flat, O ₃ transport along
4	an airway lumen occurs by convection, axial diffusion and a coupled diffusion-reaction
5	process called dispersion. Simultaneously, O_3 diffuses into the ELF where it undergoes
6	radial diffusion and chemical reaction (Figure 5-3c) (Miller, 1995). The relative
7	importance of these transport mechanisms varies among RT regions for a given level of
8	ventilation. In the URT and major bronchi, bulk airflow tends to be the predominant
9	mechanism for axial transport in the airway lumen. However, in the alveolar region of the
10	lung, diffusion is the major gas transport mechanism.
11	Gas transport in the TB region occurs by a combination of bulk flow and mixing
12	(Ultman, 1985). Mixing can occur by diffusion processes associated with the molecular
13	nature of the gas and by convection, which depends on local velocity patterns. The
14	complexity of the airway structure and surface affects the bulk airflow patterns so that not
15	all nasal and lung surfaces receive the same O ₃ exposure or dose (Miller and Kimbell,
16	<u>1995</u>). For example, it has been reported that the larger surface-to-volume ratio
17	associated with the smaller airways in women enhances local O ₃ uptake and reduces the
18	distal penetration volume of O_3 into the RT of women relative to men (Ultman et al.,
19	2004) Also it uses reported that sharpes in space spatianal area socilable for sea
	2004). Also, it was reported that changes in cross-sectional area available for gas

1	The principal influence on mining in the TD preside comes from the original selection and file
1	The principal influence on mixing in the TB region comes from the axial velocity profile
2	and diffusion. When air flows through an airway, O ₃ located near the tube center moves
3	faster than O ₃ near the tube wall where frictional forces retard the flow. This
4	non-uniformity in the radial profile of velocity gives rise to an axial spreading or
5	dispersion of the O ₃ that operates in parallel with bulk flow and axial diffusion (a process
6	caused by the ever-present Brownian motion of individual O_3 molecules). The shape of
7	the velocity profile is affected by the flow direction through bifurcating airway branches
8	(Schroter and Sudlow, 1969). The velocity profile is nearly parabolic during inhalation
9	but quite flat during exhalation. Thus, there tends to be greater axial dispersion during
10	inhalation than during exhalation. Dispersion also depends on the nature of the flow, that
11	is, whether it is laminar (i.e., streamlined) or turbulent (i.e., possessing random velocity
12	fluctuations). Because turbulent flow flattens velocity profiles, it may actually diminish
13	dispersion. In humans, turbulent flow persists only a few generations into the RT. The
14	persistence of turbulence into the RT also varies by species and flow rates. For example,
15	airflow is nonturbulent in the rat nose at any physiologic flow rate but may be highly
16	turbulent in the human nose during exercise (Miller, 1995). Diffusive forces and
17	resistance vary along the RT. Diffusive resistance increases with distal penetration into
18	the RT with a study reporting that the gas boundary layer contributes 53% of the overall
19	diffusive resistance in the URT, 78% in the proximal LRT, and 87% in the distal LRT
20	(<u>Hu et al., 1994</u>).

21Conversely, the principal mechanism of gas mixing in the lung periphery is molecular22diffusion (Engel, 1985). While moving into more distal areas of the RT, the23cross-sectional area of the airways rapidly increases and linear velocities decrease,24leading to a greater role for molecular diffusion of gases. Gas molecules close to the25alveolar-capillary membrane have almost zero convective velocity with respect to the26membrane, and this creates a substantial boundary layer resistance to O3 transfer across27the gas-eLF interface.

5.2.2.2 Target Sites for Ozone Dose

28	A primary uptake site of O_3 delivery to the lung epithelium is believed to be the
29	centriacinar region (CAR). The CAR refers to the zone at the junction of the TB airways
30	and the gas exchange region. This area is also termed the proximal alveolar region (PAR)
31	and is defined as the first generation distal to the terminal bronchioles. Contained within
32	the CAR, the respiratory bronchioles were confirmed as the site receiving the greatest O_3
33	dose (¹⁸ O mass/lung weight) in resting O ₃ exposed rhesus monkeys, when not considering
34	the nose (Plopper et al., 1998). Furthermore, the greatest cellular injury occurred in the
35	vicinity of the respiratory bronchioles and was dependent on the delivered O3 dose to

1	these tissues (see also Section $5.4.1$). However, ¹⁸ O label was detected to a lesser extent
2	in other regions of the TB airway tree, showing that O_3 is delivered to these
3	compartments as well, although in a smaller dose. These studies agree with earlier model
4	predictions showing that the tissue O ₃ dose (O ₃ flux to liquid-tissue interface) was low in
5	the trachea, increased to a maximum in the terminal bronchioles and the CAR, and then
6	rapidly decreased in the alveolar region (Miller et al., 1985). It was also predicted that the
7	net O ₃ dose (total absorption, O ₃ flux to air-liquid interface) gradually decreased with
8	distal progression from the trachea to the end of the TB region and then rapidly decreased
9	in the alveolar region. Despite the exclusion of the URT and appreciable O_3 reactions
10	with ELF constituents after the 16th generation, the results from the model agree with
11	experimental results showing that the greatest O ₃ tissue dose was received in the CAR
12	(<u>Miller et al., 1985</u>).
13	Inhomogeneity in the RT structure may affect the dose delivered to this target site.
14	Models have predicted that the farther the PAR is from the trachea, the less the O ₃ tissue
15	dose to the region. Ultman and Anjilvel (1990) and Overton and Graham (1989)
16	predicted approximately a 50 to 300% greater PAR dose for the shortest path relative to
17	the longest path in humans and rats, respectively. In addition, Mercer et al. (1991) found
18	that both path distance and ventilatory unit size affected dose. The variation of O_3 dose
19	among anatomically equivalent ventilatory units was predicted to vary as much as 6-fold,

among anatomically equivalent ventilatory units was predicted to vary as much as 6-fold, as a function of path length from the trachea. This could have implications in regional damage to the LRT, such that even though the average LRT dose may be at a level where health effects would not be predicted, local regions of the RT may receive considerably higher than average doses and therefore be at greater risk of effects.

5.2.2.3 Upper Respiratory Tract Ozone Removal and Dose

24	Total O ₃ uptake in the entire RT in rats and guinea pigs ranges from 40-54% efficient
25	(Hatch et al., 1989; Wiester et al., 1988; Wiester et al., 1987), while in humans at rest it
26	ranges from 80-95% efficient (Hu et al., 1992). The URT provides a defense against O_3
27	entering the lungs by removing half of the O_3 that will be absorbed from the airstream. In
28	both animals and humans, about 50% of the O_3 that was absorbed in the RT was removed
29	in the head (nose, mouth, and pharynx), about 7% in the larynx/trachea, and about 43% in
30	the lungs (Hu et al., 1992; Hatch et al., 1989; Miller et al., 1979). However, experimental
31	studies in dogs have reported 75-100% uptake in the URT (Yokoyama and Frank, 1972;
32	<u>Vaughan et al., 1969</u>). The fraction of O_3 taken up was inversely related to flow rate and
33	to inlet O_3 concentration (<u>Yokoyama and Frank, 1972</u> ; <u>Vaughan et al., 1969</u>). The
34	limiting factors in nasal O3 uptake were simultaneous diffusion and chemical reaction of
35	O_3 in the nasal ELF layer (Santiago et al., 2001). The ELF layer in the nose is thicker

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22

1 than in the rest of the RT, and mathematical estimates predicted that O_3 penetrates less 2 than the thickness of the ELF layer; reaction products are likely the agents damaging the 3 nasal tissue and not O_3 itself. It was hypothesized that the nasal non-linear kinetics of O_3 4 uptake fraction result from the depleting substrates in the nasal ELF becoming the 5 limiting factor of the reaction (Santiago et al., 2001).

6 Uptake efficiencies have been measured for various segments of the URT (Table 5-1). 7 Gerrity et al. (1995) reported unidirectional uptake efficiencies of O₃ inhaled from a 8 mouthpiece; of 17.6% from the mouth to vocal cords, 9.5% from the vocal cords to the 9 upper trachea (totaling 27.1%), 8.4% from the upper trachea to the main bifurcation 10 carina (totaling 35.5%), and essentially zero between the carina and the bronchus 11 intermedius (totaling 32.5%). These values are lower than those calculated by Hu et al. 12 (1992) that reported uptake efficiencies of 21, 36, 44, and 46% during a complete breath 13 in which an O_3 bolus penetrated between the mouth and the vocal cords, the upper 14 trachea, the main bifurcation carina, and the bronchus intermedius, respectively. The 15 lower efficiencies seen in Gerrity et al. (1995) may have resulted because these 16 investigators measurements were based on inhalation alone or was caused by O_3 17 scrubbing by the mouthpiece.

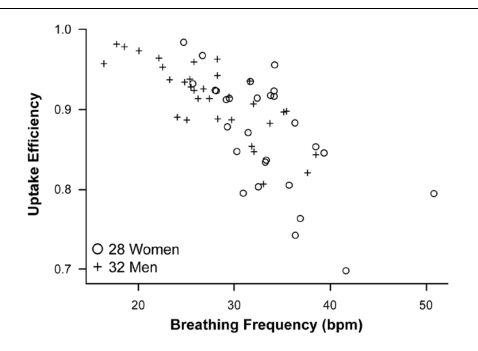
18 Past studies investigating nasal uptake of O_3 have shown that the nose partially protects 19 the LRT from damage from inspired O_3 (Santiago et al., 2001; Gerrity et al., 1988). 20 Sawyer et al. (2007) further investigated nasal uptake of O_3 in healthy adults during 21 exercise. Fractional O_3 uptake, acoustic rhinometry (AR), and nasal NO measurements 22 were taken on ten adults (8 women, 2 men) exposed to 200 ppb O₃ before and after 23 moderate exercise at two flow rates (10 and 20 L/min). The percent nasal uptake of O_3 24 was ~50% greater at 10 L/min compared to 20 L/min both pre- and post-exercise. 25 However, the inhaled O_3 dose delivered to the LRT (i.e., flow rate \times exposure 26 concentration \times (1 - nasal absorbed fraction)) was 1.6-fold greater at the higher flow than 27 at the lower flow (2.5 compared to 0.9 ppm·L/min). These results are similar to those 28 published earlier that found air pollutant retention increased with increasing airflow by 29 more than what would be predicted by just the increased partial pressure difference of the 30 gas (Aharonson et al., 1974). Prior exercise did not affect O₃ uptake at either flow rate, 31 but did significantly increase nasal volume (Vn) and AR measurements of nasal 32 cross-sectional area (minimum cross-sectional area (MCA) that corresponds to the nasal 33 valve, CSA2 that corresponds to the anterior edge of the nasal turbinates, and CSA3 that 34 corresponds to the posterior edge of the nasal turbinates) ($p \le 0.05$) (Sawyer et al., 2007). 35 Conversely, exercise decreased nasal resistance (Rn) (p < 0.01) and NO production 36 (nonsignificant, p > 0.05). The change in Vn and CSA2:MCA ratio was correlated with 37 the percent change in nasal uptake, however the overall effect was small and sensitive to 38 elimination of outliers and gender segregation.

1Overall, the majority of studies suggest that the URT removes about half of the O3 that2will be absorbed by reactions in the nasal ELF. The exact uptake efficiency will change3due to variations in flow rate and inhaled concentration.

5.2.2.4 Lower Respiratory Tract Ozone Uptake and Dose

4 Approximately 43% of the O_3 absorption occurs in the LRT of both humans and animals. 5 Models predicted that the net O_3 dose decreases distally from the trachea toward the end 6 of the TB region and then rapidly decreases in the alveolar region (Miller et al., 1985). 7 Further, these models predicted low tissue O₃ dose in the trachea and large bronchi. 8 Uptake efficiency depends on a number of variables, including O₃ exposure 9 concentration, exposure time, and breathing pattern. For breaths of similar waveforms, 10 respiratory patterns are uniquely described by breathing frequency (f_B) and tidal volume 11 (V_T) ; by minute ventilation ($\dot{V}_E = f_B \times V_T$) and f_B ; or by \dot{V}_E and V_T . Simulations from the 12 Overton et al. (1996) single-path anatomical respiratory tract model, where the upper and 13 lower respiratory tracts were modeled but uptake by the URT was not considered, 14 predicted that fractional uptake and PAR O3 dose increased with VT when fB was held 15 constant. Likewise, experimental studies found that O_3 uptake was positively correlated 16 with changes in V_T (Ultman et al., 2004; Gerrity et al., 1988). Also, O₃ exposure led to a 17 reflex mediated increase in f_B and reduction in V_T, hypothesized to be protective by 18 decreasing the dose delivered to the lung at a particular \dot{V}_{E} (Gerrity et al., 1994). Nasal O₃ 19 uptake efficiency was inversely proportional to flow rate (Santiago et al., 2001), so that 20 an increase in \dot{V}_E will increase O_3 delivery to the lower airways. At a fixed \dot{V}_E , increasing 21 V_T (corresponding to decreasing f_B) drove O_3 deeper into the lungs and increased total 22 respiratory uptake efficiency (Figure 5-4) (Ultman et al., 2004; Wiester et al., 1996a; 23 Gerrity et al., 1988). Modeling predicted a decrease in fractional uptake with increased $f_{\rm B}$ 24 when V_T was held constant, but an increase in PAR dose with increased f_B (Overton et 25 al., 1996). Similarly, increased f_B (80 - 160 bpm) and shallow breathing in rats decreased midlevel tracheal ¹⁸O content and an increased ¹⁸O content in the mainstem bronchi 26 27 (Alfaro et al., 2004). This dependence may be a result of frequency-induced alterations in 28 contact time that affects the first-order absorption rate for O_3 (Postlethwait et al., 1994). 29 Also, an association of O_3 uptake efficiency was found with \dot{V}_E and exposure time. 30 Increasing flow leads to deeper penetration of O_3 into the lung, such that a smaller 31 fraction of O_3 is absorbed in the URT and uptake shifts to the TB airways and respiratory airspaces (Nodelman and Ultman, 1999; Hu et al., 1994; Ultman et al., 1994). Hu et al. 32 33 (1994) and Ultman et al. (1994) found that O₃ absorption increased with volumetric 34 penetration (Vp) of a bolus of O_3 into the RT. Ozone uptake efficiency and Vp were not

1	affected by bolus O ₃ concentration (Kabel et al., 1994; Hu et al., 1992), indicating that
2	under these experimental conditions O_3 uptake was a linear absorption process, where the
3	diffusion and chemical reaction rates of O_3 were proportional to the O_3 concentration.
4	The absorption relationship would not be linear once interfacial mass transfer is
5	saturated. As mentioned above, a weak negative relationship between O3 concentration
6	and uptake efficiency was reported for the nasal cavities by Santiago et al. (2001). Rigas
7	et al. (2000) also found a weak but significant negative dependence of O_3 concentration
8	on RT uptake efficiency in exercising individuals. This study also found that exposure
9	time had a small but significant influence on uptake efficiency; however, this negative
10	dependence may be an artifact of progressive depletion of reactive substrates from the
11	ELF.



Note: Subjects breathed 250 ppb O_3 oronasally via a breathing mask. The uptake efficiency was well correlated with breathing frequency (r = -0.723, p <0.001) and tidal volume (not illustrated; r = 0.490, p <0.001). Source: Reprinted with permission of Health Effects Institute (<u>Ultman et al., 2004</u>).

Figure 5-4 Total ozone uptake efficiency as a function of breathing frequency at a constant minute ventilation of 30 L/min.

12	Past studies have shown that O ₃ -induced epithelial damage to the lung occurs with a
13	reproducible pattern of severity between daughter branches of individual bifurcations that
14	is dependent on the O_3 concentration-time profile of the inhaled gas. A 3-D
15	computational fluid dynamics model was created to investigate the O ₃ transport in a

1	single airway bifurcation (Taylor et al., 2007). The model consisted of one parent branch
2	and two symmetrical daughter branches with a branching angle of 90° and a sharp carinal
3	ridge. Various flow scenarios were simulated using Reynolds numbers (Re) ranging from
4	100 to 500. The Re that corresponds to a certain airway generation is dependent upon
5	both lung size and \dot{V}_{E} , such that the range in Re from 100-500 would encompass
6	generations 1-5, 3-7, and 6-10 for an adult during quiet breathing, light exertion, and
7	heavy exercise, respectively, whereas the same Re range corresponds to generations 0-4,
8	1-6, and 4-8 for a 4-year-old child. This model predicted velocity distributions that were
9	consistent with earlier work of <u>Schroter and Sudlow (1969</u>), and also reported O_3
10	concentration and wall uptake distributions. The model predicted that during inspiration,
11	the velocity and O ₃ concentration distribution were axisymmetric throughout the parent
12	branch, but skewed towards the inner wall within the daughter branches. During
13	expiration, the model predicted that the velocity and O ₃ concentration distribution was
14	slightly skewed towards the outer walls of the daughter branches. Hot spots of wall flux
15	existed at the carina during inspiration and expiration with Re >100. Additional hot spots
16	were found during expiration on the parent branch wall downstream of the branching
17	region.

18Overall O_3 inhalation uptake in humans is over 80% efficient, but the exact efficiency19that determines how much O_3 is available at longitudinally distributed compartments in20the lung is sensitive to changes in V_T , f_B , and to a minor extent, exposure time.

5.2.2.5 Mode of Breathing

21	Ozone uptake and distribution is sensitive to the mode of breathing. Variability in TB
22	airways volume had a weaker influence on O ₃ absorption during nasal breathing
23	compared to oral breathing. This could be a result of O_3 scrubbing in the nasal
24	passageways that are bypassed by oral breathing. Studies by Ultman and colleagues using
25	bolus inhalation demonstrated that O_3 uptake fraction was greater during nasal breathing
26	than during oral breathing at each Vp (e.g., 0.90 during nasal breathing and 0.80 during
27	oral breathing at 150 mL/sec and 0.45 during nasal breathing and 0.25 during oral
28	breathing at 1,000 mL/sec) (Nodelman and Ultman, 1999; Kabel et al., 1994; Ultman et
29	<u>al., 1994</u>). Therefore, oral breathing results in deeper penetration of O_3 into the RT with a
30	higher absorbed fraction in the TB and alveolar airways (Nodelman and Ultman, 1999).
31	Similar results were obtained from O ₃ uptake studies in dogs (Yokoyama and Frank,
32	<u>1972</u>). Earlier human studies suggested that oral or oronasal breathing results in a higher
33	O ₃ uptake efficiency than nasal breathing (Wiester et al., 1996a; Gerrity et al., 1988).
34	Overall, the mode of breathing may have a seemingly small effect on the RT uptake

1efficiency; however, it does play an important role in the distribution of O3 deposited in2the distal airways.

5.2.2.6 Interindividual Variability in Dose

3 Similarly exposed individuals vary in the amount of actual dose delivered to the LRT 4 (Santiago et al., 2001; Rigas et al., 2000; Bush et al., 1996). Interindividual variability 5 accounted for between 10-50% of the absolute variability in O₃ uptake measurements 6 (Santiago et al., 2001; Rigas et al., 2000). When concentration, time, and \dot{V}_E were held 7 constant, fractional absorption ranged from 0.80 to 0.91 (Rigas et al., 2000). It has been 8 hypothesized that interindividual variation in O_3 induced responses such as FEV₁ is the 9 result of interindividual variation in net dose or regional O₃ uptake among exposed 10 individuals. 11 Recent studies have reiterated the importance of intersubject variation in O_3 uptake. The 12 intersubject variability in nasal O₃ uptake determined by Sawyer et al. (2007) ranged 13 from 26.8 to 65.4% (pre- and post-exercise). A second study investigating the use of the 14 CO_2 expirogram to quantify pulmonary responses to O_3 found that intersubject variability 15 accounted for 50% of the overall variance in the study (Taylor et al., 2006). 16 Variability in net or tissue dose may be attributed to differences in the pulmonary 17 physiology, anatomy, and biochemistry. Since the URT and TB airways remove the 18 majority of inhaled O_3 before it reaches the gas exchange region, the volume and surface 19 area of these airways will influence O_3 uptake. Models predicted that fractional O_3 uptake 20 and PAR dose (flux of O₃ to the PAR surfaces divided by exposure concentration) increase with decreasing TB volume and decreasing TB region expansion. On the

21 22 contrary, alveolar expansion had minimal effect on uptake efficiency as relatively little 23 O₃ reaches the peripheral lung (Bush et al., 2001; Overton et al., 1996). Ozone uptake 24 was virtually complete by the time O₃ reaches the alveolar spaces of the lung 25 (Postlethwait et al., 1994). Experimental studies have found that differences in TB 26 volumes may account for 75% of the variation in absorption between subjects (Ultman et 27 al., 2004). In support of this concept, regression analysis showed that O_3 absorption was 28 positively correlated with anatomical dead space (V_D) and TB volume (i.e., V_D minus 29 V_{URT}), but not total lung capacity (TLC), forced vital capacity (FVC), or functional 30 residual capacity (FRC) (Ultman et al., 2004; Bush et al., 1996; Hu et al., 1994; 31 Postlethwait et al., 1994). Variability in V_D was correlated more with the variability in the 32 TB volume than the URT volume. Similarly, uptake was correlated with changes in 33 individual bronchial cross-sectional area, indicating that changes in cross-sectional area 34 available for gas diffusion are related to overall O₃ retention (Reeser et al., 2005; Ultman

1	et al., 2004). When coupled, these results suggest that the larger surface-to-volume ratio
2	associated with the smaller airways in women enhances local O_3 uptake, thereby reducing
3	the distal penetration volume of O_3 into the female respiratory system. When absorption
4	data were normalized to Vp/V_D , variability attributed to gender differences were not
5	distinguishable (<u>Bush et al., 1996</u>). These studies provide support to the RT anatomy,
6	especially the TB volume and surface area, playing a key role in variability of O ₃ uptake
7	between individuals.
8	In addition, variability between individuals is influenced by age. Overton and Graham
9	(1989) predicted that the total mass of O_3 absorbed per minute (in units of: μ g/min per
10	$[\mu g/m^3 \text{ of ambient } O_3])$ increased with age from birth to adulthood. This model predicted
11	that during quiet breathing the LRT distribution of absorbed O_3 and the CAR O_3 tissue
12	dose were not sensitive to age. However, during heavy exercise or work O ₃ uptake was
13	dependent on age. A physiologically based pharmacokinetic model simulating O_3 uptake
14	predicted that regional extraction of O3 was relatively insensitive to age, but extraction
15	per unit surface area was 2-fold to 8-fold higher in infants compared to adults, due to the
16	fact that children under age 5 have much a much smaller airway surface area in the
17	extrathoracic (nasal) and alveolar regions (Sarangapani et al., 2003). Additionally,
18	children tend to have a greater oral breathing contribution than adults at rest and during
19	exercise (Bennett et al., 2008; Becquemin et al., 1999; James et al., 1997). Normalized to
20	lung surface area, the dose rate to the lower airways of children compared to adults is
21	increased further because children breathe at higher minute ventilations relative to their
22	lung volumes.
23	Smoking history, with its known increase in mucus production, was not found to affect
24	the fractional uptake of a bolus of O_3 in apparently healthy smokers with limited smoking
25	history (<u>Bates et al., 2009</u>). Despite similar internal O_3 dose distribution, the smokers
26	exhibited greater pulmonary responses to O_3 bolus exposures, measured as FEV_1
27	decrements and increases in the normalized slope of the alveolar plateau (S_N). This was
28	contrary to previous studies conducted in smokers with a greater smoking history that
29	found decreased O_3 induced decrements in FEV_1 in smokers during continuous O_3
30	exposure (Frampton et al., 1997a; Emmons and Foster, 1991).

5.2.2.7 Physical Activity

31	Exercise increases the overall exposure of the lung to inhaled contaminants due, in most
32	part, to the increased intake of air. Thus, human studies have used exercise, at a variety of
33	activity levels, to enhance the effects of O_3 (<u>Table 5-2</u>). Further explanation of the effects
34	of physical activity on ventilation can be found in Chapters 4 and 6. <u>Table 4-5</u> presents

Table 5-2 General adult human inhalation rates by activity levels.

Activity Level	Inhalation Rate
Light	2 to 3 × resting \dot{V}_{E}^{a}
Moderate	4 to 6 × resting \dot{V}_E
Heavy	7 to 8 × resting \dot{V}_E
Very Heavy	>9 × resting \dot{V}_E

^aResting \dot{V}_E approximates 8 L/min

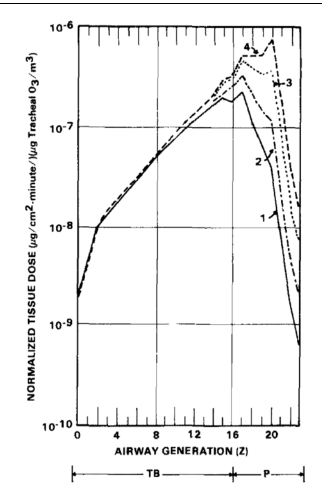
Source: <u>U.S. EPA (1986</u>).

1

2

3	As exercise increases from a light to moderate level, V_T increases. This increase in V_T is
4	achieved by encroaching upon both the inspiratory and expiratory reserve volumes of the
5	lung (Dempsey et al., 1990). After V_T reaches about 50% of the vital capacity, generally
6	during heavy exercise, further increases in ventilation are achieved by increasing f_B .
7	Ventilatory demands of very heavy exercise require airway flow rates that often exceed
8	10 times resting levels and V_T that approach 5 times resting levels (<u>Dempsey et al.</u> ,
9	<u>2008</u>).
10	This increase in V_T and flow associated with exercise in humans shifts the net O_3 dose
11	further into the periphery of the RT causing a disproportionate increase in distal lung
12	tissue dose. In addition to increasing the bulk transport of O ₃ into the lung, exercise also
13	leads to a switch from nasal to oronasal breathing. Higher ventilatory demand
14	necessitates a lower-resistance path through the mouth. Modeling heavy exercise by
15	increasing ventilatory parameters from normal respiration levels predicted a 10-fold
16	increase in total mass uptake of O_3 (<u>Miller et al., 1985</u>). This model also predicted that as
17	exercise and ventilatory demand increased, the maximum tissue dose, the O3 reaching the
18	tissues, moved distally into the RT (Figure 5-5). By increasing flow to what is common
19	in moderate or heavy exercise (respiratory flow = 45-60 L/min compared to 15 L/min),
20	the URT absorbed a smaller fraction of the O_3 (~0.50 at low flow rate to 0.10 at high flow
21	rate); however, the trachea and more distal TB airways received higher doses during
22	higher flow rates than at lower flow rates (0.65 absorbed in the lower TB airways, and
23	0.25 absorbed in the alveolar zone with high flow compared to 0.5 in the TB with almost
24	no O_3 reaching the alveolar zone at low flow) (<u>Hu et al., 1994</u>). The same shift in the O_3
25	dose distribution more distally in the lung occurred in other studies mimicking the effects
26	of exercise (Nodelman and Ultman, 1999). Also, LRT uptake efficiency was sensitive to
27	age only under exercise conditions (<u>Overton and Graham, 1989</u>). The total mass of O_3

absorbed per minute (μ g/min per [μ g/m³ of ambient O₃]) was predicted to increase with 1 2 age during heavy work or exercise. A recent study by Sawyer et al. (2007) approximated 3 that doubling minute ventilation led to only a 1.6-fold higher delivered dose rate of O_3 to 4 the lung (delivered dose was calculated as: flow rate \times [O₃ ppm] \times (100-percent nasal O₃) 5 uptake)). Past models have predicted the increase in uptake during exercise is distributed 6 unevenly in the RT compartments and regions. Tissue and mucus layer dose in the TB 7 region increased ~1.4-fold during heavy exercise compared to resting conditions, whereas 8 the alveolar region surfactant and tissue uptake increased by factors of 5.2 and 13.6, 9 respectively (Miller et al., 1985).



Note: Curve 1: $V_T = 500 \text{ mL}$; $f_B = 15 \text{ breaths/min}$. Curve 2: $V_T = 1,000 \text{ mL}$; $f_B = 15 \text{ breaths/min}$. Curve 3: $V_T = 1,750 \text{ mL}$; $f_B = 20.3 \text{ breaths/min}$. Curve 4: $V_T = 2,250 \text{ mL}$; $f_B = 30 \text{ breaths/min}$. TB = tracheobronchial region; P = pulmonary region. Source: Reprinted with permission of Elsevier (<u>Miller et al., 1985</u>).

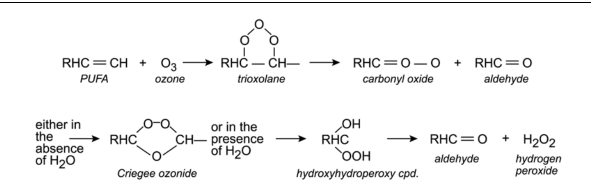
Figure 5-5 Modeled effect of exercise on tissue dose of the LRT.

5.2.2.8 Summary

1	In summary, O ₃ uptake is affected by complex interactions between a number of factors
2	including RT morphology, breathing route, frequency, and volume, physicochemical
3	properties of the gas, physical processes of gas transport, as well as the physical and
4	chemical properties of the ELF and tissue layers. The role of these processes varies
5	throughout the length of the RT and as O_3 moves from the gas into liquid compartments
6	of the RT. The primary uptake site of O_3 delivery to the lung epithelium is believed to be
7	the CAR, however inhomogeneity in the RT structure may affect the dose delivered to
8	this target site with larger path lengths leading to smaller locally delivered doses. This
9	could have implications in regional damage to the LRT, such that even though the
10	average LRT dose may be at a level where health effects would not be predicted , local
11	regions of the RT may receive considerably higher than average doses and therefore be at
12	greater risk of effects. Recent studies have provided evidence for hot spots of O ₃ flux
13	around bifurcations in airways. Experimental studies and models have suggested that the
14	net O ₃ dose gradually decreases distally from the trachea toward the end of the TB region
15	and then rapidly decreases in the alveolar region. However, the tissue O_3 dose is low in
16	the trachea, increases to a maximum in the terminal bronchioles and the CAR, and then
17	remidly decreased distally into the algorithm region
17	rapidly decreases distally into the alveolar region.
17	O_3 uptake efficiency is sensitive to a number of factors. Fractional absorption will
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5.2.3 Ozone Reactions and Reaction Products

1	Ozone dose is affected by the chemical reactions or the products of these reactions that
2	result from O ₃ exposure. The process by which O ₃ moves from the airway lumen and into
3	the ELF is related to the coupled diffusion and chemical reactions occurring in ELF is
4	called "reactive absorption". Ozone is chemically reactive with a wide spectrum of
5	biomolecules and numerous studies have evaluated the loss of specific molecules such as
6	GSH and the appearance of plausible products such as nonanal. Both in vitro and in vivo
7	studies contribute to the understanding of O ₃ reactions and reaction products.
8	Ozone may interact with many of the components in the ELF including phospholipids,
9	neutral lipids like cholesterol, free fatty acids, proteins, and low molecular weight
10	antioxidants as has been demonstrated in in vitro studies (Perez-Gil, 2008; Uppu et al.,
11	<u>1995</u>). It was estimated that 88% of the O_3 that does not come in contact with
12	antioxidants will react with unsaturated fatty acids in the ELF including those contained
13	within phospholipids or neutral lipids (Uppu et al., 1995). Ozone reacts with the double
14	bond of unsaturated fatty acids to form stable and less reactive ozonide, aldehyde, and
15	hydroperoxide reaction products via chemical reactions such as the Criegee ozonolysis
16	mechanism (Figure 5-6) (Pryor et al., 1991). Lipid ozonation products, such as the
17	aldehydes hexanal, heptanal, and nonanal, have been recovered after O_3 exposure in
18	human BAL fluid (BALF), rat BALF, isolated rat lung, and in vitro systems (Frampton et
19	al., 1999; Postlethwait et al., 1998; Pryor et al., 1996). Adducts of the aldehyde
20	4-hydroxynonenal were found in human alveolar macrophages after O3 exposure in vivo
21	(Hamilton et al., 1998). Polyunsaturated fatty acid (PUFA) reactions are limited by the
22	availability of O ₃ since lipids are so abundant in the ELF. Yields of O ₃ -induced aldehydes
23	were increased by the decrease in other substrates such as ascorbic acid (AH ₂)
24	(Postlethwait et al., 1998). Free radicals are also generated during O ₃ -mediated oxidation
25	reactions with PUFA (Pryor, 1994). These reactions are reduced by the presence of the
26	lipid-soluble free radical scavenger α-tocopherol (α-TOH) (Pryor, 1994; Fujita et al.,
27	1987; Pryor, 1976). PUFA reactions may not generate sufficient bioactive materials to
28	account for acute cell injury, however only modest amounts of products may be
29	necessary to induce cytotoxicity (Postlethwait and Ultman, 2001; Postlethwait et al.,
30	<u>1998</u>).



Note: Not all secondary reaction products are shown. Source: <u>U.S. EPA (2006b</u>).

Figure 5-6 Schematic overview of ozone interaction with PUFA in ELF and lung cells.

1	Cholesterol is the most abundant neutral lipid in human ELF. Reaction of cholesterol
2	with O ₃ results in biologically active cholesterol products such as the oxysterols,
3	β-epoxide and 6-oxo-3,5-diol (<u>Murphy and Johnson, 2008; Pulfer et al., 2005; Pulfer and</u>
4	Murphy, 2004). Product yields depend on ozonolysis conditions, however cholesterol
5	ozonolysis products form in similar abundance to phospholipid-derived ozonolysis
6	products in rat ELF (Pulfer and Murphy, 2004).
7	The ELF also contains proteins derived from blood plasma as well as proteins secreted by
8	surface epithelial cells. Ozone reactions with proteins have been studied by their in vitro
9	reactions as well as reactions of their constituent amino acids (the most reactive of which
10	are cysteine, histidine, methionine, tyrosine, and tryptophan). Ozone preferentially reacts
11	with biomolecules in the following order: thiosulfate >ascorbate >cysteine \approx methionine
12	>glutathione (Kanofsky and Sima, 1995). Rate constants for the reaction of amino acids
13	with O ₃ vary between studies due to differing reaction conditions and assumptions;
14	however aliphatic amino acids were consistently very slow to react with O ₃ (e.g., alanine:
15	25-100 moles/L/sec) (Kanofsky and Sima, 1995; Ignatenko and Cherenkevich, 1985;
16	Pryor et al., 1984; Hoigné and Bader, 1983). Uppu et al. (1995) predicted that 12% of
17	inhaled O_3 that does not react with antioxidants will react with proteins in the ELF.
18	Reactions of O ₃ with low molecular weight antioxidants have been extensively studied.
19	The consumption of antioxidants such as uric acid (UA), ascorbate (AH ₂), and reduced
20	glutathione (GSH) by O ₃ was linear with time and positively correlated with initial
21	substrate concentration and O ₃ concentration (Mudway and Kelly, 1998; Mudway et al.,
22	<u>1996</u>). Endogenous antioxidants are present in relatively high concentrations in the ELF
23	of the human airways (obtained as BALF) and display high (but not equal) intrinsic

1	reactivities toward O ₃ . In individual and in limited composite mixtures, UA was the most
2	reactive antioxidant tested, followed by AH ₂ (Mudway and Kelly, 1998). GSH was
3	consistently less reactive than UA or AH ₂ (Mudway and Kelly, 1998; Mudway et al.,
4	1996; Kanofsky and Sima, 1995). To quantify these reactions, Kermani et al. (2006)
5	evaluated the interfacial exposure of aqueous solutions of UA, AH ₂ , and GSH
6	(50-200 μ M) with O ₃ (1-5 ppm). Similar to the results of Mudway and Kelly (1998), this
7	study found the hierarchy in reactivity between O_3 and these antioxidants to be
8	$UA \ge AH_2 >> GSH$. UA and AH_2 shared a 1:1 stoichiometry with O_3 , whereas 2.5 moles of
9	GSH were consumed per mole of O ₃ . Using these stoichiometries, reaction rate constants
10	were derived $(5.8 \times 10^4 \text{ M}^{-1} \text{ sec}^{-1}, 5.5 \times 10^4 \text{ M}^{-1} \text{ sec}^{-1}, \text{ and } 57.5 \text{ M}^{-0.75} \text{ sec}^{-1} [20.9 \text{ M}^{-1} \text{ sec}^{-1}]$ for
11	the reaction of O ₃ with UA, AH ₂ , and GSH, respectively). Other studies report reactive
12	rate constants that are two to three orders of magnitude larger, however these studies used
13	higher concentrations of O ₃ and antioxidants under less physiologically relevant
14	experimental conditions (Kanofsky and Sima, 1995; Giamalva et al., 1985; Pryor et al.,
15	<u>1984</u>). However, O_3 acts through competition kinetics so the effective concentration of
16	the reactants present in the ELF will determine the reactions that occur in vivo. For
17	example, the pKa of GSH is about 8.7 so that at physiological pH very little is in the
18	reactive form of thiolate (GS ⁻). On the other hand, ascorbic acid has a pKa of about 4.2 so
19	it exists almost entirely as ascorbate (AH ⁻) in the ELF. Thus, the effective concentration
20	of GSH that is available to react with O_3 will be much lower than that of ascorbate in
21	ELF.
21 22	ELF. A series of studies used new techniques to investigate the reaction products resulting
22	A series of studies used new techniques to investigate the reaction products resulting
22 23	A series of studies used new techniques to investigate the reaction products resulting from initial air-liquid interface interactions of O_3 with ELF components
22 23 24	A series of studies used new techniques to investigate the reaction products resulting from initial air-liquid interface interactions of O_3 with ELF components (e.g., antioxidants and proteins) in ~1 millisecond (Enami et al., 2009a, b, c, 2008a, b).
22 23 24 25	A series of studies used new techniques to investigate the reaction products resulting from initial air-liquid interface interactions of O ₃ with ELF components (e.g., antioxidants and proteins) in ~1 millisecond (Enami et al., 2009a, b, c, 2008a, b). Solutions of aqueous UA, AH ₂ , GSH, α-TOH, and protein cysteines (CyS) were sprayed
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1	one-electron acceptor but very reactive O-atom donor, products of the interaction of O_3
2	with UA, AH ₂ , GSH, CyS, and α -TOH result from addition of <i>n</i> O-atoms (<i>n</i> = 1-4). These
3	products included epoxides (e.g., U-O ⁻), peroxides (e.g., U-O ₂ ⁻), and ozonides
4	(e.g., U-O ₃ ⁻). For instance, GSH was oxidized to sulfonates (GSO_3^{-}/GSO_3^{-2}), not
5	glutathione disulfide (GSSG) by O ₃ (Enami et al., 2009b). However, it is possible that
6	other oxidative species are oxidizing GSH in vivo, since sulfonates are not detected in O ₃
7	exposed ELF whereas GSSG is. This is also supported by the fact that O_3 is much less
8	reactive with GSH than other antioxidants, such that $<3\%$ of O ₃ will be scavenged by
9	GSH when in equimolar amounts with AH_2 (Enami et al., 2009b).
10	This series of studies also demonstrated that ozonolysis product yields and formation
11	were affected by pH. Acidified conditions (pH \approx 3-4), such as those that may result from
12	acidic particulate exposure or pathological conditions like asthma (pH \approx 6), decreased the
13	scavenging ability of UA and GSH for O_3 ; such that at low pH, the scavenging of O_3
14	must be taken over by other antioxidants, such as AH_2 (Enami et al., 2009b, 2008b).
15	Also, under acidic conditions (pH \approx 5), the ozonolysis products of AH ₂ shifted from the
16	innocuous dehydroascorbic acid to the more persistent products, AH ₂ ozonide and
17	threonic acid (Enami et al., 2008a). It is possible that the acidification of the ELF by
18	acidic copollutant exposure will increase the toxicity of O_3 by preventing some
19	antioxidant reactions and shifting the reaction products to more persistent compounds.
20	Cinc. FLE anistana and an interactivity in an added to be the Operation in an hotest
20	Since ELF exists as a complex mixture, it is important to look at O ₃ reactivity in substrate
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21 22	mixtures. Individual antioxidant consumption rates decreased as the substrate mixture complexity increased (e.g., antioxidant mixtures and albumin addition) (Mudway and
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- 1 al., 2005). The presence of UA or bovine serum albumin protected against lipid and 2 protein oxidation resulting from the reaction of O_3 and AH_2 (Ballinger et al., 2005). This 3 study provided evidence that antioxidants may paradoxically facilitate O₃-mediated 4 damage. This apparent contradiction should be viewed in terms of the 5 concentration-dependent role of the ELF antioxidants. Reactions between O_3 and 6 antioxidant species exhibited a biphasic concentration response, with oxidation of protein 7 and lipid occurring at lower, but not higher, concentrations of antioxidant. In this way, 8 endogenous reactants led to the formation of secondary oxidation products that were 9 injurious and also led to quenching reactions that were protective. Moreover, the formation of secondary oxidation products mediated by some antioxidants was opposed 10 by quenching reactions involving other antioxidants. 11
- 12Alterations in ELF composition can result in alterations in O_3 uptake. Bolus O_3 uptake in13human subjects can be decreased by previous continuous O_3 exposure (120-360 ppb),14possibly due to depletion of compounds able to react with O_3 (Rigas et al., 1997; Asplund15et al., 1996). Conversely, O_3 (360 ppb) bolus uptake was increased with prior NO216(360-720 ppb) or SO2 (360 ppb) exposure (Rigas et al., 1997). It was hypothesized that17this increased fractional absorption of O_3 could be due to increased production of reactive18substrates in the ELF due to oxidant-induced airway inflammation.
- 19 Besides AH₂, GSH and UA, the ELF contains numerous antioxidant substances that 20 appear to be an important cellular defense against O_3 including α -TOH, albumin, 21 ceruloplasmin, lactoferrin, mucins, and transferrin (Mudway et al., 2006; Freed et al., 22 1999). The level and type of antioxidant present in ELF varies between species, regions 23 of the RT, and can be altered by O₃ exposure. Mechanisms underlying the regional 24 variability are not well-understood. It is thought that both plasma ultrafiltrate and locally 25 secreted substances contribute to the antioxidant content of the ELF (Mudway et al., 26 2006; Freed et al., 1999). In the case of UA, the major source appears to be the plasma 27 (Peden et al., 1995). Repletion of UA in nasal lavage fluid was demonstrated during 28 sequential nasal lavage in human subjects (Mudway et al., 1999a). When these subjects, 29 exercising at a moderate level, were exposed to 200 ppb O₃ for 2 hours, nasal lavage 30 fluid UA was significantly decreased while plasma UA levels were significantly 31 increased (Mudway et al., 1999a). The finding that UA, but not AH₂ or GSH, was 32 depleted in nasal lavage fluid indicated that UA was the predominant antioxidant with 33 respect to O_3 reactivity in the nasal cavity (Mudway et al., 1999a). However, in human 34 BALF samples, the mean consumption of AH₂ was greater than UA (Mudway et al., 35 1996). In addition, concentrations of UA were increased by cholinergic stimulation of the 36 airways in human subjects, which suggested that increased mucosal gland secretions were 37 an important source (Peden et al., 1993). Using the O_3 -specific antioxidant capacity assay 38 on human nasal lavage samples, Rutkowski et al. (2011) concluded that about 30% of the

1	antioxidant capacity of the nasal liquid lining layer was attributed to UA activity.
2	Additionally, more than 50% of the subject-to-subject differences in antioxidant capacity
3	were driven by differences in UA concentration. However, day-to-day within-subject
4	variations in measured antioxidant capacity were not related to the corresponding
5	variations in UA concentration in the nasal lavage fluid. Efforts to identify the
6	predominant antioxidant(s) in other RT regions besides the nasal cavity have failed to
7	yield definitive results.
8	Regulation of AH ₂ , GSH and α -TOH concentrations within the ELF is less clear than that
9	of UA (Mudway et al., 2006). In a sequential nasal lavage study in humans, wash-out of
10	AH ₂ and GSH occurred, indicating the absence of rapidly acting repletion mechanisms
11	(Mudway et al., 1999a). Other studies demonstrated increases in BALF GSH and
12	decreases in BALF and plasma AH ₂ levels several hours following O ₃ exposure (200 ppb
13	for 2 h, while exercising at a moderate level) (Mudway et al., 2001; Blomberg et al.,
14	1999; Mudway et al., 1999b). Other investigators have demonstrated cellular uptake of
15	oxidized AH ₂ by several cell types leading to intracellular reduction and export of
16	reduced AH ₂ (Welch et al., 1995). Studies with rats exposed to 0.4-1.1 ppm O_3 for
17	1-6 hours have shown consumption of AH_2 that correlates with O_3 exposure (Gunnison
18	and Hatch, 1999; Gunnison et al., 1996; Vincent et al., 1996b).
19	A body of evidence suggests that reaction of O3 within the ELF limits its diffusive
20	transport through the ELF; direct contact of O_3 with the apical membranes of the
21	underlying epithelial cells therefore might be negligible (Ballinger et al., 2005; Connor et
22	al., 2004; Postlethwait and Ultman, 2001; Pryor, 1992). This conclusion is based on
23	computational analyses and in vitro studies. Direct confirmation using in vivo studies is
24	lacking. Nevertheless, when predicting exposure-related outcomes across species and
25	anatomic sites, whether O ₃ directly contacts the apical membranes of the epithelial cells
26	is an important consideration, given that the extracellular surface milieu of the RT
27	appreciably varies in terms of the types and concentrations of the substrates present and
28	the thickness of the ELF.
29	For O_3 or its reaction products to gain access to the underlying cellular compartments, O_3
30	must diffuse at the air-liquid interface of the airway surface and travel through the ELF
31	layer. In vitro experiments have shown that O ₃ disappearance from the gas phase depends
32	on the characteristics of the ELF substrates (Postlethwait et al., 1998; Hu et al., 1994).
33	The ELF is comprised of the airway surface lining that includes the periciliary sol layer
34	and overlying mucus gel layer, and the alveolar surface lining that includes the subphase
35	of liquid and vesicular surfactant and the continuous surfactant monolayer (Bastacky et
36	al., 1995). There is a progressive decrease in ELF thickness and increase in interfacial
37	surface with progression from the TB region to the alveolus (Mercer et al., 1992). The

progressive thinning of the ELF while moving further down the RT decreases the radial
 distance O₃ or its reaction products must travel to reach the cells lining the RT.
 Taking into account the high reactivity and low water solubility of O₃, calculations
 suggest that O₃ will not penetrate ELF layers greater than 0.1 µm without being
 transformed to other more long-lived reactive species, thus initiating a reaction cascade

transformed to other more long-lived reactive species, thus initiating a reaction cascade (Pryor, 1992). These calculations utilize the Einstein-Smoluchowski equation (Equation 5-1) that combines Fick's second law of diffusion and a stochastic view of motion to compare the half-life of O₃ in the ELF layer to the time it takes, *t*, for O₃ to travel a distance, *d*, with a diffusion coefficient of D (~2×10⁻⁵ cm²/sec).

$t = d^2/2D$

Equation 5-1

10	The transit time through an ELF layer of 10^{-5} cm was estimated to be 2.5×10^{-6} seconds.
11	The half-life of O ₃ can be approximated by dividing the pseudo-first order rate constant,
12	k_1 , into ln 2. Pryor (1992) assumed the reaction rate constant 10 ⁹ L/mol/sec for O ₃ with
13	GSH and the concentration of GSH equaled 1 mM in the ELF. Using these values and
14	neglecting reactions of O_3 with other ELF species, the half-life of O_3 would be 7×10^{-7}
15	seconds. Under these assumptions of GSH concentration and ELF thickness, the half-life
16	of O_3 is about one third of the time necessary for O_3 to diffuse through the ELF layer.
17	Further, assuming that 0.5 ppm O_3 enters the trachea and the intrapulmonary gas-phase
18	concentration is reduced only 5 fold during transport to the terminal bronchioles, by using
19	a Henry's law constant and assuming equilibrium, the ELF O3 concentration could be
20	calculated to be $<1.4 \times 10^{-9}$ M or approximately 0.0014 μ M. Further, assuming that
21	ascorbate = 100 μ M, GSH = 300 μ M, and uric acid = 250 μ M, while ignoring unsaturated
22	lipids and reactive proteins, the most facile reactants would equate to an approximately
23	500,000-fold excess over O_3 . If one then assumes a lumped reaction rate constant of
24	$10^7 \text{ M}^{-1} \text{ sec}^{-1}$, any O ₃ in solution would be consumed by reaction almost instantaneously,
25	thereby constraining its diffusion as an unreacted species to within <0.1 µm, which is less
26	than the thickness values estimated for distal airway ELF. If unsaturated lipids
27	(~ $10^6 \text{ M}^{-1} \text{ sec}^{-1}$) and proteins (for which the rate constant will vary depending on low
28	pKa thiolates and other amino acid-reactive sites) are included, the penetration depth is
29	further reduced.
30	Similarly, model calculations of the nasal cavity based on diffusion equations and
31	reaction rates of O_3 with model substrates predict an O_3 penetration distance (0.5 μ m)
32	less than the thickness of the nasal lining layer (10 μ m) (Santiago et al., 2001).
33	A computational fluid dynamics model was able to predict experimentally measured

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 O_3 uptake when nasal mucus layer thickness was considered (<u>Cohen-Hubal et al., 1996</u>), reaffirming the importance of the resistance imparted by the ELF layer in dose and lesion patterns in the nasal passage.

- 4 Despite calculations and in vitro studies suggesting that reactions of O_3 with underlying 5 epithelial cells may be negligible, there is some evidence that suggests direct interaction 6 of O₃ with epithelial cells is possible. While moving distally in the lung, the ELF 7 thickness decreases and becomes ultrathin in the alveolar region, possibly allowing for 8 direct interaction of O_3 with the underlying epithelial cells. One definitive study 9 conducted in excised rat lung measured alveolar lining layer thickness over relatively flat 10 portions of the alveolar wall to be 0.14 μ m, to be 0.89 μ m at the alveolar wall junctions, 11 and 0.09 µm over the protruding features (Bastacky et al., 1995). The area-weighted 12 average thickness of the alveolar lining fluid was found to be about 0.2 μ m and the 13 alveolar lining layer was continuous over the entire alveolar surface measured. The 14 surface appeared smooth, and no epithelial surface features or macrophage features 15 protruded above the air-liquid interface. It was noted that measurements of alveolar lining 16 layer thickness were made in lungs prepared in a state of roughly 80% of total lung capacity, and as a result, the values reported would be approaching the lowest values 17 18 possible during the respiratory cycle. However, 4% of the surface area in the alveolar 19 compartment was covered by alveolar lining fluid layer of less than 20 nm (Bastacky et 20 al., 1995), suggesting the possibility that unreacted O_3 could penetrate to the cell layer in 21 this region. Further it remains a possibility that airways macrophages may protrude into 22 the gas phase, allowing for direct contact between O_3 and airways epithelial cells.
- 23 Still, direct reaction of O_3 with alveolar epithelial cells or macrophages may be limited by 24 the presence of dipalmitoyl phosphatidylcholine (DPPC), the major component of 25 surfactant, which has been shown in vitro to inhibit uptake of O_3 into an aqueous 26 compartment containing ascorbate, glutathione, and uric acid (Connor et al., 2004). 27 Further, the amount of O₃ available to the alveolar compartment may be limited by uptake of O_3 in nasal and TB compartments (Figure 5-5). In fact, the amount of ¹⁸O 28 29 reaction product was lower in the alveolar tissues than in TB tissues of rhesus monkeys immediately following a 2 hour exposure to 18 O-labeled O₃ (0.4 and 1 ppm) (Plopper et 30 31 al., 1998). These considerations illustrate the difficulty in determining whether O_3 reacts 32 directly with cells in the alveolar compartment.
- In some cases, however, with regard to the initiating mechanisms of cellular
 perturbations, the precise reactive species that encounters the epithelia might or might not
 have specificity to O₃ per se or to its secondary oxidants. Many of the measureable
 products formed as a consequence of O₃ exposure have limited specificity to O₃, such as
 4-hydroxynonenal that is formed by autoxidation, an event that can be initiated by O₃ but

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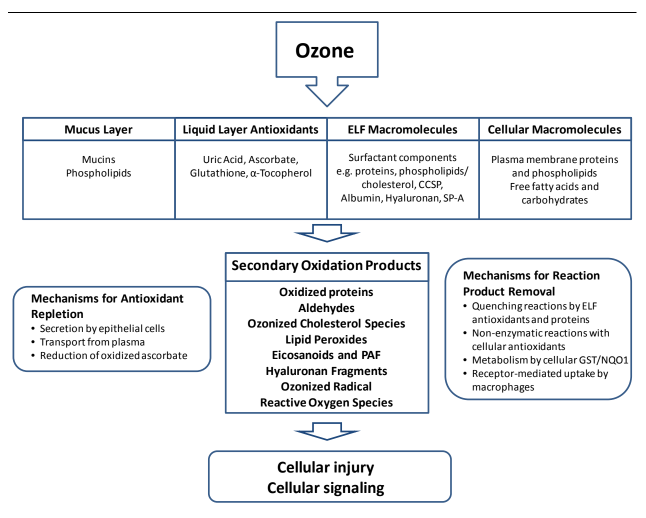
2

1	also by a multitude of other oxidants. Although some classes of lipid oxidation products
2	(e.g., specific aldehydes, cholesterol products) are specific to O_3 , measurement in either
3	BALF or in tissue does not necessarily provide insight on the compartment in which they
4	were formed (i.e., the ELF, cell membrane, intracellular space) because the ELF is a
5	dynamic compartment and, once formed, hydrophobic species can partition. Oxidation of
6	membrane components might produce similar cellular outcomes regardless of the
7	initiating oxidant. Lipid ozonides, which could be generated either within the ELF or
8	from ozonation of cell membrane unsaturated lipids, could bind to receptors, activate
9	signaling cascades, and act in other ways, making differences between pure extracellular
10	reaction and direct membrane reaction indistinguishable. Thus, in some cases
11	documenting whether O_3 per se reacts directly with cellular constituents might be
12	essential (despite the challenges of in vivo demonstrations), while in other cases precisely
13	where O ₃ reacts might be of less concern with regard to characterizing mechanisms of
14	health outcomes.
15	Thus, components of the ELE are project torgets for O and the resulting secondary

15Thus, components of the ELF are major targets for O_3 and the resulting secondary16oxidation products key mediators of toxicity in the airways (the role of reaction products17in O_3 -induced toxicity is discussed in Section 5.3). The reaction cascade resulting from18the interaction of O_3 with ELF substrates can then carry the oxidative burden deeper into19cells lining the RT to elicit the health effects observed.

5.2.3.1 Summary

20	The ELF is a complex mixture of lipids, proteins, and antioxidants that serve as the first
21	barrier and target for inhaled O_3 (Figure 5-7). The thickness of the lining fluid and mucus
22	layer is an important determinant of the dose of O_3 to the tissues. The antioxidant
23	substances present in the ELF appear in most cases to limit interaction of O_3 with
24	underlying tissues and to prevent penetration of O_3 deeper into the lung. The formation of
25	secondary oxidation products is likely related to the concentration of antioxidants present
26	and the quenching ability of the lining fluid. Mechanisms are present to replenish the
27	antioxidant substrate pools as well as to remove secondary reaction products from tissue
28	interactions. Important differences exist in the reaction rates for O_3 and these ELF
29	biomolecules and the reactivity of the resulting products. Overall, studies suggest that UA
30	and AH_2 are more reactive with O_3 than GSH, proteins, or lipids. In addition to
31	contributing to the driving force for O3 uptake, formation of secondary oxidation
32	products may lead to increased cellular injury and cell signaling (discussed in
33	Section 5.3). Studies indicate that the antioxidants might be participating in reactions
34	where the resulting secondary oxidation products might penetrate into the tissue layer and
35	lead to perturbations.



Note: Contents of this figure not discussed in Section <u>5.2</u> will be discussed in Section <u>5.3</u>. Clara cell secretory protein, CCSP; Surfactant Protein-A, SP-A; Platelet activating factor, PAF. Ozone will react with components of the ELF to produce reaction products that may lead to cellular injury and cell signaling as discussed in Section <u>5.3</u>.

Figure 5-7 Details of the ozone interaction with the airway ELF to form secondary oxidation products.

5.3 Possible Pathways/Modes of Action

5.3.1 Introduction

1	Mode of action refers to a sequence of key events and processes that result in a given
2	toxic effect (U.S. EPA, 2005). Elucidation of mechanisms provides a more detailed
3	understanding of these key events and processes (U.S. EPA, 2005). Moreover, toxicity
4	pathways describe the processes by which perturbation of normal biological processes
5	produce changes sufficient to lead to cell injury and subsequent events such as adverse

1	health effects (U.S. EPA, 2009f). The purpose of this section of Chapter 5 is to describe
2	the key events and toxicity pathways that contribute to health effects resulting from short-
3	term and long-term exposures to O ₃ . The extensive research carried out over several
4	decades in humans and in laboratory animals has yielded numerous studies on
5	mechanisms by which O ₃ exerts its effects. This section will discuss some of the
6	representative studies with particular emphasis on studies published since the 2006 O_3
7	AQCD and on studies in humans that inform biological mechanisms underlying
8	responses to O_3 .
9	It is well-appreciated that secondary oxidation products, which are formed as a result of
10	O_3 exposure, initiate numerous responses at the cellular, tissue and whole organ level of
11	the respiratory system. These responses include the activation of neural reflexes,
12	initiation of inflammation, alteration of epithelial barrier function, sensitization of
13	bronchial smooth muscle, modification of innate/adaptive immunity and airways
14	remodeling, as will be discussed below. These have the potential to result in effects on
15	other organ systems such as the cardiovascular, central nervous, hepatic and reproductive
16	systems or result in developmental effects. It has been proposed that lipid ozonides and
17	other secondary oxidation products, which are bioactive and cytotoxic in the respiratory
18	system, are responsible for systemic effects. However it is not known whether they gain
19	access to the vascular space (Chuang et al., 2009). Recent studies in animal models show
20	that inhalation of O_3 results in systemic oxidative stress. The following subsections
21	describe the current understanding of potential pathways and modes of action responsible
22	for the pulmonary and extrapulmonary effects of O ₃ exposure.

5.3.2 Activation of Neural Reflexes

23	Acute O ₃ exposure results in reversible effects on lung function parameters through
24	activation of neural reflexes. The involvement of bronchial C-fibers, a type of nociceptive
25	sensory nerve, has been demonstrated in dogs exposed through an endotracheal tube to
26	2-3 ppm O ₃ for 20-70 minutes (Coleridge et al., 1993; Schelegle et al., 1993). This vagal
27	afferent pathway was found to be responsible for O3-mediated rapid shallow breathing
28	and other changes in respiratory mechanics in O_3 -exposed dogs (<u>Schelegle et al., 1993</u>).
29	Ozone also triggers neural reflexes that stimulate the autonomic nervous system and alter
30	electrophysiologic responses of the heart. For example, bradycardia, altered HRV and
31	arrhythmia have been demonstrated in rodents exposed for several hours to 0.1-0.6 ppm
32	O ₃ (Hamade and Tankersley, 2009; Watkinson et al., 2001; Arito et al., 1990). Another
33	effect is hypothermia, which in rodents occurred subsequent to the activation of neural
34	reflexes involving the parasympathetic nervous system (Watkinson et al., 2001). Vagal
35	afferent pathways originating in the RT may also be responsible for O3-mediated

activation of nucleus tractus solitarius neurons that resulted in neuronal activation in stress-responsive regions of the central nervous system (CNS) (rats, 0.5-2.0 ppm O_3 for 1.5-120 hours) (Gackière et al., 2011).

4 Recent studies in animals provide new information regarding the effects of O_3 on reflex 5 responses mediated by bronchopulmonary C-fibers. In ex vivo mouse lungs, O₃ exposure 6 (30 µM solubilized) selectively activated a subset of C-fiber receptors that are TRPA1 7 ion channels (Taylor-Clark and Undem, 2010). TRPA1 ion channels are members of the 8 TRP family of ion channels, which are known to mediate the responses of sensory 9 neurons to inflammatory mediators (Caceres et al., 2009). In addition to TRPA1 ion 10 channels possibly playing a key role in O_3 -induced decrements in pulmonary function, 11 they may mediate allergic asthma (Caceres et al., 2009). Activation of TRPA1 ion 12 channels following O_3 exposure is likely initiated by secondary oxidation products such 13 as aldehydes and prostaglandins (Taylor-Clark and Undem, 2010) through covalent 14 modification of cysteine and lysine residues (Trevisani et al., 2007). Ozonation of 15 unsaturated fatty acids in the ELF was found to result in the generation of aldehydes 16 (Frampton et al., 1999) such as 4-hydroxynonenal and 4-oxononenal (Taylor-Clark et al., 17 2008; Trevisani et al., 2007). 4-oxononenal is a stronger electrophile than 18 4-hydroxynonenal and exhibits greater potency towards the TRPA1 channels (Taylor-19 Clark et al., 2008; Trevisani et al., 2007). In addition, PGE₂ is known to sensitize TRPA1 20 channels (Bang et al., 2007).

21 In humans exercising at a moderate level, the response to O_3 (500 ppb for 2 h) was 22 characterized by substernal discomfort, especially on deep inspiration, accompanied by 23 involuntary truncation of inspiration (Hazucha et al., 1989). This latter response led to 24 decreased inspiratory capacity and to decreased forced vital capacity (FVC) and forced 25 expiratory volume in one second (FEV_1), as measured by spirometry. These changes, 26 which occurred during O₃ exposure, were accompanied by decreased V_T and increased 27 respiratory frequency in human subjects. Spirometric changes in FEV_1 and FVC were not 28 due to changes in respiratory muscle strength (Hazucha et al., 1989). In addition, 29 parasympathetic involvement in the O₃-mediated decreases in lung volume was minimal 30 (Mudway and Kelly, 2000), since changes in FVC or symptoms were not modified by 31 treatment with bronchodilators such as atropine in human subjects exposed to 400 ppb O_3 32 for 2 hours while exercising at a heavy level (Beckett et al., 1985). However, the loss of 33 vital capacity was reversible with intravenous administration of the rapid-acting opioid 34 agonist, sufertanyl, in human subjects exercising at a moderate level and exposed to 35 $420 \text{ ppb } O_3$ for 2 hours, which indicated the involvement of opioid receptor-containing 36 nerve fibers and/or more central neurons (Passannante et al., 1998). The effects of 37 sufentaryl may be attributed to blocking C-fiber stimulation by O_3 since activation of 38 opioid receptors downregulated C-fiber function (Belvisi et al., 1992). Thus, nociceptive

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sensory nerves, presumably bronchial C-fibers, are responsible for O_3 -mediated responses in humans (<u>Passannante et al., 1998</u>). This vagal afferent pathway is responsible for pain-related symptoms and inhibition of maximal inspiration in humans (Hazucha et al., 1989).

5 There is some evidence that eicosanoids (see Section 5.3.3) play a role in the neural 6 reflex since cyclooxygenase inhibition with indomethacin (Alexis et al., 2000; Schelegle 7 et al., 1987) or ibuprofen, which also blocks some lipoxygenase activity (Hazucha et al., 8 1996), before exposure to O_3 significantly blunted the spirometric responses. These 9 studies involved exposures of 1-2 hours to 350-400 ppb O₃ in human subjects exercising 10 at light, moderate and heavy levels. In the latter study, ibuprofen treatment resulted in measurable decreases in BALF levels of PGE₂ and TXB₂ at 1-hour postexposure 11 12 (Hazucha et al., 1996). Although an earlier study demonstrated that PGE₂ stimulated 13 bronchial C-fibers (Coleridge et al., 1993; Coleridge et al., 1976) and suggested that 14 PGE₂ mediated O₃-induced decreases in pulmonary function, no correlation was observed 15 between the degree of ibuprofen-induced inhibition of BALF PGE₂ levels and blunting of 16 the spirometric response to O_3 (Hazucha et al., 1996). These results point to the 17 involvement of a lipoxygenase product. Further, as noted above, PGE_2 may play a role in 18 the neural reflex by sensitizing TRPA1 channels. A recent study in human subjects 19 exercising at a moderate to high level and exposed for 1 hour to 350 ppb O_3 also provided 20 evidence that arachidonic acid metabolites, as well as oxidative stress, contribute to 21 human responsiveness to O_3 (Alfaro et al., 2007).

22 In addition to the spirometric changes, mild airways obstruction occurred in human 23 subjects exercising at a moderate level during O_3 exposure (500 ppb for 2 hours) 24 (Hazucha et al., 1989). This pulmonary function decrement is generally measured as 25 specific airway resistance (sRaw) which is the product of airway resistance and thoracic 26 gas volume. In several studies involving human subjects exercising at a moderate to 27 heavy level and exposed for 1-4 hours to 200-300 ppb O₃, changes in sRaw correlated 28 with changes in inflammatory and injury endpoints measured 18-hours postexposure, but 29 did not follow the same time course or change to the same degree as spirometric changes 30 (i.e., FEV₁, FVC) measured during exposure (Balmes et al., 1996; Aris et al., 1993; 31 Schelegle et al., 1991). In addition, a small but persistent increase in airway resistance 32 associated with narrowing of small peripheral airways (measured as changes in 33 isovolumetric FEF_{25.75}) was demonstrated in O_3 -exposed human subjects (350 ppb for 34 130 minutes, moderate exercise level) (Weinmann et al., 1995c; Weinmann et al., 1995b). 35 A similar study (400 ppb O_3 for 2 hours in human subjects exercising at a heavy level) 36 found decreases in FEF₂₅₋₇₅ concomitant with increases in residual volume, which is 37 suggestive of small airways dysfunction (Kreit et al., 1989). In separate studies, a 38 statistically significant increase in residual volume (500 ppb for 2 hours) (Hazucha et al.,

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11989) and a statistically significant decrease in FEF25-75 (160 ppb for 7.6 hours)2(Horstman et al., 1995) were observed following O3 exposure in human subjects3exercising at moderate and light levels, respectively, providing further support for an4O3-induced effect on small airways.

5 Mechanisms underlying this rapid increase in airway resistance following O_3 exposure 6 are incompletely understood. Pretreatment with atropine decreased baseline sRaw and 7 prevented O₃-induced increases in sRaw in human subjects exercising at a heavy level 8 (400 ppb for 0.5 hours) (Beckett et al., 1985), indicating the involvement of muscarinic 9 cholinergic receptors of the parasympathetic nervous system. Interestingly, atropine 10 pretreatment partially blocked the decrease in FEV₁, but had no effect on the decrease in 11 FVC, breathing rate, tidal volume or respiratory symptoms (Beckett et al., 1985). Using a 12 β -adrenergic agonist, it was shown that smooth muscle contraction, not increased airway 13 mucus secretion, was responsible for O₃-induced increases in airway resistance (Beckett 14 et al., 1985). Thus, pulmonary function decrements measured as FEV_1 may reflect both 15 restrictive (such as decreased inspiratory capacity) and obstructive (such as 16 bronchoconstriction) type changes in airway responses. This is consistent with findings of 17 McDonnell et al. (1983) who observed a relatively strong correlation between sRaw and 18 FEV_1 (r = -0.31, p = 0.001) and a far weaker correlation between sRaw and FVC 19 (r = -0.16, p = 0.10) in human subjects exercising at a heavy level and exposed for 20 2.5 hours to 120-400 ppb O₃.

21 Furthermore, tachykinins may contribute to O₃-mediated increases in airway resistance. 22 In addition to stimulating CNS reflexes, bronchopulmonary C-fibers mediate local axon 23 responses by releasing neuropeptides such as substance P (SP), neurokinin (NK) A and 24 calcitonin gene-related peptide (CGRP). Tachykinins bind to NK receptors resulting in 25 responses such as bronchoconstriction. Recent studies in animals demonstrated that NK-1 26 receptor blockade had no effect on O3-stimulated physiologic responses such as VT and fB 27 in rats over the 8 hour exposure to 1 ppm O_3 (Oslund et al., 2008). However, SP and NK 28 receptors contributed to vagally-mediated bronchoconstriction in guinea pigs 3 days after 29 a single 4-hour exposure to 2 ppm O_3 (Verhein et al., 2011). In one human study in which 30 bronchial biopsies were performed and studied by immunohistochemistry, SP was 31 substantially diminished in submucosal sensory nerves 6 hours following O₃ exposure 32 (200 ppb for 2 hours, light exercise) (Krishna et al., 1997). A statistically significant 33 correlation was observed between loss of SP immunoreactivity from neurons in the 34 bronchial mucosa and changes in FEV₁ measured 1-hour postexposure (Krishna et al., 35 1997). Another study found that SP was increased in lavage fluid of human subjects 36 immediately after O₃ challenge (250 ppb for 1 hour, heavy exercise) (Hazbun et al., 37 1993). These results provide evidence that the increased airway resistance observed

1	following O_3 exposure is due to vagally-mediated responses and possibly by local axon
2	reflex responses through bronchopulmonary C-fiber-mediated release of SP.
3	A role for antioxidant defenses in modulating neural reflexes has been proposed given the
4	delay in onset of O ₃ -induced pulmonary function responses that has been noted in
5	numerous studies. Recently, this delay was characterized in terms of changes in f_B
6	(Schelegle et al., 2007). In humans exposed for 1-2 hours to 120-350 ppb O_3 while
7	exercising at a high level, no change in f_B was observed until a certain cumulative inhaled
8	dose of O_3 had been reached. Subsequently, the magnitude of the change in f_B was
9	correlated with the inhaled dose rate (Schelegle et al., 2007). These investigators
10	proposed that initial reactions of O_3 with ELF resulted in a time-dependent depletion of
11	ELF antioxidants, and that activation of neural reflexes occurred only after the
12	antioxidant defenses were overwhelmed (Schelegle et al., 2007).

5.3.3 Initiation of inflammation

13	As described previously (Section $5.2.3$), O ₃ mainly reacts with components of the ELF
14	and cellular membranes resulting in the generation of secondary oxidation products.
15	Higher concentrations of these products may directly injure RT epithelium. Subsequent
16	airways remodeling may also occur (Section 5.3.7) (Mudway and Kelly, 2000). Lower
17	concentrations of secondary oxidation products may initiate cellular responses including
18	cytokine generation, adhesion molecule expression, and modification of tight junctions
19	leading to inflammation and increased permeability across airway epithelium
20	(Section <u>5.3.4</u>) (Dahl et al., 2007; Mudway and Kelly, 2000).
21	An important hallmark of acute O_3 exposure in humans and animals is neutrophilic
22	airways inflammation. Although neutrophil influx into nasal airways has been
23	demonstrated in human subjects (400 ppb O_3 2 hours, heavy exercise) (Graham and
24	Koren, 1990), most studies of neutrophil influx have focused on the lower airways
25	(Hazucha et al., 1996; Aris et al., 1993). The time course of this response in the lower
26	airways and its resolution appears to be slower than that of the decrements in pulmonary
27	function in exercising human subjects (Hazucha et al., 1996). In several studies, airways
28	neutrophilia was observed by 1-3 hours, peaked by 6 hours and was returning to baseline
29	levels at 18-24 hours in human subjects exercising at a heavy level and exposed for
30	1-2 hours to 300-400 ppb O ₃ (Schelegle et al., 1991; Koren et al., 1989; Seltzer et al.,
31	1986). Neutrophils are thought to be injurious and a study in guinea pigs demonstrated
32	that the influx and persistence of neutrophils in airways following O ₃ exposure correlated
33	with the temporal profile of epithelial injury (0.26-1 ppm O_{3} , 72 hours) (<u>Hu et al., 1982</u>).
34	However, neutrophils have also been shown to contribute to repair of O ₃ -injured

1 2	epithelium in rats exposed for 8 hours to 1 ppm O ₃ , possibly by removing necrotic epithelial cells (<u>Mudway and Kelly, 2000</u> ; <u>Vesely et al., 1999</u>). Nonetheless, the degree
3	of airways inflammation due to O_3 is thought to have more important long-term
4	consequences than the more quickly resolving changes in pulmonary function since
5	airways inflammation is often accompanied by tissue injury (Balmes et al., 1996).
6	Ozone exposure results in alterations in other airways inflammatory cells besides
7	neutrophils, including lymphocytes, macrophages, monocytes and mast cells. Influx of
8	some of these cells accounts for the later (i.e., 18-20 hours) phase of inflammation
9	following O ₃ exposure. Numbers of lymphocytes and total cells in BALF were decreased
10	early after O ₃ exposure in human subjects exercising at a light to moderate level and
11	exposed for 2 hours to 200 ppb O_3 , which preceded the neutrophil influx (Mudway and
12	Kelly, 2000; Blomberg et al., 1999; Krishna et al., 1997). The decrease in total cells was
13	thought to reflect decreases in macrophages, although it was not clear whether the cells
14	were necrotic or whether membrane adhesive properties were altered making them more
15	difficult to obtain by lavage (Mudway and Kelly, 2000; Blomberg et al., 1999; Mudway
16	et al., 1999b; Frampton et al., 1997b; Pearson and Bhalla, 1997). A recent study in human
17	subjects exercising at a moderate level and exposed for 6.6 hours to 80 ppb O_3
18	demonstrated an increase in numbers of sputum monocytes and dendritic-like cells with
19	increased expression of innate immune surface proteins and antigen presentation markers
20	(Peden, 2011; Alexis et al., 2010) (see Section 6.2.3.1). An increase in submucosal mast
21	cells was observed 1.5 hours after a 2 hour-exposure to 200 ppb O ₃ (Blomberg et al.,
22	1999) and an increase in BAL mast cell number was observed 18 hours after a 4-hour
23	exposure to 220 ppb O_3 exposure in human subjects exercising at a moderate level
24	(Frampton et al., 1997b). Mast cells may play an important role in mediating neutrophil
25	influx since they are an important source of several pro-inflammatory cytokines and since
26	their influx preceded that of neutrophils in human subjects exercising at a moderate level
27	and exposed for 2 hours to 200 ppb O ₃ (Stenfors et al., 2002; Blomberg et al., 1999).
28	Further, a study using mast cell-deficient mice demonstrated decreased neutrophilic
29	inflammation in response to O_3 (1.75 ppm, 3 hours) compared with wild type mice
30	(Kleeberger et al., 1993). Influx of these inflammatory cell types in the lung is indicative
31	of O_3 -mediated activation of innate immunity as will be discussed in Section <u>5.3.6</u> .
32	Much is known about the cellular and molecular signals involved in inflammatory
33	responses to O_3 exposure (U.S. EPA, 2006b). Eicosanoids are one class of secondary
34	oxidation products that may be formed rapidly following O_3 exposure and that may
35	mediate inflammation. Eicosanoids are metabolites of arachidonic acid-a 20-carbon
36	PUFA—that are released from membrane phospholipids by phospholipase A2-mediated
37	catalysis. Activation of phospholipase A2 occurs by several cell signaling pathways and
38	may be triggered by O ₃ -mediated lipid peroxidation of cellular membranes (<u>Rashba-Step</u>

1	et al., 1997). Additionally, cellular phospholipases A2, C and D may be activated by lipid
2	ozonation products (Kafoury et al., 1998). While the conversion of arachidonic acid to
3	prostaglandins, leukotrienes and other eicosanoid products is generally catalyzed by
4	cyclooxygenases and lipoxygenases, non-enzymatic reactions also occur during oxidative
5	stress leading to the generation of a wide variety of eicosanoids and reactive oxygen
6	species. Further, the release of arachidonic acid from phospholipids is accompanied by
7	the formation of lysophospholipids that are precursors for platelet activating factors
8	(PAFs). Thus, formation of eicosanoids, reactive oxygen species and PAFs accompanies
9	O_3 -mediated lipid peroxidation.
10	In addition, secondary reaction products may stimulate macrophages to produce
11	cytokines such as IL-1, IL-6 and TNF- α that in turn activate IL-8 production by epithelial
12	cells. Although IL-8 has been proposed to play a role in neutrophil chemotaxis,
13	measurements of IL-8 in BALF from humans exposed to O ₃ found increases that were
14	too late to account for this effect (Mudway and Kelly, 2000). The time-course profiles of
15	PGE ₂ and IL-6 responses suggest that they may play a role in neutrophil chemotaxis in
16	humans (Mudway and Kelly, 2000). However, pretreatment with ibuprofen attenuated
17	O ₃ -induced increases in BALF PGE ₂ levels, but had no effect on neutrophilia in human
18	subjects exercising at a heavy level and exposed for 2 hour to 400 ppb O_3 (Hazucha et al.,
19	<u>1996</u>).
20	One set of studies in humans focused on the earliest phase of airways inflammation
20 21	
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21 22	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O_3 for 2 hours and bronchial biopsy tissues were obtained 1.5 and
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21 22 23 24 25 26 27	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O ₃ for 2 hours and bronchial biopsy tissues were obtained 1.5 and 6 hours after exposure (Bosson et al., 2009; Bosson et al., 2003; Stenfors et al., 2002; Blomberg et al., 1999). Results demonstrated upregulation of vascular endothelial adhesion molecules P-selectin and ICAM-1 at both 1.5 and 6 hours (Stenfors et al., 2002; Blomberg et al., 1999). Submucosal mast cell numbers were increased at 1.5 hours in the biopsy samples without an accompanying increase in neutrophil number (Blomberg et al.,
21 22 23 24 25 26 27 28	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O ₃ for 2 hours and bronchial biopsy tissues were obtained 1.5 and 6 hours after exposure (Bosson et al., 2009; Bosson et al., 2003; Stenfors et al., 2002; Blomberg et al., 1999). Results demonstrated upregulation of vascular endothelial adhesion molecules P-selectin and ICAM-1 at both 1.5 and 6 hours (Stenfors et al., 2002; Blomberg et al., 1999). Submucosal mast cell numbers were increased at 1.5 hours in the biopsy samples without an accompanying increase in neutrophil number (Blomberg et al., 1999). Pronounced neutrophil infiltration was observed at 6 hours in the bronchial
21 22 23 24 25 26 27 28 29	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O_3 for 2 hours and bronchial biopsy tissues were obtained 1.5 and 6 hours after exposure (Bosson et al., 2009; Bosson et al., 2003; Stenfors et al., 2002; Blomberg et al., 1999). Results demonstrated upregulation of vascular endothelial adhesion molecules P-selectin and ICAM-1 at both 1.5 and 6 hours (Stenfors et al., 2002; Blomberg et al., 1999). Submucosal mast cell numbers were increased at 1.5 hours in the biopsy samples without an accompanying increase in neutrophil number (Blomberg et al., 1999). Pronounced neutrophil infiltration was observed at 6 hours in the bronchial mucosa (Stenfors et al., 2002). Surprisingly, suppression of the NF- κ B and AP-1
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21 22 23 24 25 26 27 28 29 30 31	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O ₃ for 2 hours and bronchial biopsy tissues were obtained 1.5 and 6 hours after exposure (Bosson et al., 2009; Bosson et al., 2003; Stenfors et al., 2002; Blomberg et al., 1999). Results demonstrated upregulation of vascular endothelial adhesion molecules P-selectin and ICAM-1 at both 1.5 and 6 hours (Stenfors et al., 2002; Blomberg et al., 1999). Submucosal mast cell numbers were increased at 1.5 hours in the biopsy samples without an accompanying increase in neutrophil number (Blomberg et al., 1999). Pronounced neutrophil infiltration was observed at 6 hours in the bronchial mucosa (Stenfors et al., 2002). Surprisingly, suppression of the NF- κ B and AP-1 pathways at 1.5 hours and a lack of increased IL-8 at 1.5 or 6 hours in bronchial epithelium were observed (Bosson et al., 2009). The authors suggested that vascular
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21 22 23 24 25 26 27 28 29 30 31 32 33 34	One set of studies in humans focused on the earliest phase of airways inflammation (1-2 hours following exposure). Human subjects, exercising at a moderate level, were exposed to 200 ppb O ₃ for 2 hours and bronchial biopsy tissues were obtained 1.5 and 6 hours after exposure (Bosson et al., 2009; Bosson et al., 2003; Stenfors et al., 2002; Blomberg et al., 1999). Results demonstrated upregulation of vascular endothelial adhesion molecules P-selectin and ICAM-1 at both 1.5 and 6 hours (Stenfors et al., 2002; Blomberg et al., 1999). Submucosal mast cell numbers were increased at 1.5 hours in the biopsy samples without an accompanying increase in neutrophil number (Blomberg et al., 1999). Pronounced neutrophil infiltration was observed at 6 hours in the bronchial mucosa (Stenfors et al., 2002). Surprisingly, suppression of the NF-κB and AP-1 pathways at 1.5 hours and a lack of increased IL-8 at 1.5 or 6 hours in bronchial epithelium were observed (Bosson et al., 2009). The authors suggested that vascular endothelial adhesion molecules, rather than redox sensitive transcription factors, play key roles in early neutrophil recruitment in response to O ₃ .
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1	found that acute spirometric changes were not positively correlated with cellular and
2	biochemical indicators of inflammation (Aris et al., 1993; Schelegle et al., 1991).
3	However inflammation was correlated with changes in sRaw (Balmes et al., 1996). In
4	another study, pretreatment with ibuprofen had no effect on neutrophilia although it
5	blunted the spirometric response in human subjects exercising at heavy level and exposed
6	for 2 hours to 400 ppb O_3 (<u>Hazucha et al., 1996</u>). Taken together, results from these
7	studies indicate different mechanisms underlying the spirometric and inflammatory
8	responses to O_3 .

- 9 A common mechanism underlying both inflammation and impaired pulmonary function 10 was suggested by Krishna et al. (1997). This study, conducted in human subjects 11 exercising at a light level and exposed to 200 ppb O₃ for 2 hours, demonstrated a 12 correlation between loss of SP immunoreactivity from neurons in the bronchial mucosa 13 and numbers of neutrophils and epithelial cells (shed epithelial cells are an index of 14 injury) in the BALF 6-hours postexposure. Furthermore, the loss of SP immunoreactivity 15 was correlated with the observed changes in FEV₁. Another study found that SP was 16 increased in lavage fluid of exercising human subjects immediately after O₃ challenge 17 (250 ppb, 1 hour, heavy exercise) (Hazbun et al., 1993). SP is a neuropeptide released by 18 sensory nerves which mediates neurogenic edema and bronchoconstriction (Krishna et 19 al., 1997). Collectively, these findings suggest that O_3 -mediated stimulation of sensory 20 nerves that leads to activation of central and local axon reflexes is a common effector 21 pathway leading to impaired pulmonary function and inflammation.
- 22 Studies in animal models have confirmed many of these findings and provided evidence 23 for additional mechanisms involved in O_3 -induced inflammation. A study in mice (2 ppm 24 O_{3} , 3 hours) demonstrated that PAF may be important in neutrophil chemotaxis 25 (Longphre et al., 1999), while ICAM-1 and macrophage inflammatory protein-2 (MIP-2), 26 the rodent IL-8 homologue, have been implicated in a rat model (1 ppm O₃, 3 hours) 27 (Bhalla and Gupta, 2000). Key roles for CXCR2, a receptor for keratinocyte-derived 28 chemokine (KC) and MIP-2, and for IL-6 in O₃-mediated neutrophil influx were 29 demonstrated in mice (1 ppm O_3 , 3 hours) (Johnston et al., 2005a; Johnston et al., 2005b). 30 Activation of JNK and p38 pathways and cathepsin-S were also found to be important in 31 this response (3 ppm O₃, 3 hours) (Williams et al., 2009a; Williams et al., 2008a; 32 Williams et al., 2007a). Matrix metalloproteinase-9 (MMP-9) appeared to confer 33 protection against O_3 -induced airways inflammation and injury in mice (0.3 ppm O_3 , 34 6-72 hours) (Yoon et al., 2007). Interleukin-10 (IL-10) also appeared to be protective 35 since IL-10 deficient mice responded to O_3 exposure (0.3 ppm, 24-72 hours) with 36 enhanced numbers of BAL neutrophils, enhanced NF-κB activation and MIP-2 levels 37 compared with IL-10 sufficient mice (Backus et al., 2010).

- 1 In addition, lung epithelial cells may release ATP in response to O_3 exposure (Ahmad et 2 al., 2005). ATP and its metabolites (catalyzed by ecto-enzymes) can bind to cellular 3 purinergic receptors resulting in activation of cell signaling pathways (Picher et al., 4 2004). One such metabolite, adenine, is capable of undergoing oxidation leading to the 5 formation of UA which, if present in high concentrations, could activate inflammasomes 6 and result in caspase 1 activation and the maturation and secretion of IL-1 β and IL-18 7 (Dostert et al., 2008). A recent study in human subjects exercising at a moderate level and 8 exposed for 2 hours to 400 ppb O_3 demonstrated a correlation between ATP metabolites 9 and inflammatory markers (Esther et al., 2011), which provides some support for this 10 mechanism.
- 11 Several recent studies have focused on the role of Toll-like receptor (TLR) and its related 12 adaptor protein MyD88 in mediating O₃-induced neutrophilia. Hollingsworth et al. (2004) 13 demonstrated airways neutrophilia that was TLR4-independent following acute (2 ppm, 14 3 hours) and subchronic (0.3 ppm, 72 hours) O_3 exposure in a mouse model. However, 15 Williams et al. (2007b) found that MyD88 was important in mediating O₃-induced 16 neutrophilia in mice (3 ppm, 3 hours), with TLR4 and TLR2 contributing to the speed of 17 the response. Moreover, MyD88, TLR2 and TLR4 contributed to inflammatory gene 18 expression in this model and O₃ upregulated MyD88, TLR4 and TLR4 gene expression 19 (Williams et al., 2007a). Neutrophilic inflammation was also found to be partially 20 dependent on MyD88 in mice exposed to 1 ppm O_3 for 3 hours (Li et al., 2011).
- 21 Hyaluronan was found to mediate a later phase (24 hours) of O_3 -induced inflammation in 22 mice (Garantziotis et al., 2010; Garantziotis et al., 2009). Hyaluronan is an extracellular 23 matrix component that is normally found in the ELF as a large polymer. Exposure to 24 2 ppm O_3 for 3 hours resulted in elevated levels of soluble low molecular weight 25 hyaluronan in the BALF 24-hours postexposure (Garantziotis et al., 2010; Garantziotis et 26 al., 2009). Similar results were found in response to 3 hour exposure to 1 ppm O_3 (Li et 27 al., 2011). Ozone may have caused the depolymerization of hyaluronan to soluble 28 fragments that are known to be endogenous ligands of the CD44 receptor and TLR4 in 29 the macrophage (Jiang et al., 2005). Binding of hyaluronan fragments to the CD44 30 receptor activates hyaluronan clearance, while binding to TLR4 results in signaling 31 through MyD88 to produce chemokines that stimulate the influx of inflammatory cells 32 (Jiang et al., 2005). Activation of NF- κ B occurred in both airway epithelia and alveolar 33 macrophages 24-hours postexposure to O₃. Increases in BALF pro-inflammatory factors 34 KC, IL-1 β , MCP-1, TNF- α and IL-6 observed 24 hours following O₃ exposure were 35 found to be partially dependent on TLR4 (Garantziotis et al., 2010) while increases in 36 BAL inflammatory cells, which consisted mainly of macrophages, were dependent on 37 CD44 (Garantziotis et al., 2009). BAL inflammatory cells number and injury markers

1following O3 exposure were similar in wild-type and TLR4-deficient animals2(Garantziotis et al., 2010).

- 3 Since exposure to O_3 leads to airways inflammation characterized by neutrophilia, and 4 since neutrophil-derived oxidants often consume ELF antioxidants, concentrations of 5 ELF antioxidants have been examined during airways neutrophilia (Long et al., 2001; 6 Gunnison and Hatch, 1999; Mudway et al., 1999b). In human subjects exercising at a 7 moderate level and exposed to 200 ppb O₃ for 2 hours, UA, GSH and α-TOH levels 8 remained unchanged in BALF 6-hours postexposure while AH2 was decreased 9 significantly in both BALF and plasma (Mudway et al., 1999b). A second study 10 involving the same protocol reported a loss of AH2 from bronchial wash fluid and BALF, representing proximal and distal airway ELF respectively, as well as an increase in 11 oxidized GSH in both compartments (Mudway et al., 2001). No change was observed in 12 13 ELF UA levels in response to O₃ (Mudway et al., 2001). Further, O₃ exposure (0.8 ppm, 14 4 hours) in female rats resulted in a 50% decrease in BALF AH2 immediately 15 postexposure (Gunnison and Hatch, 1999). These studies suggested a role for AH2 and 16 GSH in protecting against the oxidative stress associated with inflammation.
- 17 The relationship between inflammation, antioxidant status and O₃ dose has also been 18 investigated. The degree of inflammation in rats has been correlated with ¹⁸O-labeled O_3 dose markers in the lower lung. In female rats exposed to 0.8 ppm O₃ for 4 hours, BAL 19 neutrophil number and ¹⁸O reaction product were directly proportional (Gunnison and 20 21 Hatch, 1999). Kari et al. (1997) observed that a 3-week caloric restriction (75%) in rats 22 abrogated the toxicity of O_3 (2 ppm, 2 hours), measured as BALF increases in protein, fibronectin and neutrophils, that was seen in normally fed rats. Accompanying this 23 resistance to O_3 toxicity was a reduction (30%) in the accumulation of ¹⁸O reaction 24 25 product in the lungs. These investigations also demonstrated an inverse relationship 26 between AH2 levels and O_3 dose and provided evidence for AH2 playing a protective 27 role following O₃ exposure in these studies. Pregnant and lactating rats had lower AH2 content in BALF and exhibited a greater increase in accumulation of ¹⁸O reaction 28 29 products compared with pre-pregnant rats in response to O_3 exposure (Gunnison and 30 Hatch, 1999). In the calorie restricted model, a 30% higher basal BALF AH2 31 concentration and a rapid accumulation of AH2 into the lungs to levels 60% above 32 normal occurred as result of O₃ exposure (Kari et al., 1997). However, this relationship 33 between AH2 levels and O₃ dose did not hold up in every study. Aging rats (9 and 34 24 months old) had 49% and 64% lower AH2 in lung tissue compared with month-old rats but the aging-induced AH2 loss did not increase the accumulation of ¹⁸O reaction 35 36 products following O₃ exposure (0.4-0.8 ppm, 2-6 hours) (Vincent et al., 1996b).

1	A few studies have examined the dose- or concentration-responsiveness of airways
2	neutrophilia in O ₃ -exposed humans (<u>Holz et al., 1999</u> ; <u>Devlin et al., 1991</u>). No
3	concentration-responsiveness was observed in healthy human subjects exposed for 1 hour
4	to 125-250 ppb O ₃ while exercising at a light level and a statistically significant increase
5	in sputum neutrophilia was observed only at the higher concentration (Holz et al., 1999).
6	However, concentration-dependent and statistically significant increases in BAL
7	neutrophils and the inflammatory mediator IL-6 were reported following exposure to 80
8	and 100 ppb O_3 for 6.6 hours in human subjects exercising at a moderate level (<u>Devlin et</u>
9	<u>al., 1991</u>). Additional evidence is provided by a meta-analysis of the O_3
10	dose-inflammatory response in controlled human exposure studies involving exposure to
11	80-600 ppb O_3 for 60-396 minutes and exercise levels ranging from light to heavy
12	(Mudway and Kelly, 2004b). Results demonstrated a linear relationship between inhaled
13	O_3 dose (determined as the product of concentration, ventilation and time) and BAL
14	neutrophils at 0-6 hours and 18-24 hours following O3 exposure (Mudway and Kelly,
15	<u>2004b</u>).

5.3.4 Alteration of Epithelial Barrier Function

- 16 Following O₃ exposure, injury and inflammation can lead to altered airway barrier 17 function. Histologic analysis has demonstrated damage to tight junctions between 18 epithelial cells, suggesting an increase in epithelial permeability. In addition, the presence 19 of shed epithelial cells in the BALF and increased epithelial permeability, which is 20 measured as the flux of small solutes, have been observed and are indicative of epithelial 21 injury. This could potentially lead to the loss of ELF solutes that could diffuse down their 22 concentration gradient from the lung to the blood. Increases in vascular permeability, as 23 measured by BALF protein and albumin, have also been demonstrated (Costa et al., 24 1985; Hu et al., 1982).
- 25 An early study in sheep measured changes in airway permeability as the flux of inhaled 26 radiolabeled histamine into the plasma (Abraham et al., 1984). Exposure of sheep to 27 0.5 ppm O_3 for 2 hours via an endotracheal tube resulted in an increased rate of histamine 28 appearance in the plasma at 1 day postexposure. Subsequently, numerous studies have measured epithelial permeability as the flux of the small solute ^{99m}Tc-DTPA that was 29 30 introduced into the air spaces in different regions of the RT. Increased pulmonary epithelial permeability, measured as the clearance of ^{99m}Tc-DTPA from lung to blood, 31 32 was demonstrated in humans 1-2 hours following a 2-hour exposure to 400 ppb O₃ while 33 exercising at a heavy level (Kehrl et al., 1987). Another study in human subjects found 34 increased epithelial permeability 19-hours postexposure to 240 ppb O_3 for 130 minutes 35 while exercising at moderate level (Foster and Stetkiewicz, 1996). Increased bronchial

1 2	permeability was also observed in dogs 1-day postexposure (0.4 ppm O_3 by endotracheal tube for 6 hours) and did not resolve for several days (Foster and Freed, 1999).
3	A role for tachykinins in mediating airway epithelial injury and decreased barrier
4	function has been suggested. Nishiyama et al. (1998) demonstrated that capsaicin, which
5	depletes nerve fibers of substance P, blocked the O3-induced increase in permeability of
6	guinea pig tracheal mucosa (0.5-3 ppm O_3 , 0.5 hours). Pretreatment with propranolol or
7	atropine failed to inhibit this response, suggesting that adrenergic and cholinergic
8	pathways were not involved. In another study, tachykinins working through NK-1 and
9	CGRP receptors were found to contribute to airway epithelial injury in O ₃ -exposed rats
10	(1 ppm, 8 hours) (<u>Oslund et al., 2009</u> , <u>2008</u>).
11	<u>Kleeberger et al. (2000</u>) evaluated genetic susceptibility to O_3 -induced altered barrier
12	function in recombinant inbred strains of mice. Lung hyperpermeability, measured as
13	BALF protein, was evaluated 72 hours after exposure to 0.3 ppm O_3 and found to be
14	associated with a functioning Tlr4 gene. This study concluded that Tlr4 was a strong
15	candidate gene for susceptibility to hyperpermeability in response to O_3 (Kleeberger et
16	al., 2000). A subsequent study by these same investigators found that Tlr4 modulated
17	mRNA levels of the Nos2 genes and suggested that the protein product of Nos2, iNOS,
18	plays an important role in O_2 -induced lung hyperpermeability (0.3 ppm, 72 hours)
19	(Kleeberger et al., 2001). More recently, HSP70 was identified as part of the TLR4
20	signaling pathway (0.3 ppm, 6-72 hours) (<u>Bauer et al., 2011</u>).
21	Antioxidants have been shown to confer resistance to O ₃ -induced injury. In a recent
22	study, lung hyperpermeability in response to O_3 (0.3 ppm, 48 hours) was unexpectedly
23	reduced in mice deficient in the glutamate-cysteine ligase modifier subunit gene
24	compared with sufficient mice (Johansson et al., 2010). Since the lungs of these mice
25	exhibited 70% glutathione depletion, protection against O3-induced injury was
26	unexpected (Johansson et al., 2010). However it was found that several other antioxidant
27	defenses, including metallothionein, were upregulated in response to O_3 to a greater
28	degree in the glutathione-deficient mice compared with sufficient mice (Johansson et al.,
29	<u>2010</u>). The authors suggested that resistance to O_3 -induced lung injury was due to
30	compensatory augmentation of antioxidant defenses (Johansson et al., 2010). Antioxidant
31	effects have also been attributed to Clara cell secretory protein (CCSP) and surfactant
32	protein A (SP-A). CCSP was found to modulate the susceptibility of airway epithelium to
33	injury in mice exposed to O_3 (0.2 or 1 ppm for 8 hours) by an unknown mechanism
34	(<u>Plopper et al., 2006</u>). SP-A appeared to confer protection against O_3 -induced airways
35	inflammation and injury in mice (2 ppm, 3 hours) (<u>Haque et al., 2007</u>).
36	Increased epithelial permeability has been proposed to play a role in allergic sensitization
37	(Matsumura, 1970), in activation of neural reflexes and in stimulation of smooth muscle

1receptors (Dimeo et al., 1981). Abraham et al. (1984) reported a correlation between2airway permeability and airways hyperresponsiveness (AHR) in O3-exposed sheep.3However a recent study in human subjects exposed to 220 ppb O3 for 135 minutes while4exercising at a light to moderate level did not find a relationship between O3-induced5changes in airway permeability and AHR (Que et al., 2011).

5.3.5 Sensitization of Bronchial Smooth Muscle

- 6 Bronchial reactivity is generally determined in terms of a response to a challenge agent. 7 Non-specific bronchial reactivity in humans is assessed by measuring the effect of 8 inhaling increasing concentrations of a bronchoconstrictive drug on lung mechanics 9 $(sRaw \text{ or } FEV_1)$. Methacholine is most commonly employed but histamine and other 10 agents are also used. Specific bronchial reactivity is assessed by measuring effects in 11 response to an inhaled allergen in individuals (or animals) already sensitized to that 12 allergen. An increase in sRaw in response to non-specific or specific challenge agents 13 indicates AHR.
- 14In addition to causing mild airways obstruction as discussed above, acute O_3 exposure15results in reversible increases in bronchial reactivity by mechanisms that are not well16understood. In one study, bronchial reactivity of healthy subjects was significantly17increased 19-hours postexposure to O_3 (120-240 ppb O_3 for 2 hours with moderate18exercise) (Foster et al., 2000). These effects may be more considerable in human subjects19with already compromised airways (Section 5.4.2.2).
- 20 Ozone may sensitize bronchial smooth muscle to stimulation through an exposure-related 21 effect on smooth muscle or through effects on the sensory nerves in the epithelium or on 22 the motor nerves innervating the smooth muscle (O'Byrne et al., 1984; O'Byrne et al., 23 1983; Holtzman et al., 1979). It is also recognized that increased bronchial reactivity can 24 be both a rapidly occurring and a persistent response to O₃ (Foster and Freed, 1999). 25 Tachykinins and secondary oxidation products of O_3 have been proposed as mediators of 26 the early response and inflammation-derived products have been proposed as mediators 27 of the later response (Foster and Freed, 1999). Furthermore, bronchial reactivity may be 28 increased as a result of O₃-mediated generation of ROS.
- 29Ozone-induced increases in epithelial permeability, which could improve access of30agonist to smooth muscle receptors, may be one mechanism of sensitization through a31direct effect on bronchial smooth muscle (Holtzman et al., 1979). As noted above, a32correlation between airway permeability and AHR has been reported in O3-exposed sheep33(Abraham et al., 1984) but not in O3-exposed human subjects (Que et al., 2011).

1	Neurally-mediated sensitization has been demonstrated. In human subjects exposed for
2	2 hours to 600 ppb O_3 while exercising at a light level, pretreatment with atropine
3	inhibited O_3 -induced AHR, suggesting the involvement of cholinergic postganglionic
4	pathways (Holtzman et al., 1979). Animal studies have demonstrated that O ₃ -induced
5	AHR involved vagally-mediated responses (rabbits, 0.2 ppm O ₃ , 72 hours) (Freed et al.,
6	<u>1996</u>) and local axon reflex responses through bronchopulmonary C-fiber-mediated
7	release of SP (guinea pigs, 0.8 ppm O_3 , 2 hours) (Joad et al., 1996). Further, pretreatment
8	with capsaicin to deplete nerve fibers of SP blocked O ₃ -mediated AHR (guinea pigs,
9	$1-2 \text{ ppm O}_3$, 2-2.25 hours) (Tepper et al., 1993). Other investigators demonstrated that SP
10	released from airway nociceptive neurons in ferrets contributed to O ₃ -induced AHR
11	(2 ppm O_3 , 3 hours) (<u>Wu et al., 2008c; Wu et al., 2003</u>).
12	Some evidence suggests the involvement of arachidonic acid metabolites and neutrophils
13	in mediating O_3 -induced AHR (<u>Seltzer et al., 1986</u> ; <u>Fabbri et al., 1985</u>). Increased BAL
14	neutrophils and cyclooxygenase products were found in one study demonstrating AHR in
15	human subjects exercising at a heavy level immediately postexposure to 600 ppb O_3 for
16	2 hours (<u>Seltzer et al., 1986</u>). Another study found that ibuprofen pretreatment had no
17	effect on AHR in human subjects exercising at a heavy level following exposure to
18	400 ppb O_3 for 2 hours, although spirometric responses were blunted (<u>Hazucha et al.</u> ,
19	<u>1996</u>). This study measured arachidonic acid metabolites and provided evidence that that
20	the arachidonic acid metabolites whose generation was blocked by ibuprofen,
21	(i.e., prostaglandins, thromboxanes and some leukotrienes) did not play a role in AHR.
22	Experiments in dogs exposed for 2 hours to 2.1 ppm O ₃ demonstrated a close correlation
23	between O ₃ -induced AHR and airways neutrophilic inflammation measured in tissue
24	biopsies (Holtzman et al., 1983). Furthermore, the increased AHR observed in dogs
25	following O ₃ exposure (3 ppm, 2 hours) was inhibited by neutrophil depletion (<u>O'Byrne</u>
26	et al., 1983) and by pre-treatment with inhibitors of arachidonic acid metabolism. In one
27	of these studies, indomethacin pre-treatment did not prevent airways neutrophilia in
28	response to O_3 (3 ppm, 2 hours) providing evidence that the subset of arachidonic acid
29	metabolites whose generation was inhibitable by the cyclooxygenase inhibitor
30	indomethacin (i.e., prostaglandins and thromboxanes) was not responsible for neutrophil
31	influx (O'Byrne et al., 1984). It should be noted that these studies did not measure
32	whether the degree to which the inhibitor blocked arachidonic acid metabolism and thus
33	their results should be interpreted with caution. Taken together, these findings suggest that
34	arachidonic acid metabolites may be involved in the AHR response following O_3
35	exposure in dogs. Studies probing the role of neutrophils in mediating the AHR response
36	have provided inconsistent results (Al-Hegelan et al., 2011).
37	Evidence for cytokine and chemokine involvement in the AHR response to O_3 has been
38	described. Some studies have suggested a role for TNF- α (mice, 0.5 and 2 ppm O ₃ ,

1	3 hours) (Cho et al., 2001; Shore et al., 2001) and IL-1 (mice and ferrets, 2 ppm O ₃ ,
2	3 hours) (Wu et al., 2008c; Park et al., 2004). The latter study found that SP expression in
3	airway neurons was upregulated by IL-1 that was released in response to O_3 . Other
4	studies in mice have demonstrated a key role for CXCR2, the chemokine receptor for the
5	neutrophil chemokines KC and MIP-2, but not for IL-6 in O ₃ -mediated AHR (1 ppm O ₃ ,
6	3 hours) (Johnston et al., 2005a; Johnston et al., 2005b). In contrast, CXCR2 and IL-6
7	were both required for neutrophil influx in this model (Johnston et al., 2005a; Johnston et
8	al., 2005b), as discussed above. Williams et al. (2008b) demonstrated that the Th2
9	cytokine IL-13 contributed to AHR, as well as to airways neutrophilia, in mice (3 ppm
10	O ₃ , 3 hours).

- 11Other studies have focused on the role of TLR4. Hollingsworth et al. (2004) measured12AHR, as well as airways neutrophilia, in mice 6 and 24 hours following acute (2 ppm O313for 3 hours) and subchronic (0.3 ppm for 3 days) exposure to O3. TLR4 is a key14component of the innate immune system and is responsible for the immediate
- 15 inflammatory response seen following challenge with endotoxin and other pathogen-16 associated substances. In this study, a functioning TLR4 was required for the full AHR 17 response following O₃ exposure but not for airways neutrophilia (Hollingsworth et al., 18 2004). These findings are complemented by an earlier study demonstrating that O_3 effects 19 on lung hyperpermeability required a functioning TLR4 (mice, 0.3 ppm O₃, 72 hours) 20 (Kleeberger et al., 2000). Williams et al. (2007b) found that TLR2, TLR4 and the TLR 21 adaptor protein MyD88 contributed to AHR in mice (3 ppm O₃, 3 hours). Ozone was also 22 found to upregulate MyD88, TLR4 and TLR4 gene expression in this model (Williams et 23 al., 2007b). Furthermore, a recent study reported O_3 -induced AHR that required TLR4
 - and MyD88 in mice exposed to 1 ppm O_3 for 3 hours (Li et al., 2011).
- 25 A newly recognized mechanistic basis for O₃-induced AHR is provided by studies 26 focusing on the role of hyaluronan following O_3 exposure in mice (Garantziotis et al., 27 2010; Garantziotis et al., 2009). Hyaluronan is an extracellular matrix component that is 28 normally found in the ELF as a large polymer. Briefly, TLR4 and CD44 were found to 29 mediate AHR in response to O_3 and hyaluronan. Exposure to 2 ppm O_3 for 3 hours 30 resulted in enhanced AHR and elevated levels of soluble low molecular weight 31 hyaluronan in the BALF 24-hours postexposure (Garantziotis et al., 2010; Garantziotis et 32 al., 2009). Ozone may have caused the depolymerization of hydronan to soluble 33 fragments that are known to be endogenous ligands of the CD44 receptor and TLR4 in 34 the macrophage (Jiang et al., 2005). In the two recent studies, O₃-induced AHR was 35 attenuated in CD44 and TLR4-deficient mice (Garantziotis et al., 2010; Garantziotis et 36 al., 2009). Hyaluronan fragment-mediated stimulation of AHR was found to require 37 functioning CD44 receptor and TLR4 (Garantziotis et al., 2010; Garantziotis et al., 2009). 38 In contrast, high-molecular-weight hyaluronan blocked AHR in response to O₃

1	(Garantziotis et al., 2009). In another study high-molecular-weight hyaluronan enhanced
2	repair of epithelial injury (Jiang et al., 2005). These studies provide a link between innate
3	immunity and the development of AHR following O_3 exposure, and indicate a role for
4	TLR4 in increasing airways responsiveness. While TLR4-dependent responses usually
5	involve activation of NF- κ B and the upregulation of proinflammatory factors, the precise
6	mechanisms leading to AHR are unknown (<u>Al-Hegelan et al., 2011</u>).
7	In guinea pigs, AHR was found to be mediated by different pathways at 1- and 3-days
8	postexposure to a single exposure of O_3 (2 ppm for 4 hours) (Verhein et al., 2011; Yost et
9	al., 2005). At 1 day, AHR was due to activation of airway parasympathetic nerves rather
10	than to an exposure-related effect on smooth muscle (<u>Yost et al., 2005</u>). This effect
11	occurred as a result of O ₃ -stimulated release of major basic protein from eosinophils
12	(<u>Yost et al., 2005</u>). Major basic protein is known to block inhibitory M2 muscarinic
13	receptors that normally dampen acetylcholine release from parasympathetic nerves (<u>Yost</u>
14	et al., 2005). The resulting increase in acetylcholine release caused an increase in smooth
15	muscle contraction following O_3 exposure (<u>Yost et al., 2005</u>). Eosinophils played a
16	different role 3-days postexposure to O_3 in guinea pigs (<u>Yost et al., 2005</u>). Ozone-
17	mediated influx of eosinophils into lung airways resulted in a different population of cells
18	present 3-days postexposure compared to those present at 1 day (Yost et al., 2005). At
19	this time point, eosinophil-derived major basic protein increased smooth muscle
20	responsiveness to acetylcholine which also contributed to AHR (Yost et al., 2005).
21	However, the major effect of eosinophils was to protect against vagal hyperreactivity
22	(Yost et al., 2005). The authors suggested that these beneficial effects were due to the
23	production of nerve growth factor (Yost et al., 2005). Further work by these investigators
24	demonstrated a key role for IL-1 β in mediating AHR 3-days postexposure to O ₃ (Verhein
25	et al., 2011). In this study, IL-1 β increased nerve growth factor and SP that acted through
26	the NK1 receptor to cause vagally-mediated bronchoconstriction (Verhein et al., 2011).
27	The mechanism by which SP caused acetylcholine release from parasympathetic nerves
28	following O_3 exposure was not determined (<u>Verhein et al., 2011</u>). Taken together, the
29	above study results indicate that mechanisms involved in O3-mediated AHR can vary
30	over time postexposure and that eosinophils and SP can play a role. Results of this animal
31	model may provide some insight into allergic airways disease in humans that is
32	characterized by eosinophilia (Section <u>5.4.2.2</u>).

5.3.6 Modification of Innate/Adaptive Immune System Responses

33	Host defense depends on effective barrier function and on innate immunity and adaptive
34	immunity (<u>Al-Hegelan et al., 2011</u>). The effects of O_3 on barrier function in the airways
35	was discussed above (Section $5.3.4$). This section focuses on the mechanisms by which

1 O_3 impacts innate and adaptive immunity. Both tissue damage and foreign pathogens are 2 triggers for the activation of the innate immune system. This results in the influx of 3 inflammatory cells such as neutrophils, mast cells, basophils, eosinophils, monocytes and 4 dendritic cells and the generation of cytokines such as TNF- α , IL-1, IL-6, KC and IL-17. 5 Further, innate immunity encompasses the actions of complement and collections, and 6 the phagocytic functions of macrophages, neutrophils and dendritic cells. Airway 7 epithelium also contributes to innate immune responses. Innate immunity is highly 8 dependent on cell signaling networks involving TLR4. Adaptive immunity provides 9 immunologic memory through the actions of B and T-cells. Important links between the 10 two systems are provided by dendritic cells and antigen presentation. Recent studies 11 demonstrate that exposure to O_3 modifies cells and processes which are required for 12 innate immunity, contributes to innate-adaptive immune system interaction and primes 13 pulmonary immune responses to endotoxin.

14 Ozone exposure of human subjects resulted in recruitment of activated innate immune 15 cells to the airways. Healthy individuals were exposed to 80 ppb O_3 for 6.6 hours while 16 exercising at a moderate level and airways inflammation was characterized in induced 17 sputum 18-hours postexposure (Alexis et al., 2010). Previous studies demonstrated that 18 induced sputum contains liquid and cellular constituents of the ELF from central 19 conducting airways (Alexis et al., 2001b) and also identified these airways as a site of 20 preferential O₃ absorption during exercise (Hu et al., 1994). Ozone exposure resulted in 21 increased numbers of neutrophils, airway monocytes and dendritic-like cells in sputum 22 (Alexis et al., 2010). In addition, increased expression of cell surface markers 23 characteristic of innate immunity and antigen presentation (i.e., CD-14 and HLA-DR) 24 was demonstrated on airway monocytes (Alexis et al., 2010). Enhanced antigen 25 presentation contributes to exaggerated T-cell responses and promotes Th2 inflammation 26 and an allergic phenotype (Lay et al., 2007). Upregulation of pro-inflammatory cytokines 27 was also demonstrated in sputum of O3-exposed subjects (Alexis et al., 2010). One of 28 these cytokines, IL-12p70, correlated with numbers of dendritic-like cells in the sputum, 29 and is an indicator of dendritic cell activation (Alexis et al., 2010). These authors have 30 previously reported that exposure of human subjects exercising at a light to moderate 31 level to 400 ppb O_3 for 2 hours resulted in activation of monocytes and macrophages 32 (Lay et al., 2007), which could play a role in exacerbating existing asthma by activating 33 allergen-specific memory T-cells. The current study confirms these findings and extends 34 them by suggesting a potential mechanism whereby O_3 -activated dendritic cells could 35 stimulate naïve T-cells to promote the development of asthma (Alexis et al., 2010). A 36 companion study by these same investigators (described in detail in Section 5.4.2.1) 37 provides evidence of dendritic cell activation, measured as increased expression of HLA-38 DR, in a subset of the human subjects (GSTM1 null) exposed to 400 ppb O_3 for 2 hours 39 while exercising at a light to moderate level (Alexis et al., 2009). Since dendritic cells are a link between innate and adaptive immunity, these studies provide evidence for an O₃-mediated interaction between the innate and adaptive immune systems.

- 3 Another recent study linked O₃-mediated activation of the innate immune system to the 4 development of non-specific AHR in a mouse model (Pichavant et al., 2008). Repeated 5 exposure to 1 ppm O₃ for 3 hours (3 days over a 5 day period) induced non-specific AHR 6 measured 24 hours following the last exposure (Pichavant et al., 2008). This response 7 was found to require NKT-cells, which are effector lymphocytes of innate immunity, as 8 well as IL-17 and airways neutrophilia (Pichavant et al., 2008). Since glycolipids such as 9 galactosyl ceramide are ligands for the invariant CD1 receptor on NKT-cells and serve as 10 endogenous activators of NKT-cells, a role for O₃-oxidized lipids in activating NKT-cells was proposed (Pichavant et al., 2008). The authors contrasted this innate immunity 11 12 pathway with that of allergen-provoked specific AHR which involves adaptive immunity, 13 the cytokines IL-4, IL -13, IL-17, and airways eosinophilia (Pichavant et al., 2008). 14 Interestingly, NKT-cells were required for both the specific AHR provoked by allergen 15 and the non-specific AHR provoked by O₃ (Pichavant et al., 2008). Different cytokine 16 profiles of the NKT-cells from allergen and O₃-exposed mice were proposed to account 17 for the different pathways (Pichavant et al., 2008). More recently, NKT-cells have been 18 found to function in both innate and adaptive immunity (Vivier et al., 2011).
- 19 An interaction between allergen and O_3 in the induction of nonspecific AHR was shown 20 in another animal study (Larsen et al., 2010). Mice were sensitized with the aerosolized 21 allergen OVA on 10 consecutive days followed by exposure to O_3 (0.1-0.5 ppm for 22 3 hours) (Larsen et al., 2010). While allergen sensitization alone did not alter airways
- responsiveness to a nonspecific challenge, O_3 exposure of sensitized mice resulted in 24 nonspecific AHR at 6- and 24-hours postexposure (Larsen et al., 2010). The effects of O_3 25 on AHR were independent of airways eosinophilia and neutrophilia (Larsen et al., 2010). 26 However, OVA pretreatment led to goblet cell metaplasia which was enhanced by O_3 27 exposure (Larsen et al., 2010). It should be noted that OVA sensitization using only 28 aerosolized antigen in this study is less common than the usual procedure for OVA 29 sensitization achieved by one or more initial systemic injections of OVA and adjuvant 30 followed by repeated inhalation exposure to OVA. This study also points to an interaction 31 between innate and adaptive immune systems in the development of the AHR response.
- 32 Furthermore, O₃ was found to act as an adjuvant for allergic sensitization (Hollingsworth 33 et al., 2010). Oropharyngeal aspiration of OVA on day 0 and day 6 failed to lead to 34 allergic sensitization unless mice were first exposed to 1 ppm O₃ for 2 hours 35 (Hollingsworth et al., 2010). The O3-mediated response involved Th2 (IL-4, IL-5 and 36 IL-9) and Th17 cytokines (IL-17) and was dependent on a functioning TLR4 37 (Hollingsworth et al., 2010). Ozone exposure also activated OVA-bearing dendritic cells

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in the thoracic lymph nodes, as measured by the presence of the CD86 surface marker, which suggests naïve T-cell stimulation and the involvement of Th2 pathways (Hollingsworth et al., 2010). Thus the adjuvant effects of O_3 may be due to activation of both innate and adaptive immunity.

5 Priming of the innate immune system by O_3 was reported by Hollingsworth et al. (2007). 6 In this study, exposure of mice to 2 ppm O_3 for 3 hours led to nonspecific AHR at 24-7 and 48-hours postexposure, an effect which subsided by 72 hours (Hollingsworth et al., 8 2007). However, in mice treated with aerosolized endotoxin immediately following O_3 9 exposure, AHR was greatly enhanced at 48-and 72-hours postexposure (Hollingsworth et 10 al., 2007). In addition, O_3 pre-exposure was found to reduce the number of inflammatory 11 cells in the BALF, to increase cytokine production and total protein in the BALF and to 12 increase systemic IL-6 following exposure to endotoxin (Hollingsworth et al., 2007). 13 Furthermore, O₃ stimulated the apoptosis of alveolar macrophages 24-hours 14 postexposure, an effect which was greatly enhanced by endotoxin treatment. Apoptosis of 15 circulating blood monocytes was also observed in response to the combined exposures 16 (Hollingsworth et al., 2007). Ozone pre-exposure enhanced the response of lung 17 macrophages to endotoxin (Hollingsworth et al., 2007). Taken together, these findings 18 demonstrated that O_3 exposure increased innate immune responsiveness to endotoxin. 19 The authors attributed these effects to the increased surface expression of TLR4 and 20 increased signaling in macrophages observed in the study (Hollingsworth et al., 2007). It 21 was proposed that the resulting decrease in airway inflammatory cells could account for 22 O₃-mediated decreased clearance of bacterial pathogens observed in numerous animal 23 models (Hollingsworth et al., 2007).

24 More recently, these authors demonstrated that hyaluronan contributed to the O₃-primed 25 response to endotoxin (Li et al., 2010). In this study, exposure of mice to 1 ppm O_3 for 26 3 hours resulted in enhanced responses to endotoxin, which was mimicked by 27 intratracheal instillation of hyaluronan fragments (Li et al., 2010). Hyaluronan, like O₃, 28 was also found to induce TLR4 receptor peripheralization in the macrophage membrane 29 (Li et al., 2010; Hollingsworth et al., 2007), an effect which is associated with enhanced 30 responses to endotoxin. This study and previous ones by the same investigators showed 31 elevation of BALF hyaluronan in response to O_3 exposure (Garantziotis et al., 2010; Li et 32 al., 2010; Garantziotis et al., 2009), providing evidence that the effects of O_3 on innate 33 immunity are at least in part mediated by hyaluronan fragments. The authors note that 34 excessive TLR4 signaling can lead to lung injury and suggest that O₃ may be responsible 35 for an exaggerated innate immune response which may underlie lung injury and 36 decreased host defense (Li et al., 2010).

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1	Activation or upregulation of the immune system has not been reported in all studies.
2	Impaired antigen-specific immunity was demonstrated following subacute O ₃ exposure
3	(0.6 ppm, 10 h/day for 15 days) in mice (Feng et al., 2006). Specifically, O ₃ exposure
4	altered the lymphocyte subset and cytokine profile and impacted thymocyte early
5	development leading to immune dysfunction. Further, recent studies demonstrated SP-A
6	oxidation in mice exposed for 3-6 hours to 2 ppm O ₃ . SP-A is an important innate
7	immune protein which plays a number of roles in host defense including acting as
8	opsonin for the recognition of some pathogens (Haque et al., 2009). These investigations
9	found that O ₃ -mediated carbonylation of purified SP-A was associated with impaired
10	macrophage phagocytosis in vitro (Mikerov et al., 2008c). In addition, O ₃ exposure
11	(2 ppm for 3 hours) in mice was found to increase susceptibility to pneumonia infection
12	in mice through an impairment of SP-A dependent phagocytosis (Mikerov et al., 2008b;
13	<u>Mikerov et al., 2008a</u>). Furthermore, early life exposure to O_3 in infant monkeys followed
14	by a recovery period led to hyporesponsiveness to endotoxin (Maniar-Hew et al., 2011),
15	as discussed below and in Section $5.4.2.4$ and Section $7.2.3.1$.

16 Taken together, results of recent studies provide evidence that O₃ alters host
17 immunologic response and leads to immune system dysfunction through its effects on
18 innate and adaptive immunity.

5.3.7 Airways Remodeling

19	The nasal airways, conducting airways and distal airways (i.e., respiratory bronchioles or
20	CAR depending on the species) have all been identified as sites of O3-mediated injury
21	and inflammation (Mudway and Kelly, 2000). At all levels of the RT, loss of sensitive
22	epithelial cells, degranulation of secretory cells, proliferation of resistant epithelial cells
23	and neutrophilic influx have been observed as a result of O_3 exposure (Mudway and
24	Kelly, 2000; Cho et al., 1999). An important study (Plopper et al., 1998) conducted in
25	adult rhesus monkeys (0.4 and 1.0 ppm O_3 for 2 hours at rest) found that 1 ppm O_3
26	resulted in the greatest epithelial injury in the respiratory bronchioles immediately
27	postexposure although injury was observed at all of the RT sites studied except for the
28	lung parenchyma. Exposure to 0.4 ppm O_3 resulted in epithelial injury only in the
29	respiratory bronchioles. Initial cellular injury correlated with site-specific O3 dose since
30	the respiratory bronchioles received the greatest O_3 dose (¹⁸ O mass/lung weight) and
31	sustained the greatest initial cellular injury. The respiratory bronchioles were also the site
32	of statistically significant GSH reduction. In addition, a study in isolated perfused rat
33	lungs found greater injury in conducting airways downstream of bifurcations where local
34	doses of O_3 were higher (<u>Postlethwait et al., 2000</u>).

1	In addition to the degree of initial injury, the degree of airways inflammation due to O_3
2	may have important long-term consequences since airways inflammation may lead to
3	tissue injury (Balmes et al., 1996). Persistent inflammation and injury, observed in animal
4	models of chronic and intermittent exposure to O ₃ , are associated with airways
5	remodeling, including mucous cell metaplasia of nasal transitional epithelium (Harkema
6	et al., 1999; Hotchkiss et al., 1991) and bronchiolar metaplasia of alveolar ducts
7	(Mudway and Kelly, 2000). Fibrotic changes such as deposition of collagen in the
8	airways and sustained lung function decrements especially in small airways have also
9	been demonstrated as a response to chronic O_3 exposure (Mudway and Kelly, 2000;
10	Chang et al., 1992). These effects, described in detail in Section 7.2.3.1, have been
11	demonstrated in rats exposed to levels of O ₃ as low as 0.25 ppm. Mechanisms responsible
12	for the resolution of inflammation and the repair of injury remain to be clarified and there
13	is only a limited understanding of the biological processes underlying long-term
14	morphological changes. However, a recent study in mice demonstrated a key role for the
15	TGF- β signaling pathway in the deposition of collagen in the airways wall following
16	chronic intermittent exposure to 0.5 ppm O_3 (Katre et al., 2011). Studies in infant
17	monkeys have also documented effects of chronic intermittent exposure to 0.5 ppm O_3 on
18	the developing lung and immune system. Extensive discussion of this topic is found in
19	Section $5.4.2.4$ (Lifestage) and in Section $7.2.3.1$.
20	It should be noted that repeated exposure to O ₃ results in attenuation of some O ₃ -induced
21	responses, including those associated with the activation of neural reflexes
22	(e.g., decrements in pulmonary function), as discussed in Section 5.3.2. However,

numerous studies demonstrate that some markers of injury and inflammation remain
increased during multi-day exposures to O₃. Mechanisms responsible for attenuation, or
the lack thereof, are incompletely understood.

5.3.8 Systemic Inflammation and Oxidative/Nitrosative Stress

27 been proposed that lipid oxidation products resulting from reaction of O_3 with lipids it	
	1
28 the ELF are responsible for systemic effects, however it is not known whether they ga	in
29 access to the vascular space (<u>Chuang et al., 2009</u>). Alternatively, extrapulmonary rele	ise
30 of diffusible mediators may initiate or propagate inflammatory responses in the vascu	ar
31 or systemic compartments (<u>Cole and Freeman, 2009</u>). A role for O_3 in modulating	
32 endothelin, a potent vasoconstrictor, has also been proposed. Studies in rats found tha	
33 exposure to 0.4 and 0.8 ppm O ₃ induced endothelin system genes in the lung and	
34 increased circulating levels of endothelin (<u>Thomson et al., 2006</u> ; <u>Thomson et al., 2005</u>).
35 Systemic oxidative stress may be suggested by studies in humans which reported	

1	associations between O_3 exposure and levels of plasma 8-isoprostanes and the presence
2	of peripheral blood lymphocyte micronuclei (<u>Chen et al., 2007a; Chen et al., 2006a</u>).
2	However, plasma isoprostanes are not a direct measure of systemic oxidative stress since
4	they are stable and can be generated in any compartment before diffusion into the
5	vascular space. Evidence of O_3 -mediated systemic oxidative stress is better provided by
6	new animal studies described below.
7	Ozone-induced perturbations of the cardiovascular system were recently investigated in
8	young mice and monkeys (<u>Chuang et al., 2009</u>) and in rats (<u>Kodavanti et al., 2011;</u>
9	Perepu et al., 2010) (see Section 6.3.3 and Section 7.3.1.2). These are the first studies to
10	suggest that systemic oxidative stress and inflammation play a mechanistic role in
11	O_3 -induced effects on the systemic vascular and heart. Exposure to 0.5 ppm O_3 for 5 days
12	resulted in oxidative/nitrosative stress, vascular dysfunction and mitochondrial DNA
13	damage in the aorta (<u>Chuang et al., 2009</u>). Chronic exposure to 0.8 ppm O ₃ resulted in an
14	enhancement of inflammation and lipid peroxidation in the heart following an ischemia-
15	reperfusion challenge (Perepu et al., 2010). In addition, chronic intermittent exposure to
16	0.4 ppm O_3 increased aortic levels of mRNA for biomarkers of oxidative stress,
17	thrombosis, vasoconstriction and proteolysis and aortic lectin-like oxidized-low density
18	lipoprotein receptor-1(LOX-1) mRNA and protein levels (Kodavanti et al., 2011). The
19	latter study suggests a role for circulating oxidized lipids in mediating the effects of O_3 .
20	Systemic inflammation and oxidative/nitrosative stress may similarly affect other organ
• •	
21	systems as well as the plasma compartment. Circulating cytokines have the potential to
21 22	systems as well as the plasma compartment. Circulating cytokines have the potential to enter the brain through diffusion and active transport and to contribute to
22	enter the brain through diffusion and active transport and to contribute to
22 23	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the
22 23 24	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (<u>Block and Calderón-Garcidueñas, 2009</u>) (see Section <u>6.4</u> and
22 23 24 25	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (<u>Block and Calderón-Garcidueñas, 2009</u>) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (<u>Block and Calderón-Garcidueñas</u> ,
22 23 24 25 26	 enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (<u>Block and Calderón-Garcidueñas, 2009</u>) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (<u>Block and Calderón-Garcidueñas, 2009</u>). Vagal afferent pathways originating in the RT may also be responsible for
22 23 24 25 26 27	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (<u>Block and Calderón-Garcidueñas, 2009</u>) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (<u>Block and Calderón-Garcidueñas,</u> <u>2009</u>). Vagal afferent pathways originating in the RT may also be responsible for O ₃ -mediated activation of nucleus tractus solitarius neurons which resulted in neuronal
22 23 24 25 26 27 28	 enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O₃-mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O₃ for
22 23 24 25 26 27 28 29	 enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section 6.4 and Section 7.5). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O₃-mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O₃ for 1.5-120 hours) (Gackière et al., 2011). Recent studies have demonstrated O₃-induced
22 23 24 25 26 27 28 29 30	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (<u>Block and Calderón-Garcidueñas</u> , 2009) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (<u>Block and Calderón-Garcidueñas</u> , 2009). Vagal afferent pathways originating in the RT may also be responsible for O ₃ -mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O ₃ for 1.5-120 hours) (<u>Gackière et al., 2011</u>). Recent studies have demonstrated O ₃ -induced brain lipid peroxidation, cytokine production in the brain and upregulated expression of
22 23 24 25 26 27 28 29 30 31	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O ₃ -mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O ₃ for 1.5-120 hours) (Gackière et al., 2011). Recent studies have demonstrated O ₃ -induced brain lipid peroxidation, cytokine production in the brain and upregulated expression of VEGF in rats (0.5 ppm O ₃ , 3 hours or 0.25-0.5 ppm O ₃ , 4 h/day, 15-60 days) (Guevara-
22 23 24 25 26 27 28 29 30 31 32	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O ₃ -mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O ₃ for 1.5-120 hours) (Gackière et al., 2011). Recent studies have demonstrated O ₃ -induced brain lipid peroxidation, cytokine production in the brain and upregulated expression of VEGF in rats (0.5 ppm O ₃ , 3 hours or 0.25-0.5 ppm O ₃ , 4 h/day, 15-60 days) (Guevara- Guzmán et al., 2009; Araneda et al., 2008; Pereyra-Muñoz et al., 2006). Further,
22 23 24 25 26 27 28 29 30 31 32 33	 enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section 6.4 and Section 7.5). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O₃-mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O₃ for 1.5-120 hours) (Gackière et al., 2011). Recent studies have demonstrated O₃-induced brain lipid peroxidation, cytokine production in the brain and upregulated expression of VEGF in rats (0.5 ppm O₃, 3 hours or 0.25-0.5 ppm O₃, 4 h/day, 15-60 days) (Guevara-Guzmán et al., 2009; Araneda et al., 2008; Pereyra-Muñoz et al., 2006). Further, O₃-induced oxidative stress resulted in increased plasma lipid peroxides (0.25 ppm,
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22 23 24 25 26 27 28 29 30 31 32 33 34 35	enter the brain through diffusion and active transport and to contribute to neuroinflammation, neurotoxicity, cerebrovascular damage and a break-down of the blood brain barrier (Block and Calderón-Garcidueñas, 2009) (see Section <u>6.4</u> and Section <u>7.5</u>). They can also activate neuronal afferents (Block and Calderón-Garcidueñas, 2009). Vagal afferent pathways originating in the RT may also be responsible for O ₃ -mediated activation of nucleus tractus solitarius neurons which resulted in neuronal activation in stress-responsive regions of the CNS in rats (0.5 or 2 ppm O ₃ for 1.5-120 hours) (<u>Gackière et al., 2011</u>). Recent studies have demonstrated O ₃ -induced brain lipid peroxidation, cytokine production in the brain and upregulated expression of VEGF in rats (0.5 ppm O ₃ , 3 hours or 0.25-0.5 ppm O ₃ , 4 h/day, 15-60 days) (<u>Guevara- Guzmán et al., 2009</u> ; <u>Araneda et al., 2008</u> ; <u>Pereyra-Muñoz et al., 2006</u>). Further, O ₃ -induced oxidative stress resulted in increased plasma lipid peroxides (0.25 ppm, 4h/day, 15-60 days) (<u>Santiago-López et al., 2010</u>), which was correlated with damage to specific brain regions (<u>Pereyra-Muñoz et al., 2006</u>).

1	steroidogenesis is blocked by oxidative stress (Diemer et al., 2003). It has been proposed
2	that lipid peroxidation of sperm plasma membrane may lead to impaired sperm mobility
3	and decreased sperm quality (Agarwal et al., 2003). Further, it has been proposed that
4	oxidative stress may damage DNA in the sperm nucleus and lead to apoptosis and a
5	decline in sperm counts (Agarwal et al., 2003). One study reported an association
6	between O ₃ exposure and semen quality and suggested oxidative stress as an underlying
7	mechanism (Sokol et al., 2006). Additional evidence is required to substantiate this link.
8	A role for plasma antioxidants in modulating O ₃ -induced respiratory effects was
9	suggested by a recent study (Aibo et al., 2010). In this study, pretreatment of rats with a
10	high dose of acetaminophen resulted in increased levels of plasma cytokines and the
11	influx of inflammatory cells into the lung following O ₃ exposure (0.25-0.5 ppm, 6 hours)
12	(<u>Aibo et al., 2010</u>). These effects were not observed in response to O_3 alone.
13	Furthermore, acetaminophen-induced liver injury was exacerbated by O3 exposure. A
14	greater increase in hepatic neutrophil accumulation and greater alteration in gene
15	expression profiles was observed in mice exposed to O_3 and acetaminophen compared
16	with either exposure alone (Aibo et al., 2010). Although not measured in this study,
17	glutathione depletion in the liver is known to occur in acetaminophen toxicity. Since liver
18	glutathione is the source of plasma glutathione, acetaminophen treatment may have
19	lowered plasma glutathione levels and altered the redox balance in the vascular
20	compartment. These findings indicate interdependence between RT, plasma and liver
21	responses to O_3 , possibly related to glutathione status.

5.3.9 Impaired Alveolar-Arterial Oxygen Transfer

22	O3 may impair alveolar-arterial oxygen transfer and reduce the supply of arterial oxygen
23	to the myocardium. This may have a greater impact in individuals with compromised
24	cardiopulmonary systems. Gong et al. (1998) provided evidence of a small decrease in
25	arterial oxygen saturation in human subjects exposed for 3 hours to 300 ppb O_3 while
26	exercising at a light to moderate level. In addition, <u>Delaunois et al. (1998</u>) demonstrated
27	pulmonary vasoconstriction in O ₃ -exposed rabbits (0.4 ppm, 4 hours). Although of
28	interest, the contribution of this pathway to O ₃ -induced cardiovascular effects remains
29	uncertain.

5.3.10 Summary

30This section summarizes the modes of action and toxicity pathways resulting from O331inhalation (Figure 5-8). These pathways provide a mechanistic basis for the health effects

1	which are described in detail in Chapters 6 and 7. Three distinct short-term responses
2	have been well-characterized in humans challenged with O_3 : decreased pulmonary
3	function, airways inflammation, and increased bronchial reactivity. In addition, O_3
4	exposure exacerbates, and possibly also causes, asthma and allergic airways disease in
5	humans. Animal studies have demonstrated airways remodeling and fibrotic changes in
6	response to chronic and intermittent O ₃ exposures and a wide range of other responses.
7	While the RT is the primary target tissue, cardiovascular and other organ effects occur
8	following short- and long-term exposures of animals to O ₃ .

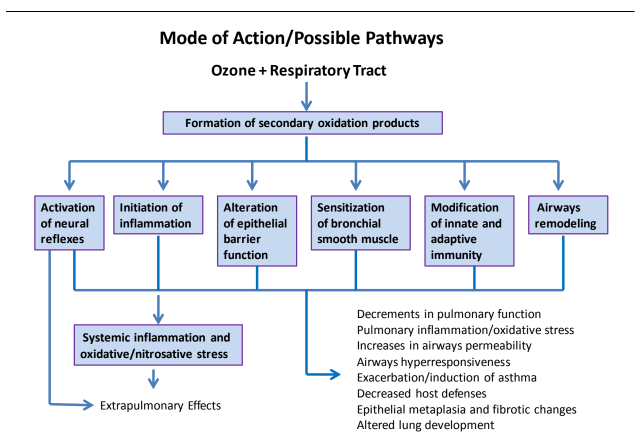


Figure 5-8 The modes of action/possible pathways underlying the health effects resulting from inhalation exposure to ozone.

9	The initial key event in the toxicity pathway of O ₃ is the formation of secondary oxidation
10	products in the RT. This mainly involves direct reactions with components of the ELF.
11	The resulting secondary oxidation products transmit signals to the epithelium,
12	nociceptive sensory nerve fibers and, if present, dendritic cells, mast cells and
13	eosinophils. Thus, the effects of O_3 are mediated by components of ELF and by the

multiple cell types found in the RT. Further, oxidative stress is an implicit part of this initial key event.

- 3 Another key event in the toxicity pathway of O_3 is the activation of neural reflexes which 4 lead to decrements in pulmonary function (see Section 6.2.1). Evidence is accumulating 5 that secondary oxidation products are responsible for this effect. Eicosanoids have been 6 implicated in humans, while both eicosanoids and aldehydes are effective in animal 7 models. Different receptors on bronchial C-fibers have been shown to mediate separate 8 effects of O_3 on pulmonary function. Nociceptive sensory nerves are involved in the 9 involuntary truncation of inspiration which results in decreases in FVC, FEV₁, tidal 10 volume and pain upon deep inspiration. Opioids block these responses while atropine has 11 only a minimal effect. New evidence in an animal model suggests that TRPA1 receptors 12 on bronchial C-fibers mediate this pathway. Ozone exposure also results in activation of 13 vagal sensory nerves and a mild increase in airway obstruction measured as increased 14 sRaw. Atropine and β -adrenergic agonists greatly inhibit this response in humans 15 indicating that the airways obstruction is due to bronchoconstriction. Other studies in 16 humans implicated SP release from bronchial C-fibers resulting in airway narrowing due 17 to either neurogenic edema or bronchoconstriction. New evidence in an animal model 18 suggests that the SP-NK receptor pathway caused bronchoconstriction following O₃ 19 exposure. Activation of neural reflexes also results in extrapulmonary effects such as 20 bradycardia.
- 21 Initiation of inflammation is also a key event in the toxicity pathway of O_3 . Secondary 22 oxidation products, as well as chemokines and cytokines elaborated by airway epithelial 23 cells and macrophages, have been implicated in the initiation of inflammation. Vascular 24 endothelial adhesion molecules may also play a role. Work from several laboratories 25 using human subjects and animal models suggest that O₃ triggers the release of 26 tachykinins such as SP from airway sensory nerves which could contribute to 27 downstream effects including inflammation (see Section 6.2.3 and Section 7.2.4). 28 Airways neutrophilia has been demonstrated in BALF, mucosal biopsy and induced 29 sputum samples. Influx of mast cells, monocytes and macrophages also occur. 30 Inflammation further contributes to O₃-mediated oxidative stress. Recent investigations 31 show that O_3 exposure leads to the generation of hydronan fragments from high 32 molecular weight polymers of hyaluronan normally found in the ELF in mice. 33 Hyaluronan activates TLR4 and CD44-dependent signaling pathways in macrophages, 34 and results in an increased number of macrophages in the BALF. Activation of these 35 pathways occurs later than the acute neutrophilic response suggesting that they may 36 contribute to longer-term effects of O₃. The mechanisms involved in clearing 37 O₃-provoked inflammation remain to be clarified. It should be noted that inflammation,

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1 as measured by airways neutrophilia, is not correlated with decrements in pulmonary 2 function as measured by spirometry. 3 A fourth key event in the toxicity pathway of O_3 is alteration of epithelial barrier 4 function. Increased permeability occurs as a result of damage to tight junctions between 5 epithelial cells subsequent to O₃-induced injury and inflammation. It may play a role in 6 allergic sensitization and in AHR (see Section 6.2.2, Section 6.2.6, and Section 7.2.5). 7 Tachykinins mediate this response while antioxidants may confer protection. Genetic 8 susceptibility has been associated with a functioning *Tlr4* and *Nos2* genes. 9 A fifth key event in the toxicity pathway of O_3 is the sensitization of bronchial smooth 10 muscle. Increased bronchial reactivity can be both a rapidly occurring and a persistent 11 response. The mechanisms responsible for early and later AHR are not well-understood 12 (see Section 6.2.2). One proposed mechanism of sensitization, O_3 -induced increases in 13 epithelial permeability, would improve access of agonist to smooth muscle receptors. The 14 evidence for this mechanism is not consistent. Another proposed mechanism, for which 15 there is greater evidence, is neurally-mediated sensitization. In humans exposed to O_3 , 16 atropine blocked the early AHR response indicating the involvement of cholinergic 17 postganglionic pathways. Animal studies demonstrated that O₃-induced AHR involved 18 vagally-mediated responses and local axon reflex responses through bronchopulmonary 19 C-fiber-mediated release of SP. Later phases of increased bronchial reactivity may 20 involve the induction of IL-1 β which in turn upregulates SP production. In guinea pigs, eosinophil-derived major basic protein contributed to the stimulation of cholinergic 21 22 postganglionic pathways. A novel role for hyaluronan in mediating the later phase effects 23 O₃-induced AHR has recently been demonstrated. Hyaluronan fragments stimulated AHR 24 in a TLR4- and CD44 receptor-dependent manner. Tachykinins and secondary oxidation 25 products of O_3 have been proposed as mediators of the early response and inflammation-26 derived products have been proposed as mediators of the later response. Inhibition of 27 arachidonic acid metabolism was ineffective in blocking O₃-induced AHR in humans 28 while in animal models mixed results were found. Other cytokines and chemokines have 29 been implicated in the AHR response to O₃ in animal models. 30 A sixth key event in the toxicity pathway of O_3 is the modification of innate/adaptive 31 immunity. While the majority of evidence for this key event comes from animal studies, 32 there are several studies suggesting that this pathway may also be relevant in humans. 33 Ozone exposure of human subjects resulted in recruitment of activated innate immune 34 cells to the airways. This included macrophages and monocytes with increased 35 expression of cell surface markers characteristic of innate immunity and antigen 36 presentation, the latter of which could contribute to exaggerated T-cell responses and the 37 promotion of an allergic phenotype. Evidence of dendritic cell activation was observed in

1	GSTM1 null human subjects exposed to O_3 , suggesting O_3 -mediated interaction between
2	the innate and adaptive immune systems. Animal studies further linked O_3 -mediated
3	activation of the innate immune system to the development of nonspecific AHR,
4	demonstrated an interaction between allergen and O_3 in the induction of nonspecific
5	AHR, and found that O_3 acted as an adjuvant for allergic sensitization through the
6	activation of both innate and adaptive immunity. Priming of the innate immune system by
7	O_3 was reported in mice. This resulted in an exaggerated response to endotoxin which
8	included enhanced TLR4 signaling in macrophages. Ozone-mediated impairment of the
9	function of SP-A, an innate immune protein, has also been demonstrated. Taken together
10	these studies provide evidence that O_3 can alter host immunologic response and lead to
11	immune system dysfunction. These mechanisms may underlie the exacerbation and
12	induction of asthma (see Section $6.2.6$ and Section $7.2.1$), as well as decreases in host
13	defense (see Section $6.2.5$ and Section $7.2.6$).
14	Another key event in the toxicity pathway of O_3 is airways remodeling. Persistent
15	inflammation and injury, which are observed in animal models of chronic and
16	intermittent exposure to O_3 , are associated with morphologic changes such as mucous
17	cell metaplasia of nasal epithelium, bronchiolar metaplasia of alveolar ducts and fibrotic
18	changes in small airways (see Section 7.2.3). Mechanisms responsible for these responses
19	are not well-understood. However a recent study in mice demonstrated a key role for the
20	TGF- β signaling pathway in the deposition of collagen in the airway wall following
21	chronic intermittent exposure to O_3 . Chronic intermittent exposure to O_3 has also been
22	shown to result in effects on the developing lung and immune system.
23	Systemic inflammation and vascular oxidative/nitrosative stress are also key events in the
24	toxicity pathway of O_3 . Extrapulmonary effects of O_3 occur in numerous organ systems,
25	including the cardiovascular, central nervous, reproductive and hepatic systems (see
26	Section 6.3 to Section 6.5 and Section 7.3 to Section 7.5). It has been proposed that lipid
27	oxidation products resulting from reaction of O3 with lipids and/or cellular membranes in
28	the ELF are responsible for systemic responses, however it is not known whether they
29	gain access to the vascular space. Alternatively, release of diffusible mediators from the
30	lung into the circulation may initiate or propagate inflammatory responses in the vascular
31	or in systemic compartments.

5.4 Interindividual Variability in Response

32	Responses to O_3 exposure are variable within the population (<u>Mudway and Kelly, 2000</u>).
33	Some studies have shown a large range of pulmonary function responses to O_3 among
34	healthy young adults (i.e., 4 hours to 200 ppb O_3 or for 1.5 hours to 420 ppb O_3 while

1	exercising at a moderate level) (Hazucha et al., 2003; Balmes et al., 1996). Since
2	individual responses were relatively consistent across time in these studies, it was thought
3	that responsiveness reflected an intrinsic characteristic of the subject (Mudway and Kelly,
4	2000). Other studies have shown that age and body mass index may influence
5	responsiveness to O _{3.} In human subjects exercising moderately and exposed to 420 ppb
6	O_3 for 1.5 hours, older adults were generally not responsive to O_3 (Hazucha et al., 2003),
7	while obese young women appeared to be more responsive than lean young women
8	(Bennett et al., 2007). In another study, the observed lack of spirometric responsiveness
9	in one group of human subjects was not attributable to the presence of endogenous
10	endorphins, which could vary between individuals and which could potentially block C-
11	fiber stimulation by O_3 (420 ppb, 2 hours, moderate exercise (Passannante et al., 1998).
12	Other responses to O ₃ have also been characterized by a large degree of interindividual
13	variability. For example, interindividual variability in the neutrophilic response has been
14	noted in human subjects (Holz et al., 1999; Devlin et al., 1991; Schelegle et al., 1991).
15	One study demonstrated a 3-fold difference in airways neutrophilia, measured as percent
16	of total cells in proximal BALF, among human subjects exposed to 300 ppb O_3 for 1 hour
17	while exercising at a heavy level (Schelegle et al., 1991). Another study reported a
18	20-fold difference in BAL neutrophils following exposure to $80-100$ ppb O ₃ for 6.6 hours
19	in human subjects exercising at a moderate level (Devlin et al., 1991). In contrast,
20	reproducibility of intraindividual responses to 1-hour exposure to 250 ppb O_3 in human
21	subjects exercising at a light level, measured as sputum neutrophilia, was demonstrated
22	by Holz et al. (1999). While the basis for the observed interindividual variability in
23	responsiveness to O ₃ is not clear, both dosimetric and mechanistic factors are likely to
24	contribute and will be discussed below.

5.4.1 Dosimetric Considerations

25	Two studies have investigated the correlation of O_3 uptake with the pulmonary function
26	responses to O_3 exposure (<u>Reeser et al., 2005</u> ; <u>Gerrity et al., 1994</u>). These studies found
27	that the large subject-to-subject variability in ΔFEV_1 response to O_3 does not appear to
28	have a dosimetric explanation. Reeser et al. (2005) found no significant relationship
29	between ΔFEV_1 and fractional absorption of O_3 using the bolus method. Contrary to
30	previous findings, the percent change in dead space volume of the respiratory tract
31	(% ΔV_D) did not correlate with O ₃ uptake, possibly due to the contraction of dead space
32	caused by airway closure. Gerrity et al. (1994) found that intersubject variability in FEV_1
33	and airway resistance was not related to differences in the O_3 dose delivered to the lower
34	airways, whereas minute ventilation was predictive of FEV ₁ decrement. No study has yet
35	demonstrated that subjects show a consistent pattern of O_3 retention when re-exposed

1over weeks of time, as has been shown to be the case for the FEV1 response, or that2within-subject variation in FEV1 response is related to fluctuations in O3 uptake.3However, these studies did not control for the differences in conducting airway volume4between individuals. By controlling for conducting airway volume, it may be possible to5estimate how much of the intersubject variation in FEV1 response at a given O3 exposure6is due to actual inter-individual variability in dose.

5.4.2 Mechanistic Considerations

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This section considers mechanistic factors that may contribute to variability in responses between individuals. It has been proposed that some of the variability may be genetically determined (<u>Yang et al., 2005a</u>). Besides gene-environment interactions, other factors such as pre-existing diseases and conditions, nutritional status, lifestage, attenuation, and co-exposures may also contribute to inter-individual variability in responses to O_3 and are discussed below.

5.4.2.1 Gene-environment Interactions

13 The pronounced interindividual variation in responses to O₃ infers that genetic 14 background may play an important role in responsiveness to O_3 (Cho and Kleeberger, 15 2007; Kleeberger et al., 1997) (see also Section 8.4). Strains of mice which are prone or 16 resistant to O₃-induced effects have been used to systematically identify candidate 17 susceptibility genes. Using these recombinant inbred strains of mice and exposures to 18 0.3 ppm O₃ for up to 72 hours, genome wide linkage analyses (also known as positional 19 cloning) demonstrated quantitative trait loci for O₃-induced lung inflammation and 20 hyperpermeability on chromosome 17 (Kleeberger et al., 1997) and chromosome 4 21 (Kleeberger et al., 2000), respectively. More specifically, these studies found that Tnf, 22 whose protein product is the inflammatory cytokine TNF- α , and Tlr4, whose protein 23 product is TLR4, were candidate susceptibility genes (Kleeberger et al., 2000; Kleeberger 24 et al., 1997). Other studies, which used targeted deletion, identified genes encoding iNOS 25 and heat shock proteins as TLR4 effector genes (Bauer et al., 2011; Kleeberger et al., 26 2001) and found that IL-10 protects against O₃-induced pulmonary inflammation (Backus 27 et al., 2010). Investigations in inbred mouse strains found that differences in expression 28 of certain proteins, such as CCSP (1.8 ppm O_3 for 3 hours) (Broeckaert et al., 2003) and 29 MARCO (0.3 ppm O_3 for up to 48 hours) (Dahl et al., 2007), were responsible for 30 phenotypic characteristics, such as epithelial permeability and scavenging of oxidized 31 lipids, respectively, which confer sensitivity to O_3 .

1 Genetic polymorphisms have received increasing attention as modulators of O₃-mediated 2 effects. Functionally relevant polymorphisms in candidate susceptibility genes have been 3 studied at the individual and population level in humans, and also in animal models. 4 Genes whose protein products are involved in antioxidant defense/oxidative stress and 5 xenobiotic metabolism, such as glutathione-S-transferase M1 (GSTM1) and 6 NADPH:quinone oxidoreductase 1 (NQO1), have also been a major focuses of these 7 efforts. This is because oxidative stress resulting from O_3 exposure is thought to 8 contribute to the pathogenesis of asthma, and because xenobiotic metabolism detoxifies 9 secondary oxidation products formed by O₃ which contribute to oxidative stress (Islam et 10 al., 2008). TNF- α is of interest since it is linked to a candidate O₃ susceptibility gene and 11 since it plays a key role in initiating airways inflammation (Li et al., 2006d). 12 Polymorphisms of genes coding for GSTM1, NOO1 and TNF- α have been associated 13 with altered risk of O₃-mediated effects (Li et al., 2006d; Yang et al., 2005a; Romieu et 14 al., 2004b; Corradi et al., 2002; Bergamaschi et al., 2001). Additional studies have 15 focused on functional variants in other genes involved in antioxidant defense such as 16 catalase (CAT), myeloperoxidase, heme oxygenase (HMOX-1) and manganese 17 superoxide dismutase (MnSOD) (Wenten et al., 2009; Islam et al., 2008). These studies 18 are discussed below.

19 GSTM1 is a phase II antioxidant enzyme which is transcriptionally regulated by 20 NF-e2-related factor 2-antioxidant response element (Nrf2-ARE) pathway. A large 21 proportion (40-50%) of the general public (across ethnic populations) has the 22 GSTM1-null genotype, which has been linked to an increased risk of health effects due to 23 exposure to air pollutants (London, 2007). A role for GSTs in metabolizing electrophiles 24 such as 4-hydroxynonenal, which is a secondary oxidation product resulting from O_3 25 exposure, has been demonstrated (Awasthi et al., 2004). A recent study found that the 26 GSTM1 genotype modulated the time course of the neutrophilic inflammatory response 27 following acute O_3 exposure (400 ppb for 2 hours with light to moderate exercise) in 28 healthy adults (Alexis et al., 2009). In GSTM1-null and -sufficient subjects, O₃-induced 29 sputum neutrophilia was similar at 4 hours. However, neutrophilia resolved by 24 hours 30 in sufficient subjects but not in GSTM1-null subjects. In contrast, no differences in 31 24 hour sputum neutrophilia were observed between GSTM1-null and -sufficient human 32 subjects exposed to 60 ppb O_3 for 2 hours with moderate exercise (Kim et al., 2011). It is 33 not known whether the effect seen at the higher exposure level (Alexis et al., 2009) was 34 due to the persistence of pro-inflammatory stimuli, impaired production of 35 downregulators or impaired neutrophil apoptosis and clearance. However, a subsequent 36 in vitro study by these same investigators found that GSTM1 deficiency in airway 37 epithelial cells enhanced IL-8 production in response to 0.4 ppm O₃ for 4 hours (Wu et 38 al., 2011). Furthermore, NF- κ B activation was required for O₃-induced IL-8 production 39 (Wu et al., 2011). Since IL-8 is a potent neutrophil activator and chemotaxin, this study

provides additional evidence for the role of GSTM1 as a modulator of inflammatory responses due to O_3 exposure.

- 3 In addition, O_3 exposure increased the expression of the surface marker CD14 in airway 4 neutrophils of GSTM1-null subjects to a greater extent than in sufficient subjects (Alexis 5 et al., 2009). Furthermore, differences in airway macrophages were noted between the 6 GSTM1-sufficient and -null subjects. Numbers of airway macrophages were decreased at 7 4 and 24 hours following O_3 exposure in GSTM1-sufficient subjects (Alexis et al., 2009). 8 Airway macrophages in GSTM1-null subjects were greater in number and found to have 9 greater oxidative burst and phagocytic capability than those of sufficient subjects. Airway 10 macrophages and dendritic cells from GSTM1-null subjects exposed to O₃ expressed 11 higher levels of the surface marker HLA-DR, suggesting activation of the innate immune 12 system (Alexis et al., 2009). These differences in inflammatory responses between the 13 GSTM1-null and -sufficient subjects may provide biological plausibility for the 14 differences in O₃-mediated effects reported in controlled human exposure studies (Corradi et al., 2002; Bergamaschi et al., 2001). It should also be noted that GSTM1 15 16 genotype did not affect the acute pulmonary function (i.e., spirometric) response to O_3 17 which provides additional evidence for separate mechanisms underlying the effects of O_3 18 on pulmonary function and inflammation in adults (Alexis et al., 2009). However, 19 GSTM1-null asthmatic children were previously found to be more at risk of O₃-induced 20 effects on pulmonary function than GSTM1-sufficient asthmatic children (Romieu et al., 21 2004b).
- 22 Another enzyme involved in the metabolism of secondary oxidation products is NQO1. 23 NQO1 catalyzes the 2-electron reduction by NADPH of quinones to hydroquinones. 24 Depending on the substrate, it is capable of both protective detoxification reactions and 25 redox cycling reactions resulting in the generation of reactive oxygen species. A recent 26 study using NQO1-null mice demonstrated that NQO1 contributes to O3-induced 27 oxidative stress, AHR and inflammation following a 3-hour exposure to 1 ppm O₃ 28 (Voynow et al., 2009). These experimental results may provide biological plausibility for 29 the increased biomarkers of oxidative stress and increased pulmonary function 30 decrements observed in O_3 -exposed individuals bearing both the wild-type NOO1 gene 31 and the null GSTM1 gene (Corradi et al., 2002; Bergamaschi et al., 2001).
- 32Besides enzymatic metabolism, other mechanisms participate in the removal of33secondary oxidation products formed as a result of O3 inhalation. One involves34scavenging of oxidized lipids via the macrophage receptor with collagenous structure35(MARCO) expressed on the cell surface of alveolar macrophages. A recent study36demonstrated increased gene expression of MARCO in the lungs of an O3-resistant C3H37mouse strain (HeJ) but not in an O3-sensitive, genetically similar strain (OuJ) (Dahl et al.,

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1	<u>2007</u>). Upregulation of MARCO occurred in mice exposed to 0.3 ppm O_3 for
2	24-48 hours; inhalation exposure for 6 hours at this concentration was insufficient for this
3	response. Animals lacking the MARCO receptor exhibited greater inflammation and
4	injury, as measured by BAL neutrophils, protein and isoprostanes, following exposure to
5	0.3 ppm O ₃ (Dahl et al., 2007). MARCO also protected against the inflammatory effects
6	of oxidized surfactant lipids (Dahl et al., 2007). Scavenging of oxidized lipids may limit
7	O ₃ -induced injury since ozonized cholesterol species formed in the ELF (mice, 0.5-3 ppm
8	O ₃ , 3 hours) (Pulfer et al., 2005; Pulfer and Murphy, 2004) stimulated apoptosis and
9	cytotoxicity in vitro (Gao et al., 2009b; Sathishkumar et al., 2009; Sathishkumar et al.,
10	<u>2007b; Sathishkumar et al., 2007a</u>).
11	Two studies reported relationships between <i>TNF</i> promoter variants and O ₃ -induced
12	effects in humans. In one study, O ₃ -induced change in lung function was significantly
13	lower in adult subjects with TNF promoter variants -308A/A and -308G/A compared with
14	adult subjects with the variant -308G/G (Yang et al., 2005a). This response was
15	modulated by a specific polymorphism of LTA (Yang et al., 2005a), a previously
16	identified candidate susceptibility gene whose protein product is lymphotoxin- α
17	(Kleeberger et al., 1997). In the second study, an association between the TNF promoter
18	variant -308G/G and decreased risk of asthma and lifetime wheezing in children was
19	found (Li et al., 2006d). The protective effect on wheezing was modulated by ambient O_3
20	levels and by GSTM1 and GSTP1 polymorphisms. The authors suggested that the
21	TNF-308 G/G genotype may have a protective role in the development of childhood
22	asthma (<u>Li et al., 2006d</u>).
23	Similarly, a promoter variant of the gene HMOX-1, consisting of a smaller number of
24	(GT)n repeats, was associated with a reduced risk for new-onset asthma in non-Hispanic
25	white children (Islam et al., 2008). The number of $(GT)_n$ repeats in this promoter has
26	been shown to be inversely related to the inducibility of HMOX-1. A modulatory effect of
27	O_3 was demonstrated since the beneficial effects of this polymorphism were seen only in
28	children living in low O_3 communities (Islam et al., 2008). This study also identified an
29	association between a polymorphism of the CAT gene and increased risk of new-onset
30	asthma in Hispanic children; however no modulation by O_3 was seen (Islam et al., 2008).
31	No association was observed in this study between a MnSOD polymorphism and asthma
32	(<u>Islam et al., 2008</u>).
33	Studies to date indicate that some variability in individual responsiveness to O ₃ may be
34	accounted for by functional genetic polymorphisms. Further, the effects of
35	gene-environment interactions may be different in children and adults.

5.4.2.2 Pre-existing Diseases and Conditions

1	Pre-existing diseases and conditions can alter the response to O_3 exposure. For example,
2	responsiveness to O ₃ , as measured by spirometry, is decreased in smokers and individuals
3	with COPD (U.S. EPA, 2006b). Asthma and allergic diseases are of major importance in
4	this discussion. In individuals with asthma, there is increased responsiveness to
5	bronchoconstrictor challenge. This results from a combination of structural and
6	physiological factors including increased airway inner-wall thickness, smooth muscle
7	responsiveness and mucus secretion. Although inflammation is likely to contribute, its
8	relationship to AHR is not clear (U.S. EPA, 2006b). However, some asthmatics have
9	higher baseline levels of neutrophils, lymphocytes, eosinophils and mast cells in
10	bronchial washes and bronchial biopsy tissue (Stenfors et al., 2002). It has been proposed
11	that enhanced sensitivity to O_3 is conferred by the presence of greater numbers of resident
12	airway inflammatory cells in disease states such as asthma (Mudway and Kelly, 2000).
13	In order to determine whether asthmatics exhibit greater responses to O ₃ , several earlier
14	studies compared pulmonary function in asthmatic and non-asthmatic subjects following
15	O_3 exposure. Some also probed mechanisms which could account for enhanced
16	sensitivity. While the majority focused on measurements of FEV_1 and FVC and found no
17	differences between the two groups following exposures of 2-4 hours to 125-250 ppb O_3
18	or to a 30-minute exposure to 120-180 ppb O_3 by mouthpiece in human subjects
19	exercising at a light to moderate level (<u>Stenfors et al., 2002</u> ; <u>Mudway et al., 2001</u> ; <u>Holz et</u>
20	al., 1999; Scannell et al., 1996; Koenig et al., 1987; Linn et al., 1978), there were notable
21	exceptions. In one study, greater airways obstruction in asthmatics compared with non-
22	asthmatic subjects was observed immediately following a 2-hour exposure to 400 ppb O ₃
23	while exercising at a heavy level (Kreit et al., 1989). These changes were measured as
24	statistically significant greater decreases in FEV_1 and in $FEF_{25.75}$ (but not in FVC) in the
25	absence of a bronchoconstrictor challenge (Kreit et al., 1989). These results suggest that
26	this group of asthmatics responded to O_3 -exposure with a greater degree of vagally-
27	mediated bronchoconstriction compared with the non-asthmatics. A second study
28	demonstrated a statistically significant greater decrease in FEV_1 and in FEV_1/FVC (but
29	not in FVC) in asthmatics compared with non-asthmatics exposed to 160 ppb O_3 for
30	7.6 hours with light exercise (Horstman et al., 1995). These responses were accompanied
31	by wheezing and inhaler use in the asthmatics (Horstman et al., 1995). Aerosol bolus
32	dispersion measurements demonstrated a statistically significant greater change in
33	asthmatics compared with non-asthmatics, which was suggestive of O ₃ -induced small
34	airway dysfunction (Horstman et al., 1995). Furthermore, a statistically significant
35	correlation was observed between the degree of baseline airway status and the FEV_1
36	response to O_3 in the asthmatic subjects (Horstman et al., 1995). A third study found
37	similar decreases in FVC and FEV_1 in both asthmatics and non-asthmatics exposed to

1	400 ppb O_3 for 2 hours with light exercise (Alexis et al., 2000). However, a statistically
2	significant decrease in FEF ₇₅ , a measure of small airway function, was observed in
3	asthmatics but not in non-asthmatics (<u>Alexis et al., 2000</u>). Taken together, these latter
4	studies indicate that while the magnitude of restrictive type spirometric decline was
5	similar in asthmatics and non-asthmatics, that obstructive type changes
6	(i.e., bronchoconstriction) were greater in asthmatics. Further, asthmatics exhibited
7	greater sensitivity to O_3 in terms of small airways function.
8	Since asthma exacerbations occur in response to allergens and/or other triggers, some
9	studies have focused on O ₃ -induced changes in AHR following a bronchoconstrictor
10	challenge. No difference in sensitivity to methacholine bronchoprovocation was observed
11	between asthmatics and non-asthmatics exposed to 400 ppb O ₃ for 2 hours while
12	exercising at a heavy level (Kreit et al., 1989). However, increased bronchial reactivity to
13	inhaled allergens was demonstrated in mild allergic asthmatics exposed to 160 ppb for
14	7.6 hours, 250 ppb for 3 hours and 120 ppb for 1 hour while exercising at a light level or
15	at rest (Kehrl et al., 1999; Jorres et al., 1996; Molfino et al., 1991) and in allergen-
16	sensitized guinea pigs following O ₃ exposure (1 ppm, 1 hour) (<u>Sun et al., 1997</u>). Similar,
17	but modest, responses were reported for individuals with allergic rhinitis (Jorres et al.,
18	1996). Further, the contractile response of isolated airways from human donor lung
19	tissue, which were sensitized and challenged with allergen, was increased by
20	pre-exposure to 1 ppm O_3 for 20 (<u>Roux et al., 1999</u>). These studies provide support for
21	O ₃ -mediated enhancement of responses to allergens in allergic subjects.
22	In terms of airways neutrophilia, larger responses were observed in asthmatics compared
23	to non-asthmatics subjects, who were exercising at a light to moderate level and exposed
24	to O_3 , in some (<u>Balmes et al., 1997</u> ; <u>Scannell et al., 1996</u> ; <u>Basha et al., 1994</u>) but not all
25	(Mudway et al., 2001) of the earlier studies. While each of these studies involved
26	exposure of exercising human subjects to 200 ppb O_3 , the duration of exposure was
27	longer (i.e., 4-6 hours) in the former studies than in the latter study (2 hours). Further,
28	statistically significantly increases in myeloperoxidase levels (an indicator of neutrophil
29	activation) in bronchial washes was observed in mild asthmatics compared with non-
30	asthmatics, despite no difference in O_3 -stimulated neutrophil influx between the 2 groups
31	following exposure to 200 ppb O_3 for 2 hours with moderate exercise (Stenfors et al.,
32	2002). A more recent study found that atopic asthmatic subjects exhibited an enhanced
33	inflammatory response to O_3 (400 ppb, 4 hours, with light to moderate exercise)
34	(<u>Hernandez et al., 2010</u>). This response was characterized by greater numbers of
35	neutrophils, higher levels of IL-6, IL-8 and IL-1 β and greater macrophage cell-surface
36	expression of TLR4 and IgE receptors in induced sputum compared with healthy
37	subjects. This study also reported a greater increase in hyaluronan in atopic subjects and
38	atopic asthmatics compared with healthy subjects following O_3 exposure. Animal studies
-	

- have previously reported that hyaluronic acid activates TLR4 signaling and results in
 AHR (see Section 5.3.5). Furthermore, levels of IL-10, a potent anti-inflammatory
 cytokine, were greatly reduced in atopic asthmatics compared to healthy subjects. These
 results provide evidence that innate immune and adaptive responses are different in
 asthmatics and healthy subjects exposed to O₃.
- 6 Eosinophils may be an important modulator of responses to O_3 in asthma and allergic 7 airways disease. Eosinophils and associated proteins are thought to affect muscarinic 8 cholinergic receptors which are involved in vagally-mediated bronchoconstriction 9 (Mudway and Kelly, 2000). Studies described in Section 5.3.5 which demonstrated a key 10 role of eosinophils in O₃-mediated AHR may be relevant to human allergic airways 11 disease which is characterized by airways eosinophilia (Yost et al., 2005). Furthermore, 12 O_3 exposure sometimes results in airways eosinophilia in allergic subjects or animal 13 models. For example, eosinophilia of the nasal and other airways was observed in 14 individuals with pre-existing allergic disease following O_3 inhalation (160 ppb, 7.6 hours 15 with light exercise and 270 ppb, 2 hours with moderate exercise) (Vagaggini et al., 2002; 16 Peden et al., 1997). Further, O_3 exposure (0.5 ppm, 8 hours/day for 1-3 days) increased 17 allergic responses, such as eosinophilia and augmented intraepithelial mucosubstances, in 18 the nasal airways of ovalbumin (OVA)-sensitized rats (Wagner et al., 2002). In contrast, 19 Stenfors et al. (2002) found no stimulation of eosinophil influx measured in bronchial 20 washes and BALF of mild asthmatics following exposure to a lower concentration 21 (200 ppb, 2 hours, with moderate exercise) of O_3 .
- 22 The role of mast cells in O₃-mediated asthma exacerbations has been investigated. Mast 23 cells are thought to play a key role in O_3 -induced airways inflammation, since airways 24 neutrophilia was decreased in mast cell-deficient mice exposed to O_3 (Kleeberger et al., 25 1993). However, another study found that mast cells were not involved in the 26 development of increased bronchial reactivity in O_3 -exposed mice (Noviski et al., 1999). 27 Nonetheless, mast cells release a wide variety of important inflammatory mediators 28 which may lead to asthma exacerbations (Stenfors et al., 2002). A large increase in mast 29 cell number in bronchial submucosa was observed in non-asthmatics and a significant 30 decrease in mast cell number in bronchial epithelium was observed in mild asthmatics 31 6 hours following exposure to 200 ppb O₃ for 2 hours during mild exercise (Stenfors et 32 al., 2002). While these results point to an O_3 -mediated flux in bronchial mast cell 33 populations which differed between the non-asthmatics and mild asthmatics, 34 interpretation of these findings is difficult. Furthermore, mast cell number did not change 35 in airway lavages in either group in response to O₃ (Stenfors et al., 2002)
- Cytokine profiles in the airways have been investigated as an indicator of O₃ sensitivity.
 Differences in epithelial cytokine expression were observed in bronchial biopsy samples

1	in non-asthmatic and asthmatic subjects both at baseline and 6-hours postexposure to
2	200 ppb O_3 for 2 hours with moderate exercise (Bosson et al., 2003). The asthmatic
3	subjects had a higher baseline expression of IL-4 and IL-5 compared to non-asthmatics.
4	In addition, expression of IL-5, IL-8, GM-CSF, and ENA-78 in asthmatics was increased
5	significantly following O_3 exposure compared to non-asthmatics (Bosson et al., 2003).
6	Some of these (IL-4, IL-5 and GM-CSF) are Th2-related cytokines or neutrophil
7	chemoattractants, and play a role in IgE production, airways eosinophilia and suppression
8	of Th1-cytokine production (Bosson et al., 2003). These findings suggest a link between
9	adaptive immunity and enhanced responses of asthmatics to O_3 .
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10	A further consideration is the compromised status of ELF antioxidants in disease states
11	such as asthma (Mudway and Kelly, 2000). This could possibly be due to ongoing
12	inflammation which causes antioxidant depletion or to abnormal antioxidant transport or
13	synthesis (Mudway and Kelly, 2000). For example, basal levels of AH2 were
14	significantly lower and basal levels of oxidized GSH and UA were significantly higher in
15	bronchial wash fluid and BALF of mild asthmatics compared with healthy control
16	subjects (Mudway et al., 2001). Differences in ELF antioxidant content have also been
17	noted between species. These observations have led to the suggestion that the amount and
18	composition of ELF antioxidants, the capacity to replenish antioxidants in the ELF or the
19	balance between beneficial and injurious interactions between antioxidants and O_3 may
20	contribute to O_3 sensitivity, which varies between individuals and species (Mudway et
21	al., 2006; Mudway and Kelly, 2000; Mudway et al., 1999a). The complexity of these
22	interactions was demonstrated by a study in which a 2-hour exposure to 200 ppb O_3 ,
23	while exercising at a moderate level, resulted in similar increases in airway neutrophils
24	and decreases in pulmonary function in both mild asthmatics and healthy controls,
25	despite differences in ELF antioxidant concentrations prior to O_3 exposure (Mudway et
26	al., 2001). Further, the O ₃ -induced increase in oxidized GSH and decrease in AH2
27	observed in ELF of healthy controls was not observed in mild asthmatics (Mudway et al.,
28	2001). While the authors concluded that basal AH2 and oxidized GSH concentrations
29	were not predictive of responsiveness to O ₃ , they also suggested that the increased basal
30	UA concentrations in the mild asthmatics may have played a protective role (Mudway et
31	al., 2001). Thus compensatory mechanisms resulting in enhanced total antioxidant
32	capacity may play a role in modulating responses to O ₃ .
33	Collectively these older and more recent studies provide insight into mechanisms which
34	may contribute to enhanced responses of asthmatic and atopic individuals following O ₃
35	exposure. Greater airways inflammation and/or greater bronchial reactivity have been
36	demonstrated in asthmatics compared to non-asthmatics. This pre-existing inflammation
37	and altered baseline bronchial reactivity may contribute to the enhanced
38	bronchoconstriction seen in asthmatics exposed to O ₃ . Furthermore, O ₃ -induced

1	inflammation may contribute to O3-mediated AHR. An enhanced neutrophilic response to
2	O3 has been demonstrated in some asthmatics. A recent study in humans provided
3	evidence for differences in innate immune responses related to TLR4 signaling between
4	asthmatics and healthy subjects. Animal studies have demonstrated a role for eosinophil-
5	derived proteins in mediating the effects of O ₃ . Since airways eosinophilia occurs in both
6	allergic humans and allergic animal models, this pathway may underlie the exacerbation
7	of allergic asthma by O_3 . In addition, differences have been noted in epithelial cytokine
8	expression in bronchial biopsy samples of healthy and asthmatic subjects. A Th2
9	phenotype, indicative of adaptive immune system activation and enhanced allergic
10	responses, was observed before O_3 exposure and was increased by O_3 exposure in
11	asthmatics. These findings support links between innate and adaptive immunity and
12	sensitivity to O ₃ -mediated effects in asthmatics and allergic airways disease.

- 13 In addition to asthma and allergic diseases, obesity may alter responses to O_3 . While O_3 is 14 a trigger for asthma, obesity is a known risk factor for asthma (Shore, 2007). The 15 relationship between obesity and asthma is not well understood but recent investigations 16 have focused on alterations in endocrine function of adipose tissue in obesity. It is 17 thought that the increases in serum levels of factors produced by adipocytes 18 (i.e., adipokines), such as cytokines, chemokines, soluble cytokine receptors and energy 19 regulating hormones, may contribute to the relationship between obesity and asthma. 20 Some of these same mechanisms may be relevant to insulin resistant states such as 21 metabolic syndrome.
- 22 In a re analysis of the data of Hazucha et al. (2003), increasing body mass index in 23 young women was associated with increased O₃ responsiveness, as measured by 24 spirometry following a 1.5-hour exposure to 420 ppb O_3 while exercising at a moderate 25 level (Bennett et al., 2007). In several mouse models of obesity, airways were found to be 26 innately more hyperresponsive and responded more vigorously to acute O_3 exposure than 27 lean controls (Shore, 2007). Pulmonary inflammatory and injury in response to O_3 were 28 also enhanced (Shore, 2007). It was postulated that oxidative stress resulting from 29 obesity-related hyperglycemia could account for these effects (Shore, 2007). However, 30 responses to O_3 in the different mouse models are somewhat variable and depend on 31 whether exposures are acute or subacute. For example, diet-induced obesity augmented 32 inflammation and injury, as measured by BALF markers, and enhanced AHR in mice 33 exposed acutely to O_3 (2 ppm, 3 hours) (Johnston et al., 2008). In contrast, the 34 inflammatory response following sub-acute exposure to O_3 was dampened by obesity in a 35 different mouse model (0.3 ppm, 72 hours) (Shore et al., 2009). It is not known whether 36 differences in responsiveness to O_3 are due to differences in lung development in 37 genetically-modified animals which result in smaller lungs and thus to differences in 38 inhaled dose because of the altered body mass to lung size ratio.

5.4.2.3 Nutritional Status

1	
1	A further consideration is the compromised status of ELF antioxidants in nutritional
2	deficiencies. Thus, many investigations have focused on antioxidant deficiency and
3	supplementation as modulators of O_3 -mediated effects. One study in mice found that
4	vitamin A deficiency enhanced lung injury induced by exposure to 0.3 ppm O_3 for
5	72 hours (<u>Paquette et al., 1996</u>). Ascorbate deficiency was shown to increase the effects
6	of acute (0.5-1 ppm for 4 hours), but not subacute (0.2-0.8 ppm for 7 days), O_3 exposure
7	in guinea pigs (Kodavanti et al., 1995; Slade et al., 1989). Supplementation with AH2
8	and α -TOH was protective in healthy adults who were on an AH2-deficient diet and
9	exposed to 400 ppb O_3 for 2 hours while exercising at a moderate level (<u>Samet et al.</u> ,
10	<u>2001</u>). In this study, the protective effect consisted of a smaller reduction in FEV_1
11	following O_3 exposure (<u>Samet et al., 2001</u>). However the inflammatory response (influx
12	of neutrophils and levels of IL-6) measured in BALF 1 hour after O ₃ exposure was not
13	different between supplemented and non-supplemented subjects (Samet et al., 2001).
14	Other investigators found that AH2 and α -TOH supplementation failed to ameliorate the
15	pulmonary function decrements or airways neutrophilia observed in humans exposed to
16	200 ppb O_3 for 2 hours while exercising at a moderate level (Mudway et al., 2006). It was
17	suggested that supplementation may be ineffective in the absence of antioxidant
18	deficiency (<u>Mudway et al., 2006</u>).
19	In asthmatic adults, these same dietary antioxidants reduced O ₃ -induced bronchial
19 20	In asthmatic adults, these same dietary antioxidants reduced O ₃ -induced bronchial hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore,
	· · · · · · · · · · · · · · · · · · ·
20	hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore,
20 21	hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α -tocopherol protected against pulmonary function
20 21 22	hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α -tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of
20 21 22 23	hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α -tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O ₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004;
20 21 22 23 24	hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α -tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O ₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α -tocopherol and
20 21 22 23 24 25	 hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (Romieu et al., 2002).
20 21 22 23 24 25 26	hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α -tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O ₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; <u>Romieu et al., 2002</u>). Similarly, supplementation with ascorbate, α -tocopherol and β -carotene improved pulmonary function in Mexico City street workers (<u>Romieu et al., 1998b</u>).
20 21 22 23 24 25 26 27	 hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (Romieu et al., 1998b). Protective effects of supplementation with α-tocopherol alone have not been observed in
20 21 22 23 24 25 26 27 28	 hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (Romieu et al., 1998b). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (Mudway and Kelly, 2000). Alpha-TOH
20 21 22 23 24 25 26 27 28 29	 hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (Romieu et al., 1998b). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (Mudway and Kelly, 2000). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of
20 21 22 23 24 25 26 27 28 29 30	 hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (<u>Sienra-Monge et al., 2004</u>; <u>Romieu et al., 2002</u>). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (<u>Romieu et al., 1998b</u>). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (<u>Mudway and Kelly, 2000</u>). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of allergic rhinosinusitis and lower airways allergic inflammation (rats, 1 ppm O₃ for
20 21 22 23 24 25 26 27 28 29 30 31	 hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (<u>Sienra-Monge et al., 2004</u>; <u>Romieu et al., 2002</u>). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (<u>Romieu et al., 1998b</u>). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (<u>Mudway and Kelly, 2000</u>). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of allergic rhinosinusitis and lower airways allergic inflammation (rats, 1 ppm O₃ for 2 days) (<u>Wagner et al., 2007</u>). However, protection in these same animal models was
20 21 22 23 24 25 26 27 28 29 30 31 32	 hyperresponsiveness (120 ppb, 45 min, light exercise) (Trenga et al., 2001). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (Sienra-Monge et al., 2004; Romieu et al., 2002). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (Romieu et al., 1998b). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (Mudway and Kelly, 2000). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of allergic rhinosinusitis and lower airways allergic inflammation (rats, 1 ppm O₃ for 2 days) (Wagner et al., 2007). However, protection in these same animal models was reported using γ-TOH supplementation (Wagner et al., 2009; Wagner et al., 2007).
20 21 22 23 24 25 26 27 28 29 30 31 32 33	 hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (<u>Sienra-Monge et al., 2004</u>; <u>Romieu et al., 2002</u>). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (<u>Romieu et al., 1998b</u>). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (<u>Mudway and Kelly, 2000</u>). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of allergic rhinosinusitis and lower airways allergic inflammation (rats, 1 ppm O₃ for 2 days) (<u>Wagner et al., 2007</u>). However, protection in these same animal models was reported using γ-TOH supplementation (<u>Wagner et al., 2009</u>; <u>Wagner et al., 2007</u>). Whether or not this effect was due to enhanced antioxidant status or to activated signaling
20 21 22 23 24 25 26 27 28 29 30 31 32 33 34	 hyperresponsiveness (120 ppb, 45 min, light exercise) (<u>Trenga et al., 2001</u>). Furthermore, supplementation with AH2 and α-tocopherol protected against pulmonary function decrements and nasal inflammatory responses which were associated with high levels of ambient O₃ in asthmatic children living in Mexico City (<u>Sienra-Monge et al., 2004</u>; <u>Romieu et al., 2002</u>). Similarly, supplementation with ascorbate, α-tocopherol and β-carotene improved pulmonary function in Mexico City street workers (<u>Romieu et al., 1998b</u>). Protective effects of supplementation with α-tocopherol alone have not been observed in humans experimentally exposed to O₃ (<u>Mudway and Kelly, 2000</u>). Alpha-TOH supplementation also failed to protect against O₃-induced effects in animal models of allergic rhinosinusitis and lower airways allergic inflammation (rats, 1 ppm O₃ for 2 days) (<u>Wagner et al., 2007</u>). However, protection in these same animal models was reported using γ-TOH supplementation (<u>Wagner et al., 2009</u>; <u>Wagner et al., 2007</u>). Whether or not this effect was due to enhanced antioxidant status or to activated signaling pathways is not known. Other investigators found that α-TOH deficiency led to an

1	6 hours/day for 3 days) (Vasu et al., 2010). A recent study used α -TOH transfer protein
2	null mice as a model of α -TOH deficiency and demonstrated an altered adaptive response
3	of the lung genome to O_3 exposure (<u>Vasu et al., 2010</u>). Taken together, these studies
4	provide evidence that the tocopherol system modulates O ₃ -induced responses.

5.4.2.4 Lifestage

5	Responses to O ₃ are modulated by factors associated with lifestage. On one end of the
6	lifestage spectrum is aging. The spirometric response to O_3 appears to be lost in humans
7	as they age, as was demonstrated in two studies involving exposures of human subjects
8	exercising at levels ranging from light to heavy to 420-450 ppb O_3 for 1.5-2 hours
9	(<u>Hazucha et al., 2003;</u> <u>Drechsler-Parks, 1995</u>). In mice, physiological responses to O_3
10	(600 ppb, 2 hours) were diminished with age (Hamade et al., 2010). Mechanisms
11	accounting for this effect have not been well-studied but could include altered number
12	and sensitivity of receptors, altered signaling pathways involved in neural reflexes or
13	compromised status of ELF antioxidants.
14	On the other side of the lifestage spectrum is pre/postnatal development. Critical
15	windows of development during the pre/postnatal period are associated with an enhanced
16	sensitivity to environmental toxicants. Adverse birth outcomes and developmental
17	disorders may occur as a result (Section 7.4).
18	Adverse birth outcomes may result from stressors which impact transplacental oxygen
19	and nutrient transport by a variety of mechanisms including oxidative stress, placental
20	inflammation and placental vascular dysfunction (Kannan et al., 2006). These
21	mechanisms may be linked since oxidative/nitrosative stress is reported to cause vascular
22	dysfunction in the placenta (Myatt et al., 2000). As described earlier in this chapter and in
23	Section 7.4, systemic inflammation and oxidative/nitrosative stress and modification of
24	innate and adaptive immunity are key events underlying the health effects of O_3 and as
25	such they may contribute to adverse birth outcomes. An animal toxicology study showing
26	that exposure to 2 ppm O_3 led to anorexia (<u>Kavlock et al., 1979</u>) (see Section <u>7.4.2</u>) in
27	exposed rat dams provide an additional mechanism by which O ₃ exposure could lead to
28	diminished transplacental nutrient transport. Disturbances of the pituitary-adrenocortico-
29	placental system (Ritz et al., 2000) may also impact normal intrauterine growth and
30	development. Further, restricted fetal growth may result from pro-inflammatory
31	cytokines which limit trophoblast invasion during the early stages of pregnancy (Hansen
32	et al., 2008). Direct effects on maternal health, such as risk of infection, and on fetal
33	health, such as DNA damage, have also been proposed as mechanisms underlying
34	adverse birth outcomes (Ritz et al., 2000). In addition to restricted fetal growth, preterm

- 1 birth may contribute to adverse birth outcomes. Preterm birth may result from the 2 development of premature contractions and/or premature rupture of membranes as well 3 as from disrupted implantation and placentation which results in suboptimal placental 4 function (Darrow et al., 2009; Ritz et al., 2000). Genetic mutations are thought to be an 5 important cause of placental abnormalities in the first trimester, while vascular alterations 6 may be the main cause of placental abnormalities in later trimesters (Jalaludin et al., 7 2007). Ozone-mediated systemic inflammation and oxidative stress/nitrosative stress may 8 possibly be related to these effects although there is no firm evidence.
- 9 Enhanced sensitivity to environmental toxicants during critical windows of development 10 may also result in developmental disorders. For example, normal migration and 11 differentiation of neural crest cells are important for heart development and are 12 particularly sensitive to toxic insults (Ritz et al., 2002). Further, immune dysregulation 13 and related pathologies are known to be associated with pre/postnatal environmental 14 exposures (Dietert et al., 2010). Ozone exposure is associated with developmental effects 15 in several organ systems. These include the lung and immune system (see below) and 16 neurobehavioral changes which could reflect the effect of O_3 on CNS plasticity or the hypothalamic-pituitary axis (Auten and Foster, 2011) (see Section 7.4.9). 17
- 18 The majority of developmental effects due to O_3 have been described for the respiratory 19 system (see Section 7.2.3 and 7.4.8). Since its growth and development take place during 20 both the prenatal and early postnatal periods, both prenatal and postnatal exposures to O_3 21 have been studied. Maternal exposure to 0.4-1.2 ppm O₃ during gestation resulted in 22 developmental health effects in the RT of mice (Sharkhuu et al., 2011). Recent studies 23 involving postnatal exposure to O_3 have focused on differences between developing and 24 adult animals in antioxidant defenses, respiratory physiology and sensitivity to cellular 25 injury, and on mechanisms, such as lung structural changes, antigen sensitization, 26 interaction with nitric oxide signaling, altered airway afferent innervation and loss of 27 alveolar repair capacity, by which early O_3 exposure could lead to asthma pathogenesis or 28 exacerbations in later life (Auten and Foster, 2011).
- 29 An interesting set of studies conducted over the last 10 years in the infant rhesus monkey 30 has identified numerous O₃-mediated perturbations in the developing lung and immune 31 system (Plopper et al., 2007). These investigations were prompted by the dramatic rise in 32 the incidence of childhood asthma and focused on the possible interaction of O₃ and 33 allergens in promoting remodeling of the epithelial-mesenchymal trophic unit during 34 postnatal development of the tracheobronchial airway wall. In humans, airways growth 35 during the 8-12 year period of postnatal development is not well understood. Rhesus 36 monkeys were used in these studies because the branching pattern and distribution of 37 airways in this model are more similar to humans than those of rodents are to humans. In

- 1 addition, a model of allergic airways disease, which exhibits the main features of human 2 asthma, had already been established in the adult rhesus monkey. Studies in infant 3 monkeys were designed to determine whether repeated exposure to O_3 altered postnatal 4 growth and development, and if so, whether such effects were reversible. In addition, 5 exposure to O_3 was evaluated for its potential to increase the development of allergic 6 airways disease. Exposures were to cyclic episodic O_3 over 5 months which involved 5 7 biweekly cycles of alternating filtered air and O_3 - 9 consecutive days of filtered air and 5 8 consecutive days of 0.5 ppm O_3 , 8 h/day – and to house dust mite allergen (HDMA) for 9 2 hours per day for 3 days on the last 3 days of O_3 exposure followed by 11 days of 10 filtered air.
- 11Key findings were numerous. First, baseline airway resistance and AHR in the infant12monkeys were dramatically increased by combined exposure to both HDMA and O313(Joad et al., 2006; Schelegle et al., 2003). Secondly, O3 exposure led to a large increase in14BAL eosinophils (Schelegle et al., 2003) while HDMA exposure led to a large increase of15eosinophils in airways tissue (Joad et al., 2006; Schelegle et al., 2003). Thirdly, the16growth pattern of distal airways was changed to a large extent by exposure to O3 alone
- 17 and in combination with HDMA. More specifically, longer and narrower airways resulted 18 and the number of conducting airway generations between the trachea and the gas 19 exchange area was decreased (Fanucchi et al., 2006). This latter effect was not 20 ameliorated by a recovery period of 6 months. Fourthly, exposure to both HDMA and O_3 21 altered the abundance and distribution of CD25+ lymphocytes in the airways (Miller et 22 al., 2009). Lastly, several effects were seen at the level of the epithelial mesenchymal 23 trophic unit in response to O₃. These include altered organization of the airways 24 epithelium (Schelegle et al., 2003), increased abundance of mucous goblet cells 25 (Schelegle et al., 2003), disruption of the basement membrane zone (Evans et al., 2003), 26 reduced innervation (Larson et al., 2004), increased neuroendocrine-like cells (Joad et al., 27 2006), and altered orientation and abundance of smooth muscle bundles (Plopper et al., 28 2007; Tran et al., 2004). Six months of recovery in filtered air led to reversal of some but 29 not all of these effects (Kajekar et al., 2007; Plopper et al., 2007; Evans et al., 2004). The 30 authors concluded that cyclic challenge of infant rhesus monkeys to allergen and O₃ 31 during the postnatal period compromised airway growth and development and resulted in 32 changes which favor allergic airways responses and persistent effects on the immune 33 system (Plopper et al., 2007). A more recent study in this same model reported that early 34
 - life exposure to O_3 resulted in decreased total peripheral blood leukocyte numbers and increased blood eosinophils as well as persistent effects on pulmonary and systemic innate immunity in the infant rhesus monkey model (Maniar-Hew et al., 2011).
- Furthermore, the effect of cyclic episodic O₃ exposure on nasal airways was studied in
 the infant rhesus monkey model. The three-dimensional detail of the nasal passages was

35

1	analyzed for developing predictive dosimetry models and exposure-dose-response
2	relationships (Carey et al., 2007). The authors reported that the relative amounts of the
3	five epithelial cell types in the nasal airways of monkeys remained consistent between
4	infancy and adulthood [comparing to ($\underline{\text{Gross}}$, 1987; $\underline{\text{Gross}}$, 1982)]. Cyclic episodic O ₃
5	exposure (as described in the previous paragraphs) resulted in 50-80% decreases in
6	epithelial thickness and epithelial cell volume of the ciliated respiratory and transitional
7	epithelium, confirming that these cell types in the nasal cavity were the most sensitive to
8	O3 exposure. The character and location of nasal lesions resulting from O3 exposure were
9	similar in the infant monkeys and adult monkeys similarly exposed. However, the nasal
10	epithelium of infant monkeys did not undergo nasal airway epithelial remodeling or
11	adaptation which occurs in adult animals following O ₃ -mediated injury and which may
12	protect against subsequent O ₃ challenge. The authors suggested that infant monkeys may
13	be prone to developing persistent necrotizing rhinitis following episodic longer-term
14	exposures.

5.4.2.5 Attenuation of Responses

15	Repeated daily exposure to O_3 often results in a reduction in the degree of a response,
16	i.e., an attenuation of response. This phenomenon may reflect compensatory mechanisms
17	and is not necessarily beneficial. Furthermore, there is variability among the different
18	O ₃ -related endpoints in terms of response attenuation, as will be described below. As a
19	result, attenuation of some responses occurs concomitantly with the exacerbation of
20	others.
21	In responsive individuals, a striking degree of attenuation of the FEV ₁ response occurred
22	following repeated daily exposures to O ₃ . Generally, the young O ₃ responder was no
23	longer responsive on the fourth or fifth day of consecutive daily O3 exposure
24	(200-500 ppb O_3 for 2-4 hours with light to heavy levels of exercise) and required days to
25	weeks of non-exposure in order for the subject to regain O_3 responsiveness (Christian et
26	<u>al., 1998; Devlin et al., 1997; Linn et al., 1982b; Horvath et al., 1981; Hackney et al.,</u>
27	<u>1977b</u>). This phenomena has been reported for both lung function and symptoms such as
28	upper airway irritation, nonproductive cough, substernal discomfort and pain upon deep
29	inspiration (Linn et al., 1982b; Horvath et al., 1981; Hackney et al., 1977b). Repeated
30	daily exposures also led to an attenuation of the sRaw response in moderately exercising
31	human subjects exposed for 4 hours to 200 ppb O_3 (Christian et al., 1998) and to a
32	dampened AHR response compared with a single day exposure in light exercising human
33	subjects exposed for 2 hours to 400 ppb O_3 (Dimeo et al., 1981). However, one group
34	reported persistent small airway dysfunction despite attenuation of the FEV_1 response on

1 2	the third day of consecutive O_3 exposure (250 ppb, 2 hours, with moderate exercise) (Frank et al., 2001).
3	Studies in rodents also indicated an attenuation of the physiologic response measured by
4	breathing patterns and tidal volume following five consecutive days of exposure to
5	0.35-1 ppm O_3 for 2.25 hours (Tepper et al., 1989). Attenuation of O_3 -induced
6	bradycardic responses, which also result from activation of neural reflexes, has been
7	reported in rodents (0.5-0.6 ppm O ₃ , 2-6 h/day, 3-5 days (Hamade and Tankersley, 2009;
8	Watkinson et al., 2001).
9	Multi-day exposure to O_3 has been found to decrease some markers of inflammation
10	compared with a single day exposure (Christian et al., 1998; Devlin et al., 1997). For
11	example, in human subjects exposed for 4 hours to 200 ppb O_3 during moderate exercise,
12	decreased numbers of BAL neutrophils and decreased levels of BALF fibronectin and
13	IL-6 were observed after 4 days of consecutive exposure compared with responses after
14	1 day (Christian et al., 1998). Results indicated an attenuation of the inflammatory
15	response in both proximal airways and distal lung. However markers of injury, such as
16	lactate dehydrogenase (LDH) and protein in the BALF, were not attenuated in this study
17	(<u>Christian et al., 1998</u>). Other investigators found that repeated O_3 exposure (200 ppb O_3)
18	for 4 hours on 4 consecutive days with light exercise) resulted in increased numbers of
19	neutrophils in bronchial mucosal biopsies despite decreased BAL neutrophilia (Jorres et
20	al., 2000). Other markers of inflammation, including BALF protein and IL-6 remained
21	elevated following the multi-day exposure (Jorres et al., 2000).
22	In rats, the increases in BALF levels of proteins, fibronectin, IL-6 and inflammatory cells
23	observed after one day of exposure to 0.4 ppm O_3 for 12 hours were no longer observed
24	after 5 consecutive days of exposure (Van Bree et al., 2002). A separate study in rats
25	exposed to 0.35-1 ppm O_3 for 2.25 hours for 5 consecutive days demonstrated a lack of
26	attenuation of the increase in BALF protein, persistence of macrophages in the
27	centriacinar region and histological evidence of progressive tissue injury (Tepper et al.,
28	1989). Findings that injury, measured by BALF markers or by histopathology, persist in
29	the absence of BAL neutrophila or pulmonary function decrements suggested that
30	repeated exposure to O ₃ may have serious long-term consequences such as airway
31	remodeling. In particular, the small airways were identified as a site where cumulative
32	injury may occur (Frank et al., 2001).
33	Some studies examined the recovery of responses which were attenuated by repeated O ₃
34	exposure. In a study of humans undergoing heavy exercise who were exposed for 2 hours
35	to 400 ppb O_3 for five consecutive days (<u>Devlin et al., 1997</u>), recovery of the
36	inflammatory responses which were diminished by repeated exposure required
37	10-20 days following the exposure (Devlin et al., 1997). In an animal study conducted in

1	parallel (Van Bree et al., 2002), full susceptibility to O_3 challenge following exposure to
2	O_3 for five consecutive days required 15-20 days recovery.
3	Several mechanisms have been postulated to explain the attenuation of some responses
4	observed in human subjects and animal models following repeated exposure to O ₃ . First,
5	the upregulation of antioxidant defenses (or conversely, a decrease in critical O ₃ -reactive
6	substrates) may protect against O3-mediated effects. Increases in antioxidant content of
7	the BALF have been demonstrated in rats exposed to 0.25 and 0.5 ppm O_3 for
8	several hours on consecutive days (Devlin et al., 1997; Wiester et al., 1996b; Tepper et
9	al., 1989). Second, IL-6 was demonstrated to be an important mediator of attenuation in
10	rats exposed to 0.5 ppm for 4 hours on two consecutive days (Mckinney et al., 1998).
11	Third, a protective role for increases in mucus producing cells and mucus concentrations
12	in the airways has been proposed (Devlin et al., 1997). Fourth, epithelial hyperplasia or
13	metaplasia may decrease further effects due to subsequent O_3 challenge (Carey et al.,
14	2007; Harkema et al., 1987a; Harkema et al., 1987b). These morphologic changes have
15	been observed in nasal and lower airways in monkeys exposed chronically to
16	0.15-0.5 ppm O ₃ and reflect a persistent change in epithelial architecture which may lead
17	to other long-term sequelae. Although there is some evidence to support these
18	possibilities, there is no consensus on mechanisms underlying response attenuation.
19	Recent studies demonstrating that O3 activates TRP receptors suggest that modulation of
20	TRP receptor number or sensitivity by repeated O ₃ exposures may also contribute to the
21	attenuation of responses.

22In summary, the attenuation of pulmonary function responses by repeated exposure to O323has been linked to exacerbation of O3-mediated injury. Enhanced exposure to O3 due to a24dampening of the O3-mediated truncation of inspiration may be one factor which25contributes to this relationship.

5.4.2.6 Co-exposures with Particulate Matter

	1
27 the prevalence of these pollutants in ambient air. Results are highly variable and depe	nd
28 on whether exposures are simultaneous or sequential, the type of PM employed and the	ie
29 endpoint examined. Additive and interactive effects have been demonstrated. For	
30 example, simultaneous exposure to O_3 (120 ppb for 2 hours at rest) and concentrated	
31 ambient particles (CAPs) in human subjects resulted in a diminished systemic IL-6	
32 response compared with exposure to CAPs alone (<u>Urch et al., 2010</u>). However, expos	ure
33 to O_3 alone did not alter blood IL-6 levels (<u>Urch et al., 2010</u>). The authors provided	
34 evidence that O ₃ mediated a switch to shallow breathing which may have accounted f	or

1	the observed antagonism (Urch et al., 2010). Further, simultaneous exposure to O_3
2	(114 ppb for 2 hours at rest) and CAPs but not exposure to either alone, resulted in
3	increased diastolic blood pressure in human subjects (Fakhri et al., 2009). Mechanisms
4	underlying this potentiation of response were not explored. In some strains of mice,
5	pre-exposure to O ₃ (0.5 ppm for 2 hours) modulated the effects of carbon black PM on
6	heart rate, HRV and breathing patterns (Hamade and Tankersley, 2009). Another recent
7	study in mice demonstrated that treatment with carbon nanotubes followed 12 hours later
8	by O ₃ exposure (0.5 ppm for 3 hours) resulted in a dampening of some of the pulmonary
9	effects of carbon nanotubes measured as markers of inflammation and injury in the
10	BALF (Han et al., 2008). Further, Harkema and Wagner (2005) found that epithelial and
11	inflammatory responses in the airways of rats were enhanced by co-exposure to O ₃
12	(0.5 ppm for 3 days) and LPS (used as a model of biogenic PM) or to O_3 (1 ppm for
13	2 days) and OVA (used as a model of an aeroallergen). Lastly, a recent study
14	demonstrated that maternal exposure to particulate matter (PM) resulted in augmented
15	lung inflammation, airway epithelial mucous metaplasia and AHR in young mice
16	exposed chronically and intermittently to 1 ppm O_3 (Auten et al., 2009).
17	In summary, many of the demonstrated responses to co-exposure were more than

additive. These findings are hard to interpret but demonstrate the complexity of responses
 following combined exposure to PM and O₃.

5.4.2.7 Summary

20	Collectively, these earlier and more recent studies provide some evidence for
21	mechanisms that may underlie the variability in responsiveness seen among individuals
22	(Figure 5-9). Certain functional genetic polymorphisms, pre-existing conditions and
23	diseases, nutritional status, lifestage and co-exposures contribute to altered risk of
24	O ₃ -induced effects. Attenuation of responses may also be important, but it is
25	incompletely understood, both in terms of the pathways involved and the resulting
26	consequences.

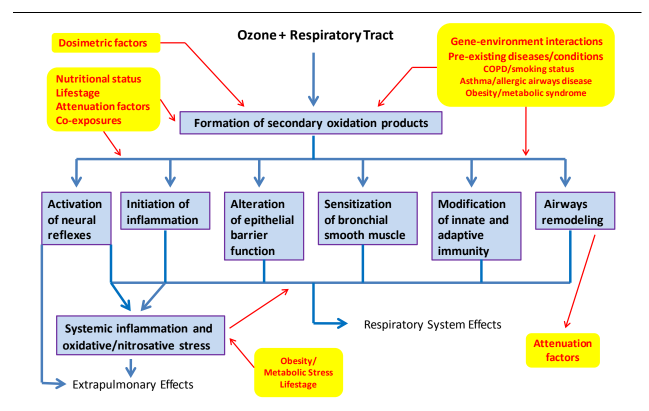


Figure 5-9 Some factors, illustrated in yellow, that likely contribute to the interindividual variability in responses resulting from inhalation of ozone.

5.5 Species Homology and Interspecies Sensitivity

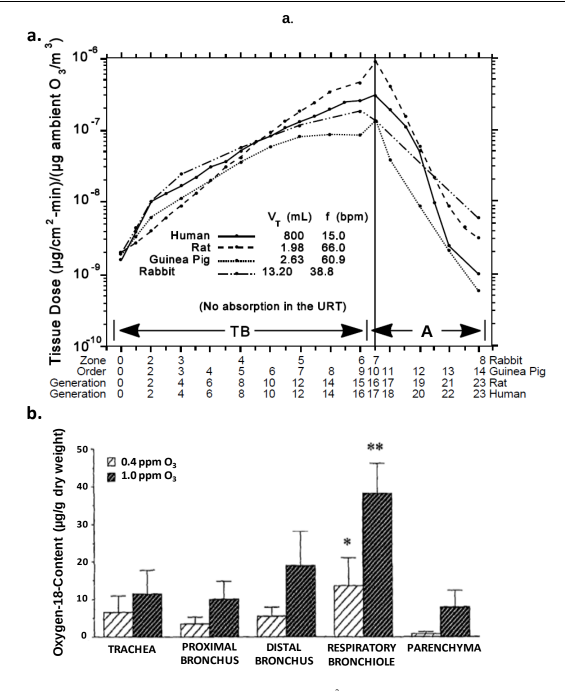
1	The previous O ₃ AQCDs discussed the suitability of animal models for comparison with
2	human O ₃ exposure and concluded that the acute and chronic functional responses of
3	laboratory animals to O_3 appear qualitatively homologous to human responses. Thus,
4	animal studies can provide important data in determining cause-effect relationships
5	between exposure and health outcome that would be impossible to collect in human
6	studies. Furthermore, animal studies add to a better understanding of the full range of
7	potential O ₃ -mediated effects.
8	Still, care must be taken when comparing quantitative dose-response relationships in
9	animal models to humans due to obvious interspecies differences. This section will
10	qualitatively describe basic concepts in species homology concerning both dose and
11	response to O_3 exposure. Overall, there have been few new publications examining
12	interspecies differences in dosimetry and response to O ₃ since the last AQCD. These
13	studies do not overtly change the conclusions discussed in the previous document.

5.5.1 Interspecies Dosimetry

1 2 3 4 5 6	As discussed in Section 5.2.1, O_3 uptake depends on complex interactions between RT morphology, breathing route, rate, and depth, physicochemical properties of the gas, physical processes of gas transport, as well as the physical and chemical properties of the ELF and tissue layers. Understanding differences in these variables between humans and experimental animals is important to interpreting delivered doses in animal and human toxicology studies.
7	Physiological and anatomical differences exist between experimental species. The
8	structure of the URT is vastly different between rodents and humans but scales according
9	to body mass. The difference in the cross-sectional shape and size of the nasal passages
10	affects bulk airflow patterns, particularly the shape of major airflow streams. The nasal
11	epithelium is lined by squamous, respiratory, or olfactory cells, depending on location.
12	The differences in airflow patterns in the URT mean that not all nasal surfaces and cell
13	types receive the same exposure to inhaled O ₃ leading to differences in local absorption
14	and potential for site-specific tissue damage. The morphology of the LRT also varies
15	within and among species. Rats and mice do not possess respiratory bronchioles;
16	however, these structures are present in humans, dogs, ferrets, cats, and monkeys.
17	Respiratory bronchioles are abbreviated in hamsters, guinea pigs, sheep, and pigs. The
18	branching structure of the ciliated bronchi and bronchioles also differs between species
19	from being a rather symmetric and dichotomous branching network of airways in humans
20	and primates to a more monopodial branching network in other mammals. In addition,
21	rodents have fewer terminal bronchioles due to a smaller lung size compared to humans
22	or canines (McBride, 1992). The cellular composition in the pulmonary region is similar
23	across mammalian species; at least 95% of the alveolar epithelial tissue is composed of
24	Type I cells. However, considerable differences exist between species in the number and
25	type of cells in the TB airways. Differences also exist in breathing route and rate.
26	Primates are oronasal breathers, while rodents are obligate nasal breathers. Past studies of
27	the effect of body size on resting oxygen consumption also suggest that rodents inhale
28	more volume of air per lung mass than primates. These distinctions as well as differences
29	in nasal structure between primates and rodents affect the amount of O_3 uptake.
30	As O_3 absorption and reactivity relies on ELF antioxidant substances (see Section 5.2.3),
31	variability in antioxidant concentrations and metabolism between species may affect dose
32	and O ₃ -induced health outcomes. The thickness of the ELF in the TB airways varies
33	among species. Mercer et al. (1992) found that the human ELF thickness in bronchi and
34	bronchioles was 6.9 and 1.8 μ m, respectively, compared to 2.6 and 1.9 μ m for the same
35	locations in the rat. Guinea pigs and mice have a lower basal activity of GSH transferase
36	and GSH peroxidase, and lower α -TOH levels in the lung compared to rats (<u>Ichinose et</u>

1	al., 1988; Sagai et al., 1987). Nasal lavage fluid analysis shows that humans have a higher
2	proportion of their nasal antioxidants as UA and low levels of AH ₂ whereas mice, rats, or
3	guinea pigs have high levels of AH_2 and undetectable levels of UA. GSH is not detected
4	in the nasal lavage of most of these species, but is present in monkey nasal lavage.
5	Guinea pigs and rats have a higher antioxidant to protein ratio in nasal lavage and BALF
6	than humans (Hatch, 1992). The BALF profile differs from the nasal lavage. Humans
7	have a higher proportion of GSH and less AH ₂ making up their BALF content compared
8	to the guinea pigs and rats (Slade et al., 1993; Hatch, 1992). Similar to the nose, rats have
9	the highest antioxidant to protein mass ratio found in BALF (Slade et al., 1993).
10	Antioxidant defenses also vary with age (Servais et al., 2005) and exposure history (Duan
11	et al., 1996). Duan et al. (1996); Duan et al. (1993) reported that differences in
12	antioxidant levels between species and lung regions did not appear to be the primary
13	factor in O_3 induced tissue injury. However, a close correlation between site-specific O_3
14	dose, the degree of epithelial injury, and reduced glutathione depletion was observed in
15	monkeys (<u>Plopper et al., 1998</u>).
16	Even with these differences humans and animals are similar in the pattern of regional O_3
17	dose distribution. As discussed for humans in Section <u>5.2.2</u> , O_3 flux to the air-liquid
18	interface of the ELF slowly decreases distally in the TB region and then rapidly decreases

interface of the ELF slowly decreases distally in the TB region and then rapidly decreases 18 19 distally in the alveolar region (Miller et al., 1985). Modeled tissue dose in the human RT, 20 representing O_3 flux to the liquid-tissue interface, is very low in the trachea, increases to 21 a maximum in the CAR, and then rapidly decreases distally in the alveolar region 22 (Figure 5-10). Similar patterns of O_3 tissue dose profiles normalized to inhaled O_3 23 concentration were predicted for rat, guinea pig, and rabbit (Miller et al., 1988; Overton 24 et al., 1987) (Figure 5-10a). Overton et al. (1987) modeled rat and guinea pig O₃ dose 25 distribution and found that after comparing two different morphometrically based 26 anatomical models for each species, considerable difference in predicted percent RT and 27 alveolar region uptakes were observed. This was due to the variability between the two 28 anatomical models in airway path distance to the first alveolated duct. As a result, the 29 overall dose profile was similar between species however the O₃ uptake efficiency varied due to RT size and path length (Section 5.2.2). A similar pattern of O_3 dose distribution 30 was measured in monkeys exposed to 0.4 and 1.0 ppm ${}^{18}O_3$ (Plopper et al., 1998) 31 (Figure 5-10b). Less ¹⁸O was measured in the trachea, proximal bronchus, and distal 32 33 bronchus than was observed in the respiratory bronchioles. Again indicating the highest 34 concentration of O₃ tissue dose is localized to the CAR, which are the respiratory bronchioles in nonhuman primates. In addition, the lowest ¹⁸O detected in the RT was in 35 the parenchyma (i.e., alveolar region), mimicking the rapid decrease in tissue O_3 dose 36 37 predicted by models for the alveolar regions of humans and other animals.

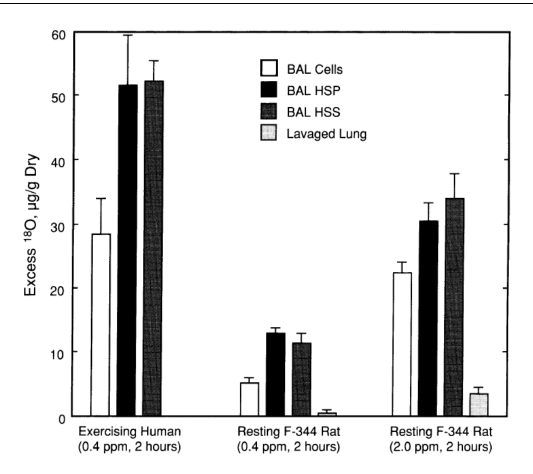


Note: Panel (a) presents the predicted tissue dose of O_3 (as μ g of O_3 per cm² of segment surface area per min, standardized to a tracheal O_3 value of 1 μ g/m³) for various regions of the rabbit, guinea pig, rat, and human RT. TB = tracheobronchial region, A = alveolar region. Panel (b) presents a comparison of excess ¹⁸O in the five regions of the TB airways of rhesus monkeys exposed to O_3 for 2h. *p <0.05 comparing the same O_3 concentration across regions. **p <0.05 comparing different O_3 concentrations in the same region.

Source: Panel (a) U.S. EPA (1996a) (b) Plopper et al. (1998)

Figure 5-10 Humans and animals are similar in the regional pattern of ozone tissue dose distribution.

1	Humans and animal models are similar in the pattern of regional O ₃ dose, but absolute
2	values differ. Hatch et al. (1994) reported that exercising humans exposed to oxygen-18
3	labeled O ₃ (400 ppb) accumulated 4-5 times higher concentrations of O ₃ reaction product
4	in BAL cells, surfactant and protein fractions compared to resting rats similarly exposed
5	(400 ppb) (Figure 5-11). The use of 18 O was specifically employed in an attempt to
6	accurately measure O_3 dose to BALF fractions and lung tissue and was normalized to the
7	dried mass of lavaged constituents. It was necessary to expose resting rats to 2 ppm O_3 to
8	achieve the same BALF accumulation of ¹⁸ O reaction product that was observed in
9	humans exposed to 400 ppb with intermittent heavy exercise ($\dot{V}_E \sim 60$ L/min). The
10	concentration of ¹⁸ O reaction product in BALF paralleled the accumulation of BALF
11	protein and cellular effects of the O3 exposure observed such that these responses to
12	$2.0 \text{ ppm } O_3$ were similar to those of the 400 ppb O_3 in exercising humans. This suggests
13	that animal data obtained in resting conditions would underestimate the reaction of O_3
14	with cells in the RT and presumably the resultant risk of effect for humans. However
15	these results should be interpreted with caution given an important limitation in the ¹⁸ O
16	labeling technique when used for interspecies comparisons. The reaction between O_3 and
17	some reactants such as ascorbate produce ¹⁸ O-labeled products that are lost during sample
18	processing. When levels of ascorbate or other such reactants vary between species, this
19	lost portion of the total ¹⁸ O-reaction products will also vary, leading to uncertainty in
20	interspecies comparisons.



Note: The excess ¹⁸O in each fraction is expressed relative to the dry weight of that fraction. Fractions assayed include cells, high speed pellet (HSP), high speed supernatant (HSS), and lavaged lung homogenates. Source: <u>Hatch et al. (1994</u>)

Figure 5-11 Oxygen-18 incorporation into different fractions of BALF from humans and rats exposed to 0.4 and 2.0 ppm ¹⁸O₃.

1	Recently, a quantitative comparison of O_3 transport in the airways of rats, dogs, and
2	humans was conducted using a three-compartment airways model, based on upper and
3	lower airway casts and mathematical calculation for alveolar parameters (Tsujino et al.,
4	2005). This one-dimensional gas transport model examined how interspecies anatomical
5	and physiological differences affect intra-airway O_3 concentrations and the amount of gas
6	absorbed. The morphological model consisted of cylindrical tubes with constant volume
7	and no airway branching patterns. Peak, real-time, and mean O_3 concentrations were
8	higher in the upper and lower airways of humans compared to rats and dogs, but lowest
9	in the alveoli of humans. The amount of O_3 absorbed was lowest in humans when
10	normalized by body weight. The intra-airway concentration decreased distally in all
11	species. Sensitivity analysis demonstrated that V_T , f_B , and upper and lower airways

1	surface area had a statistically significant impact on model results. The model is limited
2	in that it did not account for chemical reactions in the ELF or consider gas diffusion as a
3	driving force for O ₃ transport. Also, the model was run at a respiratory rate of 16/min
4	simulating a resting individual, however exercise may cause a further deviation from
5	animal models as was seen in <u>Hatch et al. (1994</u>).
6	Overall, animal models exhibit qualitatively similar patterns of O ₃ net and tissue dose
7	distribution with the largest tissue dose delivered to the CAR. However, due to
8	anatomical and biochemical RT differences the absolute values of O3 dose delivered
9	differs. Past results suggest that animal data obtained in resting conditions would
10	underestimate the O_3 reactions with cells in the BALF and presumably the resultant risk
11	of effect for humans, especially for humans during exercise.

5.5.2 Interspecies Homology of Response

12Biological response to O3 exposure broadly shows commonalities in many species.13Among rodents, non-human primates, and humans, for example, ample data suggest that14O3 induces oxidative stress, cell injury, upregulation of cytokines/chemokines,15inflammation, alterations in lung function, and disruption of normal lung growth and16development (See Chapters 6 and 7).

- 17 The effects related to early life exposures can differ appreciably across species due to the 18 maturation stage of the lung and immune systems at birth. Evidence from non-human 19 primate studies shows that early life O_3 exposure disrupts lung development producing 20 physiologic perturbations that are similar to those observed in children exposed to urban 21 air pollution (Fanucchi et al., 2006; Joad et al., 2006). Studies of O₃ effects on lung 22 surface chemistry also show some degree of homology. Lipid oxidation products specific 23 to O₃ reactions with unsaturated fatty acids have been reported, for example, in lavage 24 fluids from both rodents and humans (Frampton et al., 1999; Pryor et al., 1996). In 25 humans, the extent to which systemic effects occur is less well studied; plasma indices of 26 lipid oxidation such as isoprostanes unfortunately do not pinpoint the compartment(s) 27 where oxidative stress has transpired. That oxidative stress occurs systemically in both 28 rodents and non-human primates (Chuang et al., 2009), nevertheless, suggests that it 29 likely also occurs in humans.
- 30Despite the overall similarities in responses to O_3 among species, studies have reported31variability in the responsiveness to O_3 between and within species, as well as between32endpoints. Rodents appear to have a slightly higher tachypneic response to O_3 and are33less sensitive to changes in pulmonary function responses than humans (U.S. EPA,341996a). However, rats experience attenuation of pulmonary function and tachypneic

1	ventilatory responses, similar to humans (Wiester et al., 1996b). Hatch et al. (1986)
2	reported that guinea pigs were the most responsive to O_3 -induced inflammatory cell and
3	protein influx. Rabbits were the least responsive and rats, hamsters, and mice were
4	intermediate responders. Further analysis of this study by <u>Miller et al. (1988)</u> found that
5	the protein levels in BALF from guinea pigs increased more rapidly with predicted
6	pulmonary tissue dose than in rats and rabbits. Alveolar macrophages isolated from
7	guinea pigs and humans mounted similar qualitative and quantitative cytokine responses
8	to in vitro O_3 (0.1-1.0 ppm for 60 minutes) exposure (<u>Arsalane et al., 1995</u>).
0	to in vito O_3 (0.1-1.0 ppin for of minutes) exposure (<u>Arsalane et al., 1995</u>).
9	Also, because of their higher body surface to volume ratio, rodents can rapidly lower
10	body temperature during exposure leading to lowered O_3 dose and toxicity (Watkinson et
11	<u>al., 2003;</u> <u>Iwasaki et al., 1998;</u> <u>Slade et al., 1997</u>). In addition to lowering the O_3 dose to
12	the lungs, this hypothermic response may cause: (1) lower metabolic rate, (2) altered
13	enzyme kinetics, and (3) altered membrane function. The thermoregulatory mechanisms
14	also may affect disruption of heart rate that may lead to: (1) decreased cardiac output, (2)
15	lowered blood pressure, and (3) decreased tissue perfusion (Watkinson et al., 2003).
16	These responses have not been observed in humans except at very high exposures, thus
17	further complicating extrapolation of effects from animals to humans.
18	The degree to which O_3 induces injury and inflammation responses appears to be variable
19	between species. However, the majority of those studies did not normalize the response
20	to the dose received to account for species differences in O_3 absorption. For example,
21	Dormans et al. (1999) found that rats, mice, and guinea pigs all exhibited O_3 -induced (0.2)
22	- 0.4 ppm for 3-56 days) inflammation; however, guinea pigs were the most responsive
23	with respect to alveolar macrophage elicitation and pulmonary cell density in the
24	centriacinar region. Mice were the most responsive in terms of bronchiolar epithelial
25	hypertrophy and biochemical changes (e.g., LDH, glutathione reductase, glucose-
26	6-phosphate dehydrogenase activity), and had the slowest recovery from O_3 exposure. All
27	species displayed increased collagen in the ductal septa and large lamellar bodies in Type
28	If pneumocytes at the longest exposure and highest concentration; whereas this response
29	occurred in the rat and guinea pig at lower O_3 levels (0.2 ppm) as well. Overall, the
30	authors rated mice as most responsive, followed by guinea pigs, then rats (Dormans et al.,
31	1999). Rats were also less responsive in terms of epithelial necrosis and inflammatory
32	responses as a result of O_3 exposure (1.0 ppm for 8 hours) compared with monkeys and
33	ferrets, which manifested a similar response (<u>Sterner-Kock et al., 2000</u>). Results of this
34	study should be interpreted with caution since no dose metric was used to normalize the
35	total inhaled dose or local organ dose between species.
36	To further understand the genetic basis for age-dependent differential response to O_3 ,
37	adult (15 week old) and neonatal (15-16 day old) mice from 8 genetically diverse strains

1	were examined for O_3 -induced (0.8 ppm for 5 hours) pulmonary injury and lung
2	inflammation (Vancza et al., 2009). Ozone exposure increased polymorphonuclear
3	leukocytes (PMN) influx in all strains of neonatal mice tested, but significantly greater
4	PMNs occurred in neonatal compared to adult mice for only some sensitive strains,
5	suggesting a genetic background effect. This strain difference was not due to differences
6	in delivered dose of O_3 to the lung, evidenced by ¹⁸ O lung enrichment. The sensitivity of
7	strains for O ₃ -induced increases in BALF protein and PMNs was different for different
8	strains of mice suggesting that genetic factors contributed to heightened responses.
9	Interestingly, adult mice accumulated more than twice the levels of ¹⁸ O reaction product
10	of O_3 than corresponding strain neonates. Thus, it appeared that the infant mice showed a
11	2-fold- to 3-fold higher response than the adults when expressed relative to the
12	accumulated O_3 reaction product in their lungs. The apparent decrease in delivered O_3
13	dose in neonates could be a result of a more rapid loss of body temperature in infant
14	rodents incident to maternal separation and chamber air flow.
15	In animal studies, inhaled O ₃ concentration and exposure history rarely reflect actual
16	human environmental exposures. Generally, very high exposure concentrations are used
17	to induce murine AHR, which in some human subjects is observed at far more relevant
18	concentrations. This calls into question whether the differences in airway reactivity are
19	simply a function of differential nasopharyngeal scrubbing or whether the complexities
20	encompassing a variety of contributory biological pathways show species divergence.
21	Furthermore, in non-human primates exposed during early life, eosinophil trafficking
22	occurs, which has not been observed in rodents (unless sensitized) (Maniar-Hew et al.,
23	<u>2011</u>). This response has been shown to be persistent when O_3 challenges are
24	administered after a recovery period of ≥ 9 months during which no exposure transpired.
25	Quantitative extrapolation is challenging due to a number of uncertainties. Unfortunately,
26	many input parameters needed to conduct quantitative extrapolations across species have
27	not been obtained or currently remain undefined. It is not clear whether characterization
28	of the ELF provides the information needed to compute a profile of reaction products or
29	whether environmentally relevant exposure has altered the physicochemical interactions
30	that occur within the RT surface compartment (e.g., O ₃ diffusion through regions where
31	the ELF is thin). That systemic effects have been documented in both rodents and non-
32	human primates leads to the question of whether reaction products,
33	cytokines/chemokines, or both enter the nasopharyngeal or bronchial circulation, both of
34	which show species-dependent differences (Chuang et al., 2009; Cole and Freeman,
35	<u>2009</u>).
36	In addition, the response to O ₃ insult across species and more recent health effects such as
~-	a second s

37 immune system development are uncertain. Non-human primate studies have shown

1 hypo-responsiveness to endotoxin challenge as a consequence of exposure; whether this 2 occurs in rodents and humans is largely unknown (Maniar-Hew et al., 2011). In addition, 3 structural changes (e.g., airways remodeling, fibrogenesis) might differ appreciably 4 across species. Moreover, whether the upper airways differentially contribute to either 5 distal lung or systemic impacts has not been explored. 6 Some outcomes (e.g., inflammation) support the conclusion of homologous responses 7 across species. However, factors such as age, exposure history, diet, endogenous 8 substrate generation and homeostatic regulation, the cellular machinery that regulates 9 inflammatory cell trafficking, responses to other environmental challenges, and the 10 precise chemical species (whether ELF or cell membrane-derived) that account for 11 exposure-related initiation of pathophysiologic sequelae might differ across species, but 12 the extent of species-specific contributing factors remains unknown. Consequently, some 13 level of uncertainty cannot be dismissed. Nonetheless, if experimental animals show 14 pathophysiological consequences of exposure, assuming that qualitatively similar human 15 health impacts could occur is not unreasonable.

5.5.3 Summary

16	In summary, biological response to O ₃ exposure broadly shows commonalities in many
17	species and thus supports the use of animal studies in determining mechanistic and cause-
18	effect relationships and as supporting evidence that similar effects could occur in humans
19	if O_3 exposure is sufficient. However, there is uncertainty regarding the similarity of
20	response to ozone across species for some recently described endpoints. Differences exist
21	between species in a number of factors that influence O ₃ dosimetry and responses, such
22	as RT anatomy, breathing patterns, and ELF antioxidant concentrations and chemical
23	species. While humans and animals are similar in the pattern of regional O_3 dose
24	distribution, these differences will likely result in differences in the absolute values of
25	O_3 dose delivered throughout the RT. These considerations limit quantitative comparison
26	between species.

5.6 Chapter Summary

27Ozone is a highly reactive gas and a powerful oxidant with a short half-life. Both O328uptake and responses are dependent upon the formation of secondary reaction products in29the ELF; however more complex interactions occur. Uptake in humans at rest is 80-95%30efficient and it is influenced by a number of factors including RT morphology, breathing31route, frequency, and volume, physicochemical properties of the gas, physical processes

- 1 of gas transport, as well as the physical and chemical properties of the ELF and tissue 2 layers. In fact, even though the average LRT dose may be at a level where health effects 3 would not be predicted, local regions of the RT may receive considerably higher than 4 average doses due to RT inhomogeneity and differences in the pathlengths, and therefore 5 be at greater risk of effects. The primary uptake site of O_3 delivery to the lung epithelium 6 is believed to be the CAR, however changes in a number of factors (e.g., physical 7 activity) can alter the distribution of O_3 uptake in the RT. Ozone uptake is chemical 8 reaction-dependent and the substances present in the ELF appear in most cases to limit 9 interaction of O_3 with underlying tissues and to prevent penetration of O_3 distally into the 10 RT. Still, reactions of O_3 with soluble ELF components or possibly plasma membranes 11 result in distinct products, some of which are highly reactive and can injure and/or 12 transmit signals to RT-cells.
- 13 Thus, in addition to contributing to the driving force for O_3 uptake, formation of 14 secondary oxidation products initiates pathways that provide the mechanistic basis for 15 health effects that are described in detail in Chapters 6 and 7 and that involve the RT as 16 well as extrapulmonary systems. These pathways include activation of neural reflexes, 17 initiation of inflammation, alteration of epithelial barrier function, sensitization of 18 bronchial smooth muscle, modification of innate and adaptive immunity, airways 19 remodeling, and systemic inflammation and oxidative/nitrosative stress. With the 20 exception of airways remodeling, these pathways have been demonstrated in both 21 animals and human subjects in response to the inhalation of O₃.
- 22 Both dosimetric and mechanistic factors contribute to the understanding of 23 interindividual variability in responses to O_3 . This variability is influenced by differences 24 in RT volume and surface area, certain genetic polymorphisms, pre-existing conditions 25 and disease, nutritional status, lifestages, attenuation, and co-exposures. Some of these 26 factors also underlie differences in species homology and sensitivity. Qualitatively, 27 animal models exhibit similar patterns of O₃ net and tissue dose distribution with the 28 largest tissue dose of O₃ delivered to the CAR. However, due to anatomical and 29 biochemical RT differences, the absolute value of delivered O_3 dose differs, with animal 30 data obtained in resting conditions underestimating the dose to the RT and presumably 31 the resultant risk of effect for humans, especially humans during exercise. Even though 32 interspecies differences limit quantitative comparison between species, many short-term 33 responses of laboratory animals to O_3 appear qualitatively homologous to those of the 34 human. Furthermore, animal studies add to a better understanding of the full range of 35 potential O₃-mediated effects. Given the commonalities in many responses across 36 species, animal studies that observe O_3 -induced effects may be used as supporting 37 evidence that similar effects could occur in humans or in determining mechanistic and 38 cause-effect relationships if O₃ exposure is sufficient.

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6 INTEGRATED HEALTH EFFECTS OF SHORT-TERM OZONE EXPOSURE

6.1 Introduction

1	This chapter reviews, summarizes, and integrates the evidence for various health
2	outcomes associated with short-term (i.e., hours, days, or weeks) exposures to O ₃ .
3	Numerous controlled human exposure, epidemiologic, and toxicological studies have
4	permitted evaluation of the relationships between short-term O_3 exposure and a range of
5	endpoints related to respiratory effects (Section 6.2), cardiovascular effects (Section 6.3),
6	and mortality (Section 6.2 , Section 6.3 , and Section 6.6). A smaller number of studies
7	were available to assess the effects of O3 exposure on other physiological systems such as
8	the central nervous system (Section 6.4), liver and metabolism (Section $6.5.1$), and
9	cutaneous and ocular tissues (Section $6.5.2$). This chapter evaluates the majority of recent
10	(i.e., published since the completion of the 2006 O ₃ AQCD) short-term exposure studies;
11	however, those for birth outcomes and infant mortality are evaluated in Chapter $\frac{7}{2}$
12	(Section 7.4), because they compare associations among overlapping short- and long-
13	term exposure windows that are difficult to distinguish.
14	Within each individual section of this chapter, a brief summary of conclusions from the
15	2006 O ₃ AQCD is included along with an evaluation of recent evidence that is intended
16	to build upon the body of evidence from previous reviews. The studies evaluated are
17	organized by health endpoint (e.g., lung function, pulmonary inflammation) then by
18	scientific discipline (e.g., controlled human exposure, epidemiology, and toxicology).
19	Each major section (e.g., respiratory, cardiovascular, mortality) concludes with an
20	integrated summary of the findings and a conclusion regarding causality based upon the
21	framework described in the Preamble to this ISA. The causal determinations are
22	presented for a broad health effect category, such as respiratory effects, with coherence
23	and plausibility based on the total evidence available across disciplines and across the
24	suite of related health endpoints, including cause-specific mortality.

6.2 Respiratory Effects

25	Based on evidence integrated across controlled human exposure, epidemiologic, and
26	toxicological studies, the 2006 O ₃ AQCD concluded "that acute O ₃ exposure is causally
27	associated with respiratory system effects" (U.S. EPA, 2006b). Contributing to this
28	conclusion were the consistency and coherence across scientific disciplines for the effects

1	of short-term O_3 exposure on a variety of respiratory outcomes including "pulmonary
2	function decrements, respiratory symptoms, lung inflammation, and increased lung
3	permeability, airway hyperresponsiveness." Collectively, these findings provided
4	biological plausibility for associations in epidemiologic studies observed between short-
5	term increases in ambient O3 concentration and increases in respiratory symptoms and
6	respiratory-related hospitalizations and emergency department (ED) visits.
7	Controlled human exposure studies have provided strong and quantifiable exposure-
8	response data on the human health effects of O ₃ . The most salient observations from
9	studies reviewed in the 1996 and 2006 O ₃ AQCDs (U.S. EPA, 2006b, 1996a) included:
10	(1) young healthy adults exposed to O_3 concentrations ≥ 80 ppb develop significant
11	reversible, transient decrements in pulmonary function and symptoms of breathing
12	discomfort if minute ventilation (\dot{V}_E) or duration of exposure is increased sufficiently;
13	(2) relative to young adults, children experience similar spirometric responses but lower
14	incidence of symptoms from O ₃ exposure; (3) relative to young adults, O ₃ -induced
15	spirometric responses are decreased in older individuals; (4) there is a large degree of
16	intersubject variability in physiologic and symptomatic responses to O_{3} , but responses
17	tend to be reproducible within a given individual over a period of several months; (5)
18	subjects exposed repeatedly to O3 for several days experience an attenuation of
19	spirometric and symptomatic responses on successive exposures, which is lost after about
20	a week without exposure; and (6) acute O_3 exposure initiates an inflammatory response
21	that may persist for at least 18 to 24 hours postexposure.
22	Substantial evidence for biologically plausible O ₃ -induced respiratory morbidity has been
23	derived from the coherence between toxicological and controlled human exposure study
24	findings for parallel endpoints. For example, O3-induced lung function decrements and
25	increased airway hyperresponsiveness have been observed in both animals and humans.
26	Airway hyperresponsiveness could be an important consequence of exposure to ambient
27	O_3 because the airways are then predisposed to narrowing upon inhalation of a variety of
28	ambient stimuli. Additional airway hyperresponsiveness tends to resolve more slowly and
29	appears less subject to attenuation with repeated exposures than lung function
30	decrements. Increased permeability and inflammation have been observed in the airways
31	of humans and animals alike after O ₃ exposure, although these processes are not
32	necessarily associated with immediate changes in lung function or hyperresponsiveness.
33	Furthermore, the potential relationship between repetitive bouts of acute inflammation
34	and the development of chronic respiratory disease is unknown. Another feature of
35	O ₃ -related respiratory morbidity is impaired host defense and reduced resistance to lung
36	infection, which has been strongly supported by toxicological evidence and, to a limited
37	extent, by evidence from controlled human exposure studies. Recurrent respiratory
38	infection in early life is associated with increased incidence of asthma in humans.

1	In concordance with experimental studies, epidemiologic studies have provided clear
2	evidence for decrements in lung function related to short-term ambient O3 exposure.
3	These effects were demonstrated in healthy children attending camps, adults exercising or
4	working outdoors, and children with and without asthma (U.S. EPA, 2006b, 1996a). In
5	addition to lung function decrements, short-term increases in ambient O ₃ concentration
6	were associated with increases in respiratory symptoms (e.g., cough, wheeze, shortness of
7	breath), notably in large U.S. panel studies of children with asthma (Gent et al., 2003;
8	Mortimer et al., 2000). The evidence across disciplines for O_3 effects on a range of
9	respiratory endpoints collectively provides support for epidemiologic studies that have
10	demonstrated consistent associations between short-term increases in ambient O ₃
11	concentration and increases in respiratory hospital admissions and ED visits, specifically
12	during the summer or warm months. In contrast with other respiratory health endpoints,
13	epidemiologic evidence did not clearly support a relationship between short-term O ₃
14	exposure and respiratory mortality. Although O3 was consistently associated with
15	nonaccidental and cardiopulmonary mortality, the contribution of respiratory causes to
16	these findings was uncertain as the few studies that examined mortality specifically from
17	respiratory causes reported inconsistent associations with ambient O ₃ concentrations.
18	As will be discussed throughout this section, consistent with the strong body of evidence
19	presented in the 2006 O ₃ AQCD, recent studies continue to support associations between
20	short-term O ₃ exposure and respiratory effects, in particular, lung function decrements in
21	controlled human exposure studies, airway inflammatory responses in toxicological
22	studies, and respiratory-related hospitalizations and ED visits. Recent epidemiologic
23	studies contribute new evidence for potentially at-risk populations and associations
24	linking ambient O ₃ concentrations with biological markers of airway inflammation and
25	oxidative stress, which is consistent with the extensive evidence from controlled human
26	exposure and toxicological studies. Furthermore, extending the potential range of
27	well-established O ₃ -associated respiratory effects, recent multicity studies and a

well-established O_3 -associated respiratory effects, recent multicity studies and a multicontinent study demonstrate associations between short-term increases in ambient O_3 concentration and respiratory-related mortality.

6.2.1 Lung Function

28

29

6.2.1.1 Controlled Human Exposure

30	This section focuses on studies examining O ₃ effects on lung function and respiratory
31	symptoms in volunteers exposed, for periods of up to 8 hours, to O_3 concentrations
32	ranging from 40 to 500 ppb, while at rest or during exercise of varying intensity.

1 2 3 4 5	Responses to acute O ₃ exposures in the range of ambient concentrations include decreased inspiratory capacity; mild bronchoconstriction; rapid, shallow breathing patterns during exercise; and symptoms of cough and pain on deep inspiration (PDI). Reflex inhibition of inspiration results in a decrease in forced vital capacity (FVC) and total lung capacity (TLC) and, in combination with mild bronchoconstriction, contributes
6 7 8 9 10 11 12 13	to a decrease in the forced expiratory volume in 1 second (FEV ₁). In studies that have exposed subjects during exercise, the majority of shorter duration (\leq 4-hour exposures) studies utilized an intermittent exercise protocol in which subjects rotated between 15-minute periods of exercise and rest. A limited number of 1- to 2-hour studies, mainly focusing on exercise performance, have utilized a continuous exercise regime. A quasi continuous exercise protocol is common to prolonged exposure studies where subjects complete 50-minute periods of exercise followed by 10-minute rest periods.
14 15 16 17 18 19 20 21	The majority of controlled human exposure studies have been conducted within exposure chambers, although a smaller number of studies used a facemask to expose subjects to O_3 . Little effort has been made herein to differentiate between facemask and chamber exposures since FEV ₁ and respiratory symptom responses appear minimally affected by these exposure modalities. Similar responses between facemask and chamber exposures have been reported for exposures to 80 and 120 ppb O_3 (6.6-hour, moderate quasi continuous exercise, 40 L/min) and 300 ppb O_3 (2 h, heavy intermittent exercise, 70 L/min) (Adams, 2003a, b, 2002).
22 23 24 25 26 27 28 29 30 31 32 33	The majority of controlled human exposure studies investigating the effects O_3 are of a randomized, controlled, crossover design in which subjects were exposed, without knowledge of the exposure condition and in random order to clean filtered air (FA; the control) and, depending on the study, to one or more O_3 concentrations. The FA control exposure provides an unbiased estimate of the effects of the experimental procedures on the outcome(s) of interest. Comparison of responses following this FA exposure to those following an O_3 exposure allows for estimation of the effects of O_3 itself on an outcome measurement while controlling for independent effects of the experimental procedures. As individuals may experience small changes in various health endpoints from exercise, diurnal variation, or other effects in addition to those of O_3 during the course of an exposure, the term " O_3 -induced" is used herein to designate effects that have been corrected or adjusted for such extraneous responses as measured during FA exposures.
34 35 36 37	Spirometry, viz., FEV ₁ , is a common health endpoint used to assess effects of O_3 on respiratory health in controlled human exposure studies. In considering 6.6-hour exposures to FA, group mean FEV ₁ changes have ranged from -0.7% (<u>McDonnell et al.</u> , 1991) to 2.7% (<u>Adams</u> , 2006a). On average, across ten 6.6-hour exposure studies, there

1	has been a 1.0% (n = 279) increase in FEV ₁ (Kim et al., 2011; Schelegle et al., 2009;
2	Adams, 2006a, 2003a, 2002; Adams and Ollison, 1997; Folinsbee et al., 1994;
3	McDonnell et al., 1991; Horstman et al., 1990; Folinsbee et al., 1988). Regardless of the
4	reason for small changes in FEV_1 over the course of FA exposures, whether biologically
5	based or a systematic effect of the experimental procedures, the use of FA responses as a
6	control for the assessment of responses following O_3 exposure in randomized exposure
7	studies serves to eliminate alternative explanations other than those of O_3 itself in causing
8	the measured responses.
9	Considering FEV ₁ responses in young healthy adults, an O ₃ -induced change in FEV ₁ is
10	typically the difference between the decrement observed with O ₃ exposure and the
11	improvement observed with FA exposure. Noting that some healthy individuals
12	experience small improvements while others have small decrements in FEV1 following
13	FA exposure, investigators have used the randomized, crossover design with each subject
14	serving as their own control (exposure to FA) to discern relatively small effects with
15	certainty since alternative explanations for these effects are controlled for by the nature of
16	the experimental design. The utility of intraindividual FA control exposures becomes
17	more apparent when considering individuals with respiratory disease. The occurrence of
18	exercise-induced bronchospasm is well recognized in patients with asthma and COPD
19	and may be experienced during both FA and O_3 exposures. Absent correction for FA
20	responses, exercise-induced changes in FEV_1 could be mistaken for responses due to O_3 .
21	This biological phenomenon serves as an example to emphasize the need for a proper
22	control exposure in assessing the effects of O_3 as well as the role of this control in
23	eliminating the influence of other factors on the outcomes of interest.

Pulmonary Function Effects of Ozone Exposure in Healthy Subjects

Acute Exposure of Healthy Subjects

24	The majority of controlled human exposure studies have investigated the effects of
25	exposure to O_3 in young healthy nonsmoking adults (18-35 years of age). These studies
26	typically use fixed concentrations of O3 under carefully regulated environmental
27	conditions and subject activity levels. The magnitude of respiratory effects (decrements
28	in spirometry measurements and increases in symptomatic response) in these individuals
29	is a function of O_3 concentration (C), minute ventilation (\dot{V}_E), and exposure duration
30	(time). Any physical activity will increase minute ventilation and therefore the dose of
31	inhaled O_3 . Dose of inhaled O_3 to the lower airways is also increased due to a shift from
32	nasal to oronasal breathing with a consequential decrease in O_3 scrubbing by the upper
33	airways. Thus, the intensity of physiological response following an acute exposure will
34	be strongly associated with minute ventilation.

1	The product of $C \times \dot{V}_E \times$ time is commonly used as a surrogate for O ₃ dose to the
2	respiratory tract in controlled human exposure studies. A large body of data regarding the
3	interdependent effects of C, \dot{V}_E , and time on pulmonary responses was assessed in the
4	1986 and 1996 O_3 AQCDs (U.S. EPA, 1996a, 1986). Acute responses were modeled as a
5	function of total inhaled dose (C × \dot{V}_E × time) which was found to be a better predictor of
6	response to O_3 than C, \dot{V}_E , or time of exposure, alone, or as a combination of any two of
7	these factors. However, intake dose ($C \times \dot{V}_E \times time$) did not adequately capture the
8	temporal dynamics of pulmonary responses in a comparison between a constant (square-
9	wave) and a variable (triangular) O_3 exposure (average 120 ppb O_3 ; moderate exercise,
9 10	
	$\dot{V}_{E} = 40$ L/min; 8 hour duration) conducted by <u>Hazucha et al. (1992</u>). Recent nonlinear
11	statistical models clearly describe the temporal dynamics of FEV_1 responses as a function
12	of C, \dot{V}_E , time, and age of the exposed subject (<u>McDonnell et al., 2010</u> , <u>2007</u>).
13	For healthy young adults exposed at rest for 2 hours, 500 ppb is the lowest O ₃
14	concentration reported to produce a statistically significant O ₃ -induced group mean FEV ₁
15	decrement of 6.4% (n = 10) (Folinsbee et al., 1978) to 6.7% (n = 13) (Horvath et al.,
16	<u>1979</u>). Airway resistance was not clearly affected during at-rest exposure to these
17	O_3 concentrations. For exposures of 1-2 hours to ≥ 120 ppb O_3 , statistically significant
18	symptomatic responses and effects on FEV ₁ are observed when \dot{V}_E is sufficiently
19	increased by exercise (McDonnell et al., 1999b). For instance, 5% of young healthy
20	adults exposed to 400 ppb O ₃ for 2 hours during rest experienced pain on deep
21	inspiration. Respiratory symptoms were not observed at lower exposure concentrations
22	(120-300 ppb) or with only 1 hour of exposure even at 400 ppb. However, when exposed
23	to 120 ppb O ₃ for 2 hours during light-to-moderate intermittent exercise (\dot{V}_E of 22 -
24	35 L/min), 9% of individuals experienced pain on deep inspiration, 5% experienced
25	cough, and 4% experienced shortness of breath. With very heavy continuous exercise
26	$(\dot{V}_E = 89 \text{ L/min})$, an O ₃ -induced group mean decrement of 9.7% in FEV ₁ has been reported
27	for healthy young adults exposed for 1 hour to 120 ppb O_3 (Gong et al., 1986). Symptoms
28	are present and decrements in forced expiratory volumes and flows occur at 160-240 ppb
29	O ₃ following 1 hour of continuous heavy exercise ($\dot{V}_E \approx 55$ to 90 L/min (<u>Gong et al.</u> ,
30	1986; Avol et al., 1984; Folinsbee et al., 1984; Adams and Schelegle, 1983) and
31	following 2 hours of intermittent heavy exercise ($\dot{V}_E \approx 65-68$ L/min) (Linn et al., 1986;
32	Kulle et al., 1985; McDonnell et al., 1983). With heavy intermittent exercise (15-min
33	intervals of rest and exercise [$\dot{V}_E = 68 \text{ L/min}$]), symptoms of breathing discomfort and a
34	group mean O_3 -induced decrement of 3.4% in FEV ₁ occurred in young healthy adults
35	exposed for 2 hours to 120 ppb O_3 (McDonnell et al., 1983). ¹ Table 6-1 provides
36	examples of typical exercise protocols utilized in controlled human exposures to O_3 . The

 1 In total, subjects were exposed to O_3 for 2.5 hours. Intermittent exercise periods, however, were only conducted for the first 2 hours of exposure and ${\sf FEV}_1$ was determined 5 minutes after the exercise was completed.

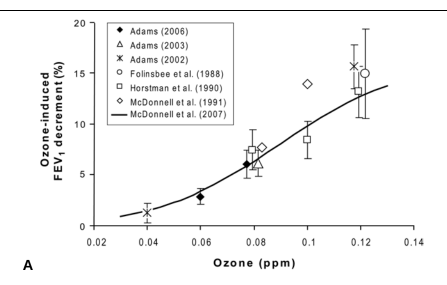
1	\dot{V}_{E} rates in this table are per body surface area (BSA) which is, on average, about 1.7 m ²
2	and 2.0 m ² for young healthy adult females and males, respectively, who participated in
3	controlled O ₃ exposure studies.

Table 6-1Activity levels used in controlled exposures of healthy young
adults to ozone.

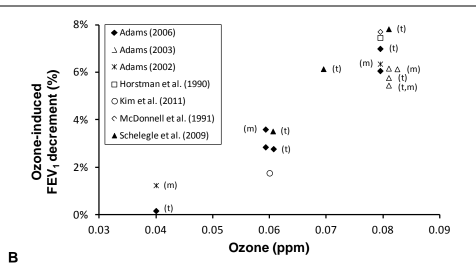
Activity ^{a,b}	Study Duration (hours)	(L/min per m ² BSA)	Heart Rate (bpm)	Treadmill Speed (mph)	Treadmill Grade (%)	Cycle (watts)
Rest	2	4	70	n.a.	n.a.	n.a.
Light quasi-continuous exercise	6.6-7.6	15	110	3.5-4.4	0	42
Moderate quasi- continuous exercise	6.6	17-23	115-130	3.3-3.5	4-5	72
Heavy intermittent exercise	1-2	27-33	160	3.5-5	10	100
Very heavy continuous exercise	1	45	160	n.a.	n.a.	260

^aBased on group mean exercise specific data provided in the individual studies. On average, subjects were 23 years of age. For exercise protocols, the minute ventilation and heart rate are for the exercise periods. Quasi-continuous exercise consists of 50 minutes of exercise periods followed by 10 minutes of rest. Intermittent exercise consists of alternating periods of 15 minutes of exercise and 15 minutes of rest.

^bReferences: <u>Horvath et al. (1979</u>) for rest; <u>Adams (2000</u>) and <u>Horstman et al. (1995</u>) for light quasi-continuous exercise, <u>2006a</u>); (2002, 2000), <u>Folinsbee et al. (1988</u>), <u>Horstman et al. (1990</u>), and <u>McDonnell et al. (1991</u>) for moderate quasi-continuous exercise; <u>Kehrl et al. (1987</u>), <u>Kreit et al. (1989</u>), and <u>McDonnell et al. (1983</u>) for heavy intermittent exercise, and <u>Gong et al. (1986</u>) for very heavy continuous exercise.



Source: Brown et al. (2008).



Top, panel A: All studies exposed subjects to a constant (square-wave) concentration in a chamber, except Adams (2002) where a facemask was used. All responses at and above 0.06 ppm were statistically significant. The McDonnell et al. (2007) curve illustrates the predicted FEV₁ decrement at 6.6 hours as a function of O_3 concentration for a 23 year-old (the average age of subjects that participated in the illustrated studies). Note that this curve was not "fitted" to the plotted data. Error bars (where available) are the standard error of responses.

Bottom, panel B: All studies used constant (square-wave) exposures in a chamber unless designated as triangular (t) and/or facemask (m) exposures. All responses at and above 0.07 ppm were statistically significant. At 0.06 ppm, <u>Adams (2006a)</u> and <u>Kim et al. (2011</u>) responses to square-wave chamber exposures were statistically significant. During each hour of the exposures, subjects were engaged in moderate quasi continuous exercise (40 L/min) for 50 minutes and rest for 10 minutes. Following the third hour, subjects had an additional 35-minute rest period for lunch. The data at 0.06, 0.08 and 0.12 ppm have been offset for illustrative purposes.

Studies appearing in the figure legends: <u>Adams (2006a</u>); (2003a, 2002), <u>Folinsbee et al. (1988</u>), <u>Horstman et al. (1990</u>), (<u>Kim et al.</u>, 2011), 2007); (<u>1991</u>), and <u>Schelegle et al. (2009</u>).

Figure 6-1 Cross-study comparison of mean ozone-induced FEV₁ decrements following 6.6 hours of exposure to ozone.

1	For prolonged (6.6 hours) exposures relative to shorter exposures, significant pulmonary
2	function responses and symptoms have been observed at lower O_3 concentrations and at a
3	moderate level of exercise ($\dot{V}_E = 40 \text{ L/min}$). The 6.6-hour experimental protocol was
4	intended to simulate the performance of heavy physical labor for a full workday
5	
	(Folinsbee et al., 1988). The results from studies using 6.6 hours of constant or square-
6	wave exposures to between 40 and 120 ppb O_3 are illustrated in Figure 6-1(A).
7	<u>Figure 6-1(B)</u> focuses on the range from 40 to 80 ppb and includes triangular exposure
8	protocols as well as facemask exposures. Exposure to 40 ppb O ₃ for 6.6 hours produces
9	small, statistically nonsignificant changes in FEV_1 that are relatively similar to responses
10	from FA exposure (<u>Adams, 2002</u>). Volunteers exposed to 60 ppb O_3 experience group
11	mean O ₃ -induced FEV ₁ decrements of about 3% (<u>Kim et al., 2011</u> ; <u>Brown et al., 2008</u> ;
12	<u>Adams, 2006a</u>) ¹ ; those exposed to 80 ppb have group mean decrements that range from 6
13	to 8% (<u>Adams, 2006a, 2003a;</u> <u>McDonnell et al., 1991;</u> <u>Horstman et al., 1990</u>); at 100 ppb,
14	group mean decrements range from 8 to 14% (McDonnell et al., 1991; Horstman et al.,
15	1990); and at 120 ppb, group mean decrements of 13 to 16% are observed (Adams, 2002;
16	Horstman et al., 1990; Folinsbee et al., 1988). As illustrated in Figure 6-1, there is a
17	smooth intake dose-response curve without evidence of a threshold for exposures
18	between 40 and 120 ppb O ₃ . This is consistent with <u>Hazucha and Lefohn (2007</u>), who
19	suggested that a randomly selected group of healthy individuals of sufficient size would
20	include hypo-, normo-, and hyper-responsive individuals such that the average response
21	would show no threshold for any spirometric endpoint. Taken together, these data
22	indicate that mean FEV ₁ is clearly decreased by 6.6-hour exposures to 60 ppb O_3 and
23	higher concentrations in subjects performing moderate exercise.
24	The time course of responses during prolonged (6.6 hours) square-wave O3 exposures
25	with moderate exercise ($\dot{V}_E = 40 \text{ L/min}$) depends on O ₃ concentration. At 120 ppb O ₃ ,
26	<u>Folinsbee et al. (1988</u>) observed that somewhat small FEV_1 decrements and symptoms of
27	breathing discomfort become apparent in healthy subjects following the second hour of
28	exposure with a more rapid change in responses between the 3rd and 5th hour of
29	exposure and a diminishing response or plateau in responses over the last hour of
30	exposure. Relative to FA, the change in FEV_1 at 120 ppb O ₃ became statistically
31	significant after 4.6 hours. Following the same exposure protocol, <u>Horstman et al. (1990</u>)
32	observed a linear increase in FEV_1 responses with time following 2 hours of exposure to
33	120 ppb O_3 that was statistically different from FA responses after 3 h. At 100 ppb O_3 ,
34	FEV_1 responses diverged from FA after 3 hours and were statistically different at 4.6
35	hours (Horstman et al., 1990). At 80 ppb O_3 , FEV ₁ responses diverged from FA after 4.6
55	$(\underline{\mathbf{rorstinum}} \mathbf{v}, \underline{\mathbf{u}}, \underline{\mathbf{ryy}}). \mathbf{r}, \mathbf{v} \mathbf{v} \mathbf{v}_{3}, \mathbf{r} \mathbf{L} \mathbf{v}_{1} \mathbf{responses} \mathbf{u} \mathbf{v} \mathbf{c} \mathbf{r} \mathbf{g} \mathbf{u} \mathbf{n} \mathbf{u} \mathbf{n} \mathbf{r} \mathbf{A} \mathbf{u} \mathbf{t} \mathbf{c} 1 4.0$

¹ <u>Adams (2006b</u>) did not find effects on FEV₁ at 60 ppb to be statistically significant. In an analysis of the <u>Adams (2006b</u>) data, even after removal of potential outliers, <u>Brown et al. (2008</u>) found the average effect on FEV₁ at 60 ppb to be small, but highly statistically significant (p < 0.002) using several common statistical tests.

1	hours and were statistically different from FA at 5.6 hours (Horstman et al., 1990).
2	Subsequently, Adams (2006a) observed FEV_1 decrements and total respiratory symptoms
3	at 80 ppb O_3 to diverge from FA responses after 3 h, but did not become statistically
4	different until 6.6 hours. At 60 ppb O ₃ , FEV ₁ responses generally tracked responses in FA
5	for the first 4.6 hours of exposure and diverged after 5.6 hours (Adams, 2006a). FEV ₁
6	responses, but not symptomatic responses, become statistically different between 60 ppb
7	O_3 and FA at 6.6 hours (Kim et al., 2011; Brown et al., 2008). At 40 ppb, FEV ₁ and
8	symptomatic responses track FA for 5.6 hours of exposure and may begin to diverge after
9	6.6 hours (Adams, 2002). In prolonged (6.6 hours) square-wave O_3 exposures between 40
10	and 120 ppb with moderate exercise ($\dot{V}_E = 40$ L/min), the time required for group mean
11	responses to differ between O_3 and FA exposures increases with decreasing O_3
12	concentration.

- 13 As opposed to constant (i.e., square-wave) concentration patterns used in the studies 14 described above, many studies conducted at the levels of 40-80 ppb have used variable 15 O_3 concentration patterns. It has been suggested that a triangular (variable concentration) 16 exposure profile can potentially lead to higher FEV_1 responses than square-wave profiles 17 despite having the same average O₃ concentration over the exposure period. Hazucha et 18 al. (1992) were the first to investigate the effects of variable versus constant 19 concentration exposures on responsiveness to O₃. In their study, volunteers were 20 randomly exposed to a triangular concentration profile (averaging 120 ppb over the 21 8-hour exposure) that increased linearly from 0-240 ppb for the first 4 hours of the 8-hour 22 exposure, then decreased linearly from 240 to 0 ppb over the next 4 hours of the 8-hour 23 exposure, and to an square-wave exposure of 120 ppb O₃ for 8 hours. While the total 24 inhaled O₃ doses at 4 hours and 8 hours for the square-wave and the triangular 25 concentration profile were almost identical, the FEV_1 responses were dissimilar. For the square-wave exposure, FEV₁ declined \sim 5% by the fifth hour and then remained at that 26 27 level. With the triangular O_3 profile, there was minimal FEV₁ response over the first 28 3 hours followed by a rapid decrease in FEV₁ to a decrement of 10.3% over the next 3 29 hours. During the seventh and eighth hours, mean FEV_1 decrement improved to 6.3% as 30 the O_3 concentration decreased from 120 to 0 ppb (mean = 60 ppb). These findings 31 illustrate that the severity of symptoms and the magnitude of spirometric responses are 32 time-dependent functions of O_3 delivery rate with periods of both effect development and 33 recovery during the course of an exposure.
- 34Subsequently, others have also demonstrated that variable concentration exposures can35elicit greater FEV1 and symptomatic responses than do square-wave exposures (Adams,362006a, b, 2003a). Adams (2006b) reproduced the findings of Hazucha et al. (1992) at37120 ppb. However, Adams (2006a); (2003a) found that responses from an 80 ppb O338(average) triangular exposure did not differ significantly from those observed in the

1 80 ppb O_3 square-wave exposure at 6.6 hours. Nevertheless, FEV₁ and symptoms were 2 significantly different from pre-exposure at 4.6 hours (when the O_3 concentration was 3 150 ppb) in the triangular exposure, but not until 6.6 hours in the square-wave exposure. 4 At the lower O_3 concentration of 60 ppb, no temporal pattern differences in FEV₁ 5 responses between square-wave and triangular exposure profiles could be discerned 6 (Adams, 2006a). However, both total symptom scores and pain on deep inspiration 7 tended to be greater following the 60 ppb triangular than the 60 ppb square-wave 8 exposure. At 80 ppb, respiratory symptoms tended to increase more rapidly during the 9 triangular than square-wave exposure protocol, but then decreased during the last hour of 10 exposure to be less than that observed with the square-wave exposure at 6.6 hours. Both 11 total symptom scores and pain on deep inspiration were significantly increased following 12 exposures to 80 ppb relative to all other exposure protocols, i.e., FA, 40, and 60 ppb 13 exposures. Following the 6.6-hour exposures, respiratory symptoms at 80 ppb were 14 roughly 2-3 times greater than those observed at 60 ppb. At 40 ppb, triangular and 15 square-wave patterns produced spirometric and subjective symptom responses similar to 16 FA exposure (Adams, 2006a, 2002).

17 For O_3 exposures of 60 ppb and greater, studies (Adams, 2006a, b, 2003a; Hazucha et al., 18 1992) demonstrate that during triangular exposure protocols, volunteers exposed during 19 moderate exercise ($\dot{V}_E = 40 \text{ L/min}$) may develop greater spirometric and/or symptomatic 20 responses during and following peak O_3 concentrations as compared to responses over 21 the same time interval of square-wave exposures. This observation is not unexpected 22 since the inhaled dose rate during peaks of the triangular protocols approached twice that 23 of the square-wave protocols, e.g., 150 ppb versus 80 ppb peak concentration. At time 24 intervals toward the end of an exposure, O₃ delivery rates for the triangular protocols 25 were less than those of square-wave. At these later time intervals, there is some recovery 26 of responses during triangular exposure protocols, whereas there is a continued 27 development of or a plateau of responses in the square-wave exposure protocols. Thus, 28 responses during triangular protocols relative to square-wave protocols may be expected 29 to diverge and be greater following peak exposures and then converge toward the end of 30 an exposure. Subsequent discussion will focus on exposures between 40 and 80 ppb 31 where FEV₁ pre-to-post responses are similar (although not identical) between triangular 32 and square-wave protocols having equivalent average exposure concentrations.

33Schelegle et al. (2009) recently investigated the effects of 6.6-hour variable O_3 exposure34protocols at mean concentrations of 60, 70, 80, and 87 ppb on respiratory symptoms and35pulmonary function in young healthy adults (16 F, 15 M; 21.4 ± 0.6 years) exposed36during moderate quasi continuous exercise ($\dot{V}_E = 40 \text{ L/min}$). The mean FEV1 (± standard37error) decrements at 6.6 hours (end of exposure relative to pre-exposure) were -38 $0.80 \pm 0.90\%$, $2.72 \pm 1.48\%$, $5.34 \pm 1.42\%$, $7.02 \pm 1.60\%$, and $11.42 \pm 2.20\%$ for

1	exposure to FA, 60, 70, 80, and 87 ppb O_3 , respectively. Statistically significant
2	decrements in FEV ₁ and increases in total subjective symptom scores ($p < 0.05$) were
3	found following exposure to mean concentrations of 70, 80, and 87 ppb O ₃ relative to FA.
4	Statistically significant effects were not found at 60 ppb. One of the expressed purposes
5	of the <u>Schelegle et al. (2009</u>) study was to determine the minimal mean O_3 concentration
6	that produces a statistically significant decrement in FEV_1 and respiratory symptoms in
7	healthy individuals completing 6.6-hour exposure protocols. At 70 ppb, Schelegle et al.
8	(2009) observed a statistically significant O_3 -induced FEV ₁ decrement of 6.1% at
9	6.6 hours and a significant increase in total subjective symptoms at 5.6 and 6.6 hours. A
10	re analysis found the FEV_1 responses at 70 ppb to be significantly different from FA
11	responses beginning at 4.6 hours of exposure (Lefohn et al., 2010a). At 60 ppb, an
12	O_3 -induced 3.5% FEV ₁ decrement was not found to be statistically significant. However,
13	this effect is similar in magnitude to the 2.9% FEV_1 decrement at 60 ppb observed by
14	Adams (2006a), which was found to be statistically significant by Brown et al. (2008).
15	More recently, Kim et al. (2011) investigated the effects of a 6.6-hour exposure to 60 ppb

16 O_3 during moderate quasi continuous exercise ($\dot{V}_E = 40 \text{ L/min}$) on pulmonary function 17 and respiratory symptoms in young healthy adults (32 F, 27 M; 25.0 ± 0.5 year) who 18 were roughly half GSTM1-null and half GSTM1-positive. Sputum neutrophil levels were also measured in a subset of the subjects (13 F, 11 M). The mean FEV₁ (± standard error) 19 20 decrements at 6.6 hours (end of exposure relative to pre-exposure) were significantly 21 different (p = 0.008) between the FA (0.002 \pm 0.46%) and O₃ (1.76 \pm 0.50%) exposures. 22 The inflammatory response following O_3 exposure was also significantly (p < 0.001) 23 increased relative to the FA exposure. Respiratory symptoms were not affected by O_3 24 exposure. There was also no significant effect of GSTM1 genotype on FEV₁ or 25 inflammatory responses to O₃.

26 Consideration of the minimal O_3 concentration producing statistically significant effects 27 on FEV_1 and respiratory symptoms (e.g., cough and pain on deep inspiration) following 28 6.6-hour exposures warrants additional discussion. As discussed above, numerous studies 29 have demonstrated statistically significant O₃-induced group mean FEV₁ decrements of 30 6-8% and an increase in respiratory symptoms at 80 ppb. Schelegle et al. (2009) have 31 now reported a statistically significant O_3 -induced group mean FEV₁ decrement of 6%, as 32 well as increased respiratory symptoms, at 70 ppb. At 60 ppb, there is information 33 available from 4 separate studies (Kim et al., 2011; Schelegle et al., 2009; Adams, 2006a,

1	
1	<u>2002</u>). ¹ The group mean O_3 -induced FEV ₁ decrements observed in these studies were
2	3.6% (facemask, square-wave) by <u>Adams (2006a</u>); (2002) ² , 2.8% (triangular exposure)
3	and 2.9% (square-wave exposure) by <u>Adams (2006a</u>), 3.5% (triangular exposure) by
4	Schelegle et al. (2009), and 1.8% (square-wave exposure) by Kim et al. (2011). Based on
5	data from these studies, at 60 ppb, the weighted-average group mean O_3 -induced FEV_1
6	decrement (i.e., adjusted for FA responses) is 2.7% (n = 150). Although not consistently
7	statistically significant, these group mean changes in FEV_1 at 60 ppb are consistent
8	among studies, i.e., none observed an average improvement in lung function following a
9	6.6-hour exposure to 60 ppb O_3 . Indeed, as was illustrated in Figure 6-1, the group mean
10	FEV ₁ responses at 60 ppb fall on a smooth intake dose-response curve for exposures
11	between 40 and 120 ppb O ₃ . Furthermore, in a re-analysis of the 60 ppb square-wave data
12	from Adams (2006a), Brown et al. (2008) found the mean effects on FEV_1 to be highly
13	statistically significant (p <0.002) using several common statistical tests even after
14	removal of 3 potential outliers. A statistically significant increase in total respiratory
15	symptoms at 60 ppb has only been reported by Adams (2006a) for a triangular exposure
16	protocol at 5.6 hours and 6.6 hours relative to baseline (not FA). Although not
17	statistically significant, there was a tendency for an increase in total symptoms and pain
18	on deep inspiration following the 60 ppb exposures (triangular and square-wave) relative
19	to those following both FA and 40 ppb exposures. The time-course and magnitude of
20	FEV ₁ responses at 40 ppb resemble those occurring during FA exposures (Adams, 2006a,
21	2002). In both of these studies, there was a tendency (not statistically significant) for a
22	small increase in total symptoms and pain on deep inspiration following the 40 ppb
23	exposures relative to those following FA. Taken together, the available evidence shows
24	that detectable effects of O_3 on group mean FEV ₁ persist down to 60 ppb, but not 40 ppb
25	in young healthy adults exposed for 6.6 hours during moderate exercise. Although group
26	mean FEV ₁ responses at 60 ppb are relatively small (2-3% mean FEV ₁ decrement), it
27	should be emphasized that there is considerable intersubject variability, with some
28	responsive individuals consistently experiencing larger than average FEV_1 responses.
29	In addition to overt effects of O ₃ exposure on the large airways indicated by spirometric
30	responses, O ₃ exposure also affects the function of the small airways and parenchymal
31	lung. Foster et al. (1997); (1993) examined the effect of O_3 on ventilation distribution. In
32	healthy adult males (n = 6; 26.7 \pm 7 years old) exposed to O ₃ (330 ppb with light
33	intermittent exercise for 2 h), there was a significant reduction in ventilation to the lower

¹ <u>Adams (2006a)</u>; (2002) both provide data for an additional group of 30 healthy subjects that were exposed via facemask to 60 ppb (square-wave) O₃ for 6.6 hours with moderate exercise ($\dot{V}_E = 23 \text{ L/min per m}^2 \text{ BSA}$). These subjects are described on page 133 of <u>Adams (2006a)</u> and pages 747 and 761 of <u>Adams (2002</u>). The FEV₁ decrement may be somewhat increased due to a target \dot{V}_E of 23 L/min per m² BSA relative to other studies with which it is listed having the target \dot{V}_E of 20 L/min per m² BSA. Based on <u>Adams (2003a, b, 2002</u>) the facemask exposure is not expect to affect the FEV₁ responses relative to a chamber exposure.

² This group average FEV₁ response is for a set of subjects exposed via facemask to 60 ppb O₃, see page 133 of <u>Adams (2006a)</u>.

1	lung (31% of lung volume) and significant increases in ventilation to the upper- and
2	middle-lung regions (Foster et al., 1993). In a subsequent study of healthy males ($n = 15$;
3	25.4 ± 2 years old) exposed to O ₃ (350 ppb with moderate intermittent exercise for 2.2 h),
4	O_3 exposure caused a delayed gas washout in addition to a 14% FEV ₁ decrement (Foster
5	et al., 1997). The pronounced slow phase of gas washout following O_3 exposure
6	represented a 24% decrease in the washout rate. A day following O ₃ exposure, 50% of
7	the subjects still had (or developed) a delayed washout relative to the pre- O_3 maneuver.
8	These studies suggest a prolonged O_3 effect on the small airways and ventilation
9	distribution in healthy young individuals.
10	There is a rapid recovery of O_3 -induced spirometric responses and symptoms; 40 to 65%
10	recovery appears to occur within about 2 hours following exposure (Folinsbee and
12	<u>Hazucha, 1989</u>). For example, following a 2-hour exposure to 400 ppb O_3 with
13	intermittent exercise, Nightingale et al. (2000) observed a 13.5% mean decrement in
14	FEV ₁ . By 3 hours postexposure, however, only a 2.7% FEV ₁ decrement persisted. Partial
15	recovery also occurs following cessation of exercise despite continued exposure to O ₃
16	(Folinsbee et al., 1977) and at low O_3 concentrations during exposure (Hazucha et al.,
17	<u>1992</u>). A slower recovery phase, especially after exposure to higher O_3 concentrations,
18	may take at least 24 hours to complete (Folinsbee and Hazucha, 2000; Folinsbee et al.,
19	<u>1993</u>). Repeated daily exposure studies at higher concentrations typically show that FEV_1
20	response to O_3 is enhanced on the second day of exposure. This enhanced response
21	suggests a residual effect of the previous exposure, about 22 hours earlier, even though
22	the pre-exposure spirometry may be the same as on the previous day. The absence of the
23	enhanced response with repeated exposure at lower O ₃ concentrations may be the result
24	of a more complete recovery or less damage to pulmonary tissues (Folinsbee et al., 1994).

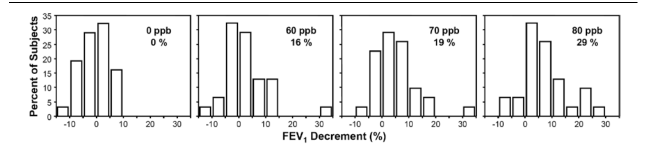
Predicted Responses in Healthy Subjects

25 Studies analyzing large data sets (hundreds of subjects) provide better predictive ability 26 of acute changes in FEV₁ at low levels of O_3 and \dot{V}_E than is possible via comparisons 27 between smaller studies. A few such studies described in the 2006 O₃ AQCD (U.S. EPA, 28 2006b) analyzed FEV₁ responses in healthy young adults (18-35 years of age) recruited 29 from the area around Chapel Hill, NC and exposed for 2 hours to O_3 concentrations of up 30 to 400 ppb at rest or with intermittent exercise (McDonnell et al., 1997; Seal et al., 1996; 31 Seal et al., 1993). McDonnell et al. (1999b) examined changes in respiratory symptoms 32 with O₃ exposure in a subset of the Chapel Hill data. In general, these studies showed that 33 FEV1 and respiratory symptom responses increase with increasing O3 concentration and 34 $\dot{V}_{\rm E}$ and decrease with increasing subject age. More recent studies expand upon these 35 analyses of FEV₁ responses to also include longer duration (up to 8 h) studies and periods 36 of recovery following exposure.

- 1 McDonnell et al. (2007) provided a nonlinear empirical model for predicting group 2 average FEV₁ responses as a function of O_3 concentration, exposure time, \dot{V}_E , and age of 3 the exposed individual. The model predicts temporal dynamics of FEV_1 change in 4 response to any set of O_3 exposure conditions that might reasonably be experienced in the 5 ambient environment. The model substantially differs from earlier statistical models in 6 that it effectively considers the concurrent processes of damage and repair, i.e., the model 7 allows effects on FEV_1 to accumulate during exposure at the same time they are reduced 8 due to the reversible nature of the effects. The model was based on response data of 9 healthy, nonsmoking, white males (n = 541), 18-35 years old, from 15 studies conducted 10 at the U.S. EPA Human Studies Facility in Chapel Hill, NC.
- 11 McDonnell et al. (2010) tested the predictive ability of the model (McDonnell et al., 12 2007) against independent data (i.e., data that were not used to fit the model) of Adams 13 (2006a); (2006b, 2003a, 2002, 2000), Hazucha et al. (1992), and Schelegle et al. (2009). 14 The model generally captured the dynamics of group average FEV₁ responses within 15 about a one percentage point of the experimental data. Consistent with Bennett et al. 16 (2007), an increased body mass index (BMI) was found to be associated with enhanced 17 FEV₁ responses to O₃ by McDonnell et al. (2010). The BMI effect is of the same order of 18 magnitude but in the opposite direction of the age effect where by FEV₁ responses 19 diminish with increasing age. Although the effects of age and BMI are relatively strong, 20 these characteristics account for only a small amount of the observed variability in 21 individual responses.
- 22 Alternatively, Lefohn et al. (2010a) proposed that FEV_1 responses to O_3 exposure might 23 be described by a cumulative integrated exposure index with a sigmoidal weighting 24 function similar to the W126 used for predicting vegetation effects (see Section 9.5). The 25 integrated exposure index is the sum of the hourly average O₃ concentrations times their 26 respective weighing factors. Based on a limited number of studies, the authors assumed 27 weighting factors ranged from near zero at 50 ppb up to approximately 1.0 for 28 concentrations at \geq 125 ppb. The concentrations of 60, 70 and 80 ppb correspond to the 29 weights of 0.14, 0.28, and 0.50, respectively. These weighting factors apply only to the case of exposure during moderate exercise ($\dot{V}_E = 20$ L/min per m² BSA). Lefohn et al. 30 31 (2010a) calculated the cumulative exposure index for the protocols used by Adams 32 (2006a); (2003a) and Schelegle et al. (2009). They found statistically significant O₃ 33 effects after 4 hours on FEV_1 at 105 ppb-hour based on Schelegle et al. (2009) and at 34 235 ppb-hour based on Adams (2006a); (2003a). Based on this analysis, the authors 35 recommended a 5-hour accumulation period to protect against O₃ effects on lung 36 function.

Intersubject Variability in Response of Healthy Subjects

1	Consideration of group mean changes is important in discerning if observed effects are
2	due to O ₃ exposure rather than chance alone. Inter-individual variability in responses is,
3	however, considerable and pertinent to assessing the fraction of the population that might
4	actually be affected during an O_3 exposure. <u>Hackney et al. (1975</u>) first recognized a wide
5	range in the sensitivity of subjects to O_3 . The range in the subjects' ages (29 to 49 years)
6	and smoking status (0 to 50 pack years) in the Hackney et al. (1975) study are now
7	understood to affect the spirometric and symptomatic responses to O ₃ . Subsequently,
8	DeLucia and Adams (1977) examined responses to O_3 in six healthy non-smokers and
9	found that two exhibited notably greater sensitivity to O ₃ . Since that time, numerous
10	studies have documented considerable variability in responsiveness to O_3 even in subjects
11	recruited to assure homogeneity in factors recognized or presumed to affect responses.
12	An individual's FEV ₁ response to a 2 hour O_3 exposure is generally reproducible over
13	several months and presumably reflects the intrinsic responsiveness of the individual to
14	O ₃ (<u>Hazucha et al., 2003; McDonnell et al., 1985b</u>). The frequency distribution of
15	individual FEV_1 responses following these relatively short exposures becomes skewed as
16	the group mean response increases, with some individuals experiencing large reductions
17	in FEV ₁ (Weinmann et al., 1995a; Kulle et al., 1985). For 2-hour exposures with
18	intermittent exercise causing a predicted average FEV ₁ decrement of 10%, individual
19	decrements ranged from approximately 0 to 40% in white males aged 18-36 years
20	(McDonnell et al., 1997). For an average FEV_1 decrement of 13%, Ultman et al. (2004)
21	reported FEV ₁ responses ranging from a 4% improvement to a 56% decrement in young
22	healthy adults (32 M, 28 F) exposed for 1 hour to 250 ppb O ₃ . One-third of the subjects
23	had FEV ₁ decrements of >15%, and 7% of the subjects had decrements of >40%. The
24	differences in FEV_1 responses did not appear to be explained by intersubject differences
25	in the fraction of inhaled O_3 retained in the lung (<u>Ultman et al., 2004</u>).



Note: During each hour of the exposures, subjects were engaged in moderate quasi continuous exercise (40 L/min) for 50 minutes and rest for 10 minutes. Following the third hour, subjects had an additional 35 minute rest period for lunch. Subjects were exposed to a triangular O_3 concentration profile having the average O_3 concentration provided in each panel. As average ozone concentration increased, the distribution of responses became asymmetric with a few individuals exhibiting large FEV₁ decrements. The percentage indicated in each panel is the portion of subjects having a FEV₁ decrement in excess of 10%. Source: Adapted with permission of American Thoracic Society (Schelegle et al., 2009).

Figure 6-2 Frequency distributions of FEV₁ decrements observed by Schelegle et al. (2009) in young healthy adults (16 F, 15 M) following 6.6-hour exposures to ozone or filtered air.

1	Consistent with the 1- to 2-hour studies, the distribution of individual responses
2	following 6.6-hour exposures becomes skewed with increasing exposure concentration
3	and magnitude of the group mean FEV_1 response (McDonnell, 1996). Figure 6-2
4	illustrates frequency distributions of individual FEV ₁ responses observed in 31 young
5	healthy adults following 6.6-hour exposures between 0 and 80 ppb. Schelegle et al.
6	(2009) found >10% FEV ₁ decrements in 16, 19, 29, and 42% of individuals exposed for
7	6.6 hours to 60, 70, 80, and 87 ppb, respectively. Just as there are differences in mean
8	decrements between studies having similar exposure scenarios (Figure 6-1 at 80 and
9	120 ppb), there are differences in the proportion of individuals affected with $>10\%$ FEV ₁
10	decrements. At 80 ppb, the proportion affected with $>10\%$ FEV ₁ decrements was 17%
11	(n = 30) by <u>Adams (2006a</u>) ¹ , 26% $(n = 60)$ by <u>McDonnell (1996</u>), and 29% $(n = 31)$ by
12	<u>Schelegle et al. (2009</u>). At 60 ppb, the proportion with $>10\%$ FEV ₁ decrements was 20%
13	(n = 30) by <u>Adams (2002</u>) ² , 3% $(n = 30)$ by <u>Adams (2006a</u>) ¹ , 16% $(n = 31)$ by <u>Schelegle</u>
14	et al. (2009), and 5% (n = 59) by Kim et al. (2011). Based on these studies, the weighted
15	average proportion of individuals with >10% FEV1 decrements is 10% following
16	exposure to 60 ppb. Due to limited data within the published papers, these proportions
17	were not corrected for responses to FA exposure where lung function typically improves
18	in healthy adults. For example, uncorrected versus O3-induced (i.e., adjusted for response

¹ Not assessed by <u>Adams (2006a)</u>, the proportion was provided in Figure 8-1B of the 2006 O₃ AQCD (U.S. EPA, 2006b).

² This information is from page 761 of <u>Adams (2002)</u>. <u>Adams (2006a, 2002)</u> both provide data for a group of 30 healthy subjects that were exposed via facemask to 60 ppb (square-wave) O₃ for 6.6 hours with moderate exercise ($\dot{V}_E = 23 \text{ L/min per m}^2 \text{ BSA}$). These subjects are described on page 133 of <u>Adams (2006a</u>) and pages 747 and 761 of <u>Adams (2002</u>). The FEV₁ decrement may be somewhat increased due to a target \dot{V}_E of 23 L/min per m² BSA relative to other studies with which it is listed having the target \dot{V}_E of 20 L/min per m² BSA. Based on <u>Adams (2003a, b, 2002</u>), similar FEV₁ responses are expected between facemask and chamber exposures.

during FA exposure) proportions of individuals having >10% FEV₁ decrements in the <u>Adams (2006a</u>)¹ study were, respectively, 3% versus 7% at 60 ppb and 17% versus 23% at 80 ppb. Thus, uncorrected proportions underestimate the actual fraction of healthy individuals affected.

5 Given considerable inter-individual variability in responses, the interpretation of 6 biologically small group mean decrements requires careful consideration. Following 7 prolonged 6.6-hour exposures to an average level of 60 ppb O_3 , data available from four 8 studies yield a weighted-average group mean O_3 -induced FEV₁ decrement (i.e., adjusted 9 for FA responses) of 2.7% (n = 150) (Kim et al., 2011; Schelegle et al., 2009; Adams, 10 2006a, 1998). The data from these studies also yield a weighted-average proportion 11 (uncorrected for FA responses) of subjects with >10% FEV₁ decrements of 10% 12 (n = 150) (<u>Kim et al., 2011; Schelegle et al., 2009; Adams, 2006a, 1998</u>). In an individual 13 with relatively "normal" lung function, with recognition of the technical and biological 14 variability in measurements, confidence can be given that within-day changes in FEV_1 of 15 \geq 5% are clinically meaningful (Pellegrino et al., 2005; ATS, 1991). Here focus is given 16 to individuals with >10% decrements in FEV₁ since some individuals in the Schelegle et 17 al. (2009) study experienced 5-10% FEV₁ decrements following exposure to FA. A 10% 18 FEV₁ decrement is also generally accepted as an abnormal response and a reasonable 19 criterion for assessing exercise-induced bronchoconstriction (Dryden et al., 2010; ATS, 20 2000a). The data are not available in the published papers to determine the O₃-induced 21 proportion for either the Adams (1998) or Schelegle et al. (2009) studies. As already 22 stated, however, this uncorrected proportion likely underestimates the actual proportion 23 of healthy individuals experiencing O_3 -induced FEV₁ decrements in excess of 10%. 24 Therefore, by considering uncorrected responses and those individuals having >10%25 decrements, 10% is an underestimate of the proportion of healthy individuals that are 26 likely to experience clinically meaningful changes in lung function following exposure 27 for 6.6 hours to 60 ppb O_3 during moderate exercise. Of the studies conducted at 60 ppb, 28 only Kim et al. (2011) reported FEV₁ decrements at 60 ppb to be statistically significant. 29 However, Brown et al. (2008) found those from Adams (2006a) to be highly statistically 30 significant. Though group mean decrements are biologically small and generally do not 31 attain statistical significance, a considerable fraction of exposed individuals experience 32 clinically meaningful decrements in lung function.

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¹ Not assessed by <u>Adams (2006a</u>), uncorrected and O₃-induced proportions are from Figures 8-1B and 8-2, respectively, of the 2006 O₃ AQCD (2006b).

Factors Modifying Responsiveness to Ozone

	Physical activity increases \dot{V}_E and therefore the dose of inhaled O_3 . Consequently, the
,	intensity of physiological response during and following an acute O ₃ exposure will be
	strongly associated with minute ventilation. Apart from inhaled O3 dose and related
	environmental factors (e.g., repeated daily exposures), individual-level factors, such as
	health status, age, gender, ethnicity, race, smoking habit, diet, and socioeconomic status
i la	(SES) have been considered as potential modulators of a physiologic response to such
	exposures.

Responses in Individuals with Pre-existing Disease

- 8 Individuals with respiratory disease are of primary concern in evaluating the health
 9 effects of O₃ because a given change in function is likely to have more impact on a
 10 person with preexisting function impairment and reduced reserve.
- 11 Possibly due to the age of subjects studied, patients with COPD performing light to 12 moderate exercise do not generally experience statistically significant pulmonary 13 function decrements following 1- and 2-hour exposures to ≤ 300 ppb O₃ (Kehrl et al., 14 1985; Linn et al., 1983; Linn et al., 1982a; Solic et al., 1982). Following a 4-hour 15 exposure to 240 ppb O₃ during exercise, Gong et al. (1997b) found an O₃-induced FEV₁ 16 decrement of 8% in COPD patients which was not statistically different from the 17 decrement of 3% in healthy subjects. Demonstrating the need for control exposures and 18 presumably due to exercise, four of the patients in the Gong et al. (1997b) study had 19 FEV₁ decrements of >14% following both the FA and O₃ exposures. Although the 20 clinical significance is uncertain, small transient decreases in arterial blood oxygen 21 saturation have also been observed in some of these studies.
- 22 Based on studies reviewed in the 1996 and 2006 O₃ AQCDs, asthmatic subjects appear to 23 be at least as sensitive to acute effects of O_3 as healthy nonasthmatic subjects. Horstman 24 et al. (1995) found the O_3 -induced FEV₁ decrement in 17 mild-to-moderate asthmatics to 25 be significantly larger than that in 13 healthy subjects (19% versus 10%, respectively) 26 exposed to 160 ppb O₃ during light exercise (\dot{V}_E of 15 L/min per m² BSA) for 7.6-hour 27 exposure. In asthmatics, a significant positive correlation between O₃-induced 28 spirometric responses and baseline lung function was observed, i.e., responses increased 29 with severity of disease. In the shorter duration study by Kreit et al. (1989), 9 asthmatics 30 also showed a considerable larger average O₃-induced FEV₁ decrement than 9 healthy 31 controls (25% vs. 16%, respectively) following exposure to 400 ppb O₃ for 2 hours with 32 moderate-heavy exercise ($\dot{V}_E = 54 \text{ L/min}$). <u>Alexis et al. (2000)</u> [400 ppb; 2 h; exercise, 33 $\dot{V}_E = 30 \text{ L/min}$] and <u>Jorres et al. (1996</u>) [250 ppb; 3 h; exercise, $\dot{V}_E = 30 \text{ L/min}$] reported a 34 tendency for slightly greater FEV₁ decrements in asthmatics than healthy subjects.

1	Several studies reported similar responses between asthmatics and healthy individuals
2	(Scannell et al., 1996; Hiltermann et al., 1995; Basha et al., 1994). The lack of differences
3	in the Hiltermann et al. (1995) [400 ppb; 2 h; exercise, $\dot{V}_E = 20$ L/min] and Basha et al.
4	(1994) [200 ppb; 6 h; exercise, $\dot{V}_E = 25$ L/min] studies was not surprising, however, given
5	extremely small sample sizes (5-6 subjects per group) and corresponding lack of
6	statistical power. Power was not likely problematic for Scannell et al. (1996) [200 ppb;
7	4 h; exercise, $\dot{V}_E \approx 44$ L/min] with 18 mild asthmatics and 81 age-matched healthy
8	controls from companion studies (Balmes et al., 1996; Aris et al., 1995). Of note,
9	Mudway et al. (2001) reported a tendency for asthmatics to have smaller O ₃ -induced
10	FEV1 decrements than healthy subjects (3% versus 8%, respectively) when exposed to
11	200 ppb O_3 for 2 hours during exercise. However, the asthmatics in (Mudway et al.,
12	2001) also tended to be older than the healthy subjects, which could partially explain
13	their smaller response since FEV_1 responses to O_3 diminish with age.
14	In a study published since the 2006 O ₃ AQCD, <u>Stenfors et al. (2010</u>) exposed persistent
15	asthmatics (n = 13; aged 33 years) receiving chronic inhaled corticosteroid therapy to
16	200 ppb O_3 for 2 hours with moderate exercise. An average O_3 -induced FEV ₁ decrement
17	of 8.4% was observed, whereas, only a 3.0% FEV ₁ decrement is predicted for similarly

10200 ppb 03 for 2 hours with inductive chereice in a verify 03 induced i EV (acceleration)17of 8.4% was observed, whereas, only a 3.0% FEV1 decrement is predicted for similarly18exposed age-matched healthy controls (McDonnell et al., 2007). Vagaggini et al. (2010)19exposed mild-to-moderate asthmatics (n = 23; 33 ± 11 years) to 300 ppb O3 for 2 hours20with moderate exercise. Although the group mean O3-induced FEV1 decrement was only214%, eight subjects were categorized as "responders" with >10% FEV1 decrements.22Baseline lung function did not differ between the responders and nonresponders23suggesting that, in contrast to Horstman et al. (1995), O3-induced FEV1 responses were24not associated with disease severity.

Lifestage

25 Children, adolescents, and young adults (<18 years of age) appear, on average, to have 26 nearly equivalent spirometric responses to O₃, but have greater responses than middle-27 aged and older adults when similarly exposed to O₃ (U.S. EPA, 1996a). Symptomatic responses to O₃ exposure, however, appear to increase with age until early adulthood and 28 29 then gradually decrease with increasing age (U.S. EPA, 1996a). For example, healthy 30 children (aged 8-11 y) exposed to 120 ppb O₃ (2.5 h; heavy intermittent exercise) 31 experienced similar spirometric responses, but lesser symptoms than similarly exposed 32 young healthy adults (McDonnell et al., 1985a). For subjects aged 18-36 years, 33 McDonnell et al. (1999b) reported that symptom responses from O_3 exposure also 34 decrease with increasing age. Diminished symptomatic responses in children and the 35 elderly might put these groups at increased risk for continued O₃ exposure, i.e., a lack of 36 symptoms may result in their not avoiding or ceasing exposure. Once lung growth and

- 1development reaches the peak (18-20 years of age in females and early twenties in2males), pulmonary function, which is at its maximum as well, begins to decline3progressively with age as does O3 sensitivity.
- 4 In healthy individuals, the fastest rate of decline in O_3 responsiveness appears between 5 the ages of 18 and 35 years (Passannante et al., 1998; Seal et al., 1996), more so for 6 females then males (Hazucha et al., 2003). During the middle age period (35-55 years), 7 O₃ sensitivity continues to decline, but at a much lower rate. Beyond this age (>55 years), 8 acute O_3 exposure elicits minimal spirometric changes. Whether the same age-dependent 9 pattern of O_3 sensitivity decline also holds for nonspirometric pulmonary function, 10 airway reactivity or inflammatory endpoints has not been determined. Although there is 11 considerable evidence that spirometric and symptomatic responses to O_3 exposure 12 decrease with age beyond young adulthood, this evidence comes from cross-sectional 13 analyses and has not been confirmed by longitudinal studies of the same individuals.

Sex

14 Several studies have suggested that physiological differences between sexes may 15 predispose females to greater O₃-induced health effects. In females, lower plasma and 16 nasal lavage fluid (NLF) levels of uric acid (the most prevalent antioxidant), the initial 17 defense mechanism of O_3 neutralization in airway surface liquid, may be a contributing 18 factor (Housley et al., 1996). Consequently, reduced absorption of O₃ in the upper 19 airways may promote its deeper penetration. Dosimetric measurements have shown that 20 the absorption distribution of O_3 is independent of sex when absorption is normalized to 21 anatomical dead space (Bush et al., 1996). Thus, a sex-related differential removal of O_3 22 by uric acid seems to be minimal. In general, the physiologic response of young healthy 23 females to O_3 exposure appears comparable to the response of young males (Hazucha et 24 al., 2003). Several studies have investigated the effects of the menstrual cycle on 25 responses to O_3 in healthy young women. In a study of 9 women exposed during exercise 26 to 300 ppb O_3 for an hour, Fox et al. (1993) found lung function responses to O_3 27 significantly enhanced during the follicular phase relative to the luteal phase. However, 28 Weinmann et al. (1995c) found no difference in responses between the follicular and 29 luteal phases as well as no significant differences between 12 males and 12 females 30 exposed during exercise to 350 ppb O_3 for 2.15 hours. Seal et al. (1996) also reported no 31 effect of menstrual cycle phase in their analysis of responses of 150 women (n = 25 per 32 exposure group; 0, 120, 240, 300, and 400 ppb O₃). Seal et al. (1996) conceded that the 33 methods used by Fox et al. (1993) more precisely defined menstrual cycle phase.

Ethnicity

1	Only two controlled human exposure studies have assessed differences in lung function
2	responses between races. Seal et al. (1993) compared lung function responses of whites
3	(93 M, 94 F) and blacks (undefined ancestry; 92 M, 93 F) exposed to a range of O_3
4	concentrations (0-400 ppb). The main effects of the sex-race group and O ₃ concentration
5	were statistically significant (both at $p < 0.001$), although the interaction between sex-race
6	group and O_3 concentration was not significant (p = 0.13). These findings indicate some
7	overall difference between the sex-race groups that is independent of O ₃ concentration,
8	i.e., the concentration-response (C-R) curves for the four sex-race groups are parallel. In
9	a multiple comparison procedure on data collapsed across all O ₃ concentrations for each
10	sex-race group, both black men and black women had significantly larger decrements in
11	FEV_1 than did white men. The authors noted that the O_3 dose per unit of lung tissue
12	would be greater in blacks and females than whites and males, respectively. It cannot be
13	ruled out that this difference in tissue dose might have affected responses to O ₃ . The
14	college students recruited for the Seal et al. (1993) study were noted by the authors as
15	probably being from better educated and SES advantaged families, thus reducing the
16	potential influence of these variables on results. In a follow-up analysis, <u>Seal et al. (1996</u>)
17	reported that, of three SES categories, individuals in the middle SES category showed
18	greater concentration-dependent decline in percent-predicted FEV_1 (4-5% at 400 ppb O_3)
19	than low and high SES groups. The authors did not have an "immediately clear"
20	explanation for this finding.
21	More recently, Que et al. (2011) assessed pulmonary responses in blacks of African
22	American ancestry (22 M, 24 F) and Caucasians (55 M, 28 F) exposed to 220 ppb O_3 for
23	2.25 hours (alternating 15 min periods of rest and brisk treadmill walking). On average,
24	the black males experienced a 16.8% decrement in FEV_1 following O_3 exposure which
25	was significantly larger than mean FEV_1 decrements of 6.2, 7.9, and 8.3% in black
26	females and Caucasian males and Caucasian females, respectively. In the study by <u>Seal et</u>
27	al. (1993), there was potential that the increased FEV_1 decrements in blacks relative to
28	whites were due to increased O ₃ tissue doses since exercise rates were normalized to
29	BSA. Differences in O_3 tissue doses between the races should not have occurred in the

30 31

31 32

Smoking

Que et al. (2011) study because exercise rates were normalized to lung volume (viz.,

attributable to systematically larger O₃ tissue doses in blacks relative to whites.

6-8 times FVC). Thus, the increased mean FEV₁ decrement in black males is not likely

33Smokers are less responsive to O3 for some (but not all) health endpoints than34nonsmokers. Spirometric and plethysmographic pulmonary function decline, respiratory35symptoms, and nonspecific airway hyperreactivity of smokers to O3 were all weaker than

1 data reported for nonsmokers. However, the time course of development and recovery of 2 these effects as well their reproducibility in smokers was not different from nonsmokers

3 (Frampton et al., 1997a). Another similarity between smokers and nonsmokers is that, the

- 4 inflammatory response to O₃ does not appear to depend on smoking status nor the
 - responsiveness of individuals to changes in lung function (Torres et al., 1997). Chronic airway inflammation with desensitization of bronchial nerve endings and an increased production of mucus may plausibly explain the reduced responses to O_3 in smokers

relative to nonsmokers (Frampton et al., 1997a; Torres et al., 1997).

Antioxidant supplementation

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9 The first line of defense against oxidative stress is antioxidants-rich ELF which 10 scavenges free radicals and limits lipid peroxidation. Exposure to O₃ depletes the 11 antioxidant level in nasal ELF probably due to scrubbing of O_3 (Mudway et al., 1999a), 12 however, the concentration and the activity of antioxidant enzymes either in ELF or plasma do not appear to be related to O_3 responsiveness (Samet et al., 2001; Avissar et 13 14 al., 2000; Blomberg et al., 1999). Carefully controlled studies of dietary antioxidant 15 supplementation have demonstrated some protective effects of α -tocopherol and 16 ascorbate on spirometric lung function from O_3 but not on the intensity of subjective 17 symptoms and inflammatory response including cell recruitment, activation and a release 18 of mediators (Samet et al., 2001; Trenga et al., 2001). Dietary antioxidants have also been 19 reported to attenuate O₃-induced bronchial hyperresponsiveness in asthmatics (Trenga et 20 al., 2001).

Genetic polymorphisms

21 Some studies (e.g., <u>Corradi et al., 2002</u> ; <u>Bergamaschi et al., 2001</u>) reviewed in th	ne 2006
O_3 AQCD reported that genetic polymorphisms of antioxidant enzymes may more	dulate
23 pulmonary function and inflammatory response to O ₃ challenge. It was suggested	d that
24 healthy carriers of NAD(P)H:quinone oxidoreductase wild type (NQO1wt) in	
25 combination with glutathione S-transferase μ-1 genetic deficiency (GSTM1null)	were
26 more responsive to O ₃ . <u>Bergamaschi et al. (2001</u>) reported that subjects having N	NQO1wt
and GSTM1null genotypes had increased O ₃ responsiveness (FEV ₁ decrements a	and
28 epithelial permeasibility), whereas subjects with other combinations of these gene	otypes
29 were less affected. A subsequent study from the same laboratory reported a posit	tive
30 association between O ₃ responsiveness, as characterized by the level of oxidative	e stress
31 and inflammatory mediators (8-isoprostane, LTB ₄ and TBARS) in exhaled breat	h
32 condensate and the NQO1wt and GSTM1null genotypes (Corradi et al., 2002). H	Iowever,
33 none of the spirometric endpoints (e.g., FEV_1) were affected by O_3 exposure.	

1	In a controlled exposure of mild-to-moderate asthmatics ($n = 23$; 33 ± 11 years) to
2	300 ppb O_3 for 2 hours with moderate exercise, <u>Vagaggini et al. (2010</u>) found that six of
3	the subjects had a NQO1wt and GSTM1 null, but this genotype was not associated with
4	the changes in lung function or inflammatory responses to O ₃ . Kim et al. (2011) also
5	recently reported that GSTM1 genotype was not predictive of FEV1 responses in young
6	healthy adults (32 F, 27 M; 25.0 \pm 0.5 year) who were roughly half GSTM1-null and half
7	GSTM1-sufficient. Sputum neutrophil levels, measured in a subset of the subjects (13 F,
8	11 M), were also not significantly associated with GSTM1 genotype.
9	In a study of healthy volunteers with GSTM1 sufficient (n = 19; 24 ± 3) and GSTM1 null
10	$(n = 16; 25 \pm 5)$ genotypes exposed to 400 ppb O ₃ for 2 hours with exercise, <u>Alexis et al.</u>
11	(2009) found that inflammatory responses but not lung function responses to O_3 were
12	dependent on genotype. At 4 hours post-O3 exposure, both GSTM1 genotype groups had
13	significant increases in sputum neutrophils with a tendency for a greater increase in
14	GSTM1 sufficient than nulls. At 24 hours postexposure, sputum neutrophils had returned
15	to baseline levels in the GSTM1 sufficient individuals. In the GSTM1 null subjects,
16	however, sputum neutrophil levels increased from 4 hours to 24 hours and were
17	significantly greater than both baseline levels and levels at 24 hours in the GSTM1
18	sufficient individuals. Since there was no FA control in the Alexis et al. (2009) study,
19	effects of the exposure other than O_3 itself cannot be ruled out. In general, the findings
20	between studies are inconsistent.

Body Mass Index

21 In a retrospective analysis of data from 541 healthy, nonsmoking, white males between 22 the ages of 18-35 years from 15 studies conducted at the U.S. EPA Human Studies 23 Facility in Chapel Hill, NC, McDonnell et al. (2010) found that increased BMI was 24 associated with enhanced FEV1 responses to O3. The BMI effect was of the same order of 25 magnitude but in the opposite direction of the age effect where by FEV₁ responses 26 diminish with increasing age. In a similar retrospective analysis, Bennett et al. (2007) 27 found enhanced FEV₁ decrements following O₃ exposure with increasing BMI in a group 28 of 75 healthy, nonsmoking, women (age 24 ± 4 years; BMI range 15.7 to 33.4), but not 29 122 healthy, nonsmoking, men (age 25 ± 4 years; BMI range 19.1 to 32.9). In the women, 30 greater O_3 -induced FEV₁ decrements were seen in overweight (BMI >25) than in normal 31 weight (BMI from 18.5 to 25), and in normal weight than in underweight (BMI <18.5) 32 (P trend \leq 0.022). Together, these results indicate that higher BMI may be a risk factor 33 for pulmonary effects associated with O₃ exposure.

Repeated Ozone Exposure Effects

1	The attenuation of responses observed after repeated consecutive O ₃ exposures in
2	controlled human exposure studies has also been referred to in the literature as
3	"adaptation" or "tolerance" (e.g., Linn et al., 1988). In animal toxicology studies,
4	however, the term tolerance has more classically been used to describe the phenomenon
5	wherein a prior exposure to a low, nonlethal concentration of O_3 provides some
6	protection against death and lung edema at a higher, normally lethal exposure
7	concentration (see Section 9.3.5 of U.S. EPA, 1986). The term "attenuation" will be used
8	herein to refer to the reduction in responses to O_3 observed with repeated O_3 exposures in
9	controlled human exposure studies. Neither tolerance nor attenuation should be presumed
10	to imply complete protection from the biological effects of inhaled O ₃ , because
11	continuing injury still occurs despite the desensitization to some responses.
12	The attenuation of responses due to ambient O ₃ exposure was first investigated by
13	Hackney et al. (1976); (1977a). Experiencing frequent ambient O ₃ exposures, Los
14	Angeles residents were compared to groups having less ambient O3 exposure. Following
15	a controlled laboratory exposure to 370-400 ppb O_3 for 2 hours with light intermittent
16	exercise (2-2.5 times resting \dot{V}_E), the Los Angeles residents exhibited minimal FEV ₁
17	responses relative to groups having less ambient O3 exposure. Subsequently, Linn et al.
18	(1988) examined the seasonal variation in Los Angeles residents' responses to O_3
19	exposure. A group of 8 responders (3M, 5F) and 9 nonresponders (4M, 5F) were exposed
20	to 180 ppb O ₃ for 2 hours with heavy intermittent exercise ($\dot{V}_E = 35 \text{ L/min per m}^2 \text{ BSA}$)
21	on four occasions (spring, fall, winter, and the following spring). In responders, relative
22	to the first spring exposures, FEV_1 responses were attenuated in the fall and winter, but
23	returned to similar decrements the following spring. By comparison, the nonresponders,
24	on average, showed no FEV_1 decrements on any of the four occasions. In subjects
25	recruited regardless of FEV_1 responsiveness to O_3 from the area around Chapel Hill, NC,
26	no seasonal effect of ambient O_3 exposure on FEV_1 responses following chamber
27	exposures to O_3 has been observed (<u>Hazucha et al., 2003</u> ; <u>McDonnell et al., 1985b</u>).
28	Based on studies reviewed in previous O3 AQCDs, several conclusions can be drawn
29	about repeated 1- to 2-h O_3 exposures. Repeated exposures to O_3 causes enhanced
30	(i.e., greater decrements) FVC and FEV_1 responses on the second day of exposure. The
31	enhanced response appears to depend to some extent on the magnitude of the initial
32	response (<u>Horvath et al., 1981</u>). Small responses to the first O_3 exposure are less likely to
33	result in an enhanced response on the second day of O_3 exposure (Folinsbee et al., 1994).
34	With continued daily exposures (i.e., beyond the second day) there is a substantial (or
35	even total) attenuation of pulmonary function responses, typically on the third to
36	fifth days of repeated O_3 exposure. This attenuation of responses is lost in 1 week (<u>Kulle</u>

1	et al., 1982; Linn et al., 1982b) or perhaps 2 weeks (Horvath et al., 1981) without O_3
2	exposure. In temporal conjunction with pulmonary function changes, symptoms induced
3	by O3 (e.g., cough, pain on deep inspiration, and chest discomfort), are also increased on
4	the second exposure day but are attenuated with repeated O_3 exposure thereafter (U.S.
5	EPA, 1998b; Foxcroft and Adams, 1986; Linn et al., 1982b; Folinsbee et al., 1980). In
6	longer-duration (4-6.6 hours), lower-concentration studies that do not cause an enhanced
7	second-day response, the attenuation of response to O3 appears to proceed more rapidly
8	(<u>Folinsbee et al., 1994</u>).

9 Consistent with other investigators, Frank et al. (2001) found FVC and FEV₁ decrements 10 to be significantly attenuated following four consecutive days of exposure to O₃ 11 (250 ppb, 2 h). However, the effects of O_3 on the small airways (assessed by a combined 12 index of isovolumetric forced expiratory flow between 25 and 75% of vital capacity 13 [FEF₂₅₋₇₅] and flows at 50% and 75% of FVC) showed a persistent functional reduction 14 from Day 2 through Day 4. Notably, in contrast to FVC and FEV₁ which exhibited a 15 recovery of function between days, there was a persistent effect of O_3 on small airways 16 function such that the baseline function on Day 2 through Day 4 was depressed relative to 17 Day 1. Frank et al. (2001) also found neutrophil (PMN) numbers in BAL remained 18 significantly higher following O_3 (24 hours after last O_3 exposure) compared to FA. 19 Markers from bronchioalveolar lavage fluid (BALF) following 4 consecutive days of 20 both 2-hour (Devlin et al., 1997) and 4-hour (Jorres et al., 2000; Christian et al., 1998) 21 exposures have indicated ongoing cellular damage irrespective of the attenuation of some 22 cellular inflammatory responses of the airways, lung function and symptoms response. 23 These data suggest that the persistent small airways dysfunction assessed by Frank et al. 24 (2001) is likely induced by both neurogenic and inflammatory mediators, since the 25 density of bronchial C-fibers is much lower in the small than large airways.

Summary of Controlled Human Exposure Studies on Lung Function

26 Responses in humans exposed to ambient O_3 concentrations include: decreased 27 inspiratory capacity; mild bronchoconstriction; rapid, shallow breathing pattern during 28 exercise; and symptoms of cough and pain on deep inspiration (U.S. EPA, 2006b, 1996a). 29 Discussed in subsequent Section <u>6.2.2.1</u> and Section <u>6.2.3.1</u>, exposure to O_3 also results 30 in airway hyperresponsiveness, pulmonary inflammation, immune system activation, and 31 epithelial injury (Que et al., 2011; Mudway and Kelly, 2004a). Reflex inhibition of 32 inspiration results in a decrease in forced vital capacity and, in combination with mild 33 bronchoconstriction, contributes to a decrease in the FEV₁. Healthy young adults exposed 34 to O_3 concentrations ≥ 60 ppb develop statistically significant reversible, transient 35 decrements in lung function and symptoms of breathing discomfort if minute ventilation 36 or duration of exposure is increased sufficiently (Kim et al., 2011; McDonnell et al.,

1	<u>2010; Schelegle et al., 2009; Brown et al., 2008; Adams, 2006a</u>). With repeated O_3
2	exposures over several days, FEV_1 and symptom responses become attenuated in both
3	healthy individuals and asthmatics, but this attenuation of responses is lost after about
4	a week without exposure (Gong et al., 1997a; Folinsbee et al., 1994; Kulle et al., 1982).
5	In contrast to the attenuation of FEV_1 responses, there appear to be persistent O_3 effects
6	on small airways function as well as ongoing cellular damage during repeated exposures.
7	There is a large degree of intersubject variability in lung function decrements
8	(McDonnell, 1996). However, these lung function responses tend to be reproducible
9	within a given individual over a period of several months indicating differences in the
10	intrinsic responsiveness of individuals (Hazucha et al., 2003; McDonnell et al., 1985b).
11	In healthy young adults, O_3 -induced decrements in FEV_1 do not appear to depend on
12	gender (Hazucha et al., 2003), body surface area or height (McDonnell et al., 1997), lung
13	size or baseline FVC (Messineo and Adams, 1990). There is limited evidence that blacks
14	may experience greater O_3 -induced decrements in FEV ₁ than age-matched whites (Que et
15	al., 2011; Seal et al., 1993). Healthy children experience similar spirometric responses
16	but lesser symptoms from O_3 exposure relative to young adults (McDonnell et al.,
17	<u>1985a</u>). On average, spirometric and symptom responses to O_3 exposure appear to decline
18	with increasing age beyond about 18 years of age (McDonnell et al., 1999b; Seal et al.,
19	<u>1996</u>). There is a tendency for slightly increased spirometric responses in individuals
20	with mild asthma and allergic rhinitis relative to healthy young adults (Jorres et al.,
21	<u>1996</u>). Spirometric responses in asthmatics appear to be affected by baseline lung
22	function, i.e., responses increase with disease severity (Horstman et al., 1995).
23	Available information on recovery of lung function following O ₃ exposure indicates that
24	an initial phase of recovery in healthy individuals proceeds relatively rapidly, with acute
25	spirometric and symptom responses resolving within about 2 to 4 hours (Folinsbee and

Hazucha, 1989). Small residual lung function effects are almost completely resolved within 24 h. One day following O_3 exposure, persistent effects on the small airways assessed by decrements in FEF₂₅₋₇₅ and altered ventilation distribution have been reported (Frank et al., 2001; Foster et al., 1997).

6.2.1.2 Epidemiology

30	The O ₃ -induced lung function decrements consistently demonstrated in controlled human
31	exposure studies (Section $6.2.1.1$) provide biological plausibility for the epidemiologic
32	evidence consistently linking short-term increases in ambient O ₃ concentration with lung
33	function decrements in diverse populations. In the 1996 and 2006 O_3 AQCDs, coherence
34	with controlled human exposure study results was found not only for epidemiologic

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1	associations observed in groups with expected higher ambient O3 exposures and higher
2	exertion levels, including children attending summer camps and adults exercising or
3	working outdoors, but also for associations observed in children and individuals with
4	asthma (U.S. EPA, 2006b, 1996a). Recent epidemiologic studies focused more on
5	children with asthma rather than groups with increased outdoor exposures or other
6	healthy populations. Whereas recent studies contributed less consistent evidence, the
7	cumulative body of evidence indicates decreases in lung function in association with
8	increases in ambient O ₃ concentration in children with asthma. Collectively, studies in
9	adults with asthma and individuals without asthma found both O3-associated decreases
10	and increases in lung function. Recent studies did provide additional data to assess
11	whether particular lags of O3 exposure were more strongly associated with decrements in
12	lung function; whether O3 associations were confounded by copollutant exposures; and
13	whether associations were modified by factors such as corticosteroid (CS) use, genetic
14	polymorphisms, and elevated BMI.

Populations with Increased Outdoor Exposures

15 Epidemiologic studies primarily use ambient O_3 concentrations to represent exposure; 16 however, few studies have accounted for time spent outdoors, which has been shown to 17 influence the relationship between ambient concentrations and individual exposures to O_3 18 (Section 4.3.3). Epidemiologic studies of individuals engaged in outdoor recreation, 19 exercise, or work are noteworthy for the likely greater extent to which ambient O_3 concentrations represent ambient O₃ exposures. Ambient O₃ concentrations, locations, 20 21 and time periods for epidemiologic studies of populations with increased outdoor 22 exposures are presented in Table 6-2. Most of these studies measured ambient O_3 at the 23 site of subjects' outdoor activity and related lung function changes to the O₃ 24 concentrations measured during outdoor activity, which have contributed to higher O_3 25 personal exposure-ambient concentration correlations and ratios (Section 4.3.3). Because 26 of improved O₃ exposure estimates, measurement of lung function before and after 27 discrete periods of outdoor activity, and examination of O₃ effects during exertion when 28 the dose of O₃ reaching the lungs may be higher due to higher ventilation and inhalation 29 of larger volumes of air, epidemiologic studies of populations with increased outdoor 30 exposures are more comparable to controlled human exposure studies. Further, these 31 epidemiologic studies provide strong evidence for respiratory effects in children and 32 adults related to ambient O₃ exposure. Similar to findings from controlled human 33 exposure studies, the collective body of epidemiologic evidence clearly demonstrates 34 decrements in lung function in association with increases in ambient O₃ exposure during 35 periods of outdoor activity (Figure 6-3 to Figure 6-5 and Table 6-3 to Table 6-5).

Table 6-2Mean and upper percentile ozone concentrations in epidemiologic
studies of lung function in populations with increased outdoor
exposures.

Study*	Location	Study Period	O ₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Thurston et al.</u> (1997)	Connecticut River Valley, CT	June 1991-1993	1-h max	83.6	Max: 160
<u>Berry et al.</u> (1991)	Mercer County, NJ	July 1988	1-h max ^ª	NR	Max: 204
<u>Spektor and</u> Lippmann (1991)	Fairview Lake, NJ	July-August 1988	1-h avg ^b	69	Max: 137
<u>Avol et al.</u> (1990)	ldyllwild, CA	June-August 1988	1-h avg⁵	94	Max: 161
<u>Burnett et al.</u> (<u>1990</u>)	Lake Couchiching, Ontario, Canada	June-July 1983	1-h avg ^b	59	Max: 95
<u>Higgins et al.</u> (1990)	San Bernardino, CA	June-July 1987	1-h avg⁵	123	Max: 245
Raizenne et al. (1989)	Lake Erie, Ontario, Canada	July-August 1986	1-h avg ^b	71	Max: 143
<u>Spektor et al.</u> (<u>1988a</u>)	Fairview Lake, NJ	July-August 1984	1-h avg⁵	53	Max: 113
<u>Neas et al.</u> (1999)	Philadelphia, PA	July-September 1993	12-h avg ^a (9 a.m 9 p.m.)	57.5 (near Camp 1) 55.9 (near Camp 2)	Max (near Camp 1): 106
<u>Nickmilder et</u> al. (2007)	Southern Belgium	July-August 2002	1-h max 8-h max	NR	Max (across 6 camps): 24.5-112.7° Max (across 6 camps): 18.9-81.1°
<u>Girardot et al.</u> (<u>2006</u>)	Great Smoky Mountain NP, TN	August-October 2002 June- August 2003	Hike-time avg (2-9 h) ^d	48.1	Max: 74.2
<u>Korrick et al.</u> (1998)	Mt. Washington, NH	Summers 1991, 1992	Hike-time avg (2-12 h) ^d	40	Max: 74
<u>Hoppe et al.</u> (2003)	Munich, Germany	Summers 1992- 1995	30-min max (1-4 p.m.)	High O_3 days: 65.9 Control O_3 days: 27.2	Max (high O ₃ days): 86
<u>Spektor et al.</u> (1988b)	Tuxedo, NY	June-August 1985	Exercise-time avg (15 - 55 min)	NR	Max: 124
<u>Selwyn et al.</u> (<u>1985</u>)	Houston, TX	May-October 1981	Exercise-time 15- min max (4-7 p.m.)	47	Max: 135
<u>Brunekreef et</u> al. (1994)	Eastern Netherlands	June-August 1981	Exercise-time avg ^a (10-145 min)	42.8 [°]	Max: 99.5°
<u>Braun-</u> F <u>ahrlander et</u> al. <u>(1994</u>)	Southern Switzerland	May-October 1989	Exercise-time 30-min avg	NR	Max: 80°
<u>Castillejos et</u> al. <u>(1995</u>)	Mexico City, Mexico	June 1990- October 1991	1-h max ^a	179	Max: 365
<u>Hoek et al.</u> (<u>1993</u>)	Wageningen, Netherlands	May-July 1989	1-h max ^ª	NR	Max: 122 ^c

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Hoppe et al.</u> (1995)	Munich, Germany	Summers 1992- 1995	30-min max (1-4 p.m.)	High O_3 days: 64 Control O_3 days: 32	Max (high O_3 days): 77
<u>Chan and Wu</u> (2005)	Taichung City, Taiwan	November- December 2001	8-h avg (9 a.m 5 p.m.) 1-h max	35.6 52.6	Max: 65.1 95.5
<u>Brauer et al.</u> (1996)	British Columbia, Canada	June-August 1993	1-h max ^a	40	Max: 84
<u>Romieu et al.</u> (1998b)	Mexico City, Mexico	March-August 1996	Work-shift avg (6 - 12 h) ^a	67.3	95th: 105.8
<u>Thaller et al.</u> (2008)	Galveston, TX	Summer 2002- 2004	1-h max	35 (median)	Max: 118

* Note: Studies presented in order of first appearance in the text of this section.

NR = not reported.

^aSome or all measurements obtained from monitors located off site of outdoor activity.

^b1-h avg, preceding lung function measurement, as reported in the pooled analysis by Kinney et al. (1996).

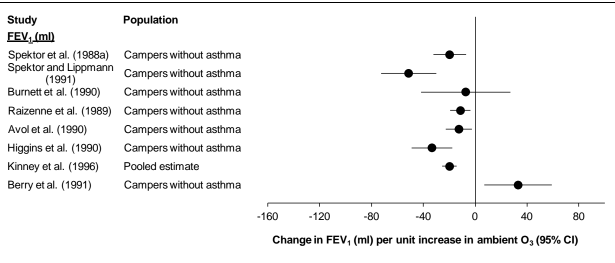
 $^{\circ}$ Concentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25 $^{\circ}$ C) and pressure (1 atm).

^dIndividual-level estimates calculated from concentrations measured in different segments of hiking trail.

Children Attending Summer Camps

1	Studies of children attending summer camps, most of which were discussed in the 1996
2	O ₃ AQCD, have provided important evidence of the impact of ambient O ₃ exposure on
3	respiratory effects in young, healthy children. In addition to the improved exposure
4	assessment as described above, these studies were noted for their daily assessment of
5	lung function by trained staff over 1- to 2-week periods in the mornings and late
6	afternoons before and after hours of outdoor activity (Thurston et al., 1997; Berry et al.,
7	1991; Spektor and Lippmann, 1991; Avol et al., 1990; Burnett et al., 1990; Higgins et al.,
8	<u>1990; Raizenne et al., 1989; Spektor et al., 1988a).</u>
9	In groups mostly comprising healthy children (ages 7-17 years), decrements in FEV_1
10	were associated consistently with increases in ambient O ₃ concentration averaged over
11	the 1-12 hours preceding lung function measurement (Figure 6-3 and Table 6-3). Kinney
12	et al. (1996) corroborated this association in a re-analysis combining 5,367 lung function
13	measurements collected from 616 healthy children from six studies (Spektor and
14	Lippmann, 1991; Avol et al., 1990; Burnett et al., 1990; Higgins et al., 1990; Raizenne et
15	al., 1989; Spektor et al., 1988a). Based on uniform statistical methods, a -20 ml (95% CI:
16	-25, -14) change in afternoon FEV_1 was estimated for a 40-ppb increase in O_3
17	concentration averaged over the 1 hour before lung function measurement (Kinney et al.,
18	<u>1996</u>) (all effect estimates are standardized to increments specific to the O_3 averaging
19	time as detailed in Section 2.1). All of the studies in the pooled analysis were conducted

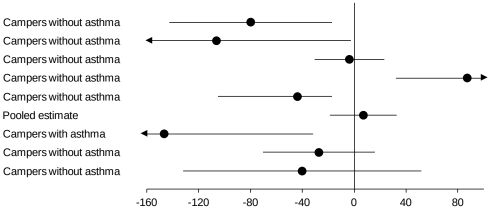
1	during summer months but were diverse in locations examined (i.e., Northeast U.S.,
2	Canada, California), range in ambient concentrations of O_3 (presented within Table 6-2)
3	and other pollutants measured, and magnitudes of association observed. Study-specific
4	effect estimates ranged between a 0.76 and 48 mL decrease or a 0.3% to 2.2% decrease in
5	study mean FEV ₁ per 40-ppb increase in 1-h avg O_3 .
6	Among camp studies included the pooled analysis plus others, associations for peak
7	expiratory flow (PEF) were more variable than were those for FEV_1 , as indicated by the
8	wider range in effect estimates and wider 95% CIs (Figure 6-3 and Table 6-3).
9	Nonetheless, in most cases, increases in ambient O3 concentration were associated with
10	decreases in PEF. The largest O3-associated decrease in PEF (mean 2.8% decline per
11	40-ppb increase in 1-h max O ₃) was found in a group of campers with asthma, in whom
12	an increase in ambient O ₃ concentration also was associated with increases in chest
13	symptoms and bronchodilator use (Thurston et al., 1997).
14	For both FEV_1 and PEF, the magnitude of association was not related to the study mean
15	ambient 1-h avg or max O_3 concentration. With exclusion of results from <u>Spektor and</u>
16	Lippmann (1991), larger O_3 -associated FEV ₁ decrements were found in populations with
17	lower mean FEV ₁ . No trend was found with mean PEF. Sufficient data were not provided
18	to assess whether the temporal variability in O_3 concentrations, activity levels of subjects,
19	or associations with other pollutants contributed to between-study heterogeneity in O_3
20	effect estimates.





Spektor et al. (1988a)
Burnett et al. (1990)
Raizenne et al. (1989)
Avol et al. (1990)
Higgins et al. (1990)
Kinney et al. (1996)
Thurston et al. (1997)
Neas et al. (1999)
Berry et al. (1991)

Campers without asthma Campers without asthma Campers without asthma Campers without asthma Pooled estimate Campers with asthma Campers without asthma Campers without asthma



Change in PEF (ml/sec) per unit increase in ambient O₃ (95% Cl)

Note: Results generally are presented in order of increasing mean ambient O₃ concentration. Effect estimates are from singlepollutant models and are standardized to a 40-ppb increase for 1-h avg or 1-h max O₃ concentration and a 30-ppb increase for 12-h avg O₃ concentration.

Changes in FEV₁ (mL) or PEF (mL/sec) in association with ambient Figure 6-3 ozone concentrations among children attending summer camp.

Study	Location	Population, Mean FEV₁ (mL) or PEF (mL/sec)	Standardized Percent Change (95% CI) ^a	Standardized Effect Estimate (95% CI) ^a
FEV ₁				(mL)
<u>Spektor et al.</u> (1988a)	Fairview Lake, NJ	91 campers without asthma ages 8-15 yr, 2,140	-0.93 (-1.5, -0.35) ^b	-20.0 (-32.5, -7.5) ^b
<u>Spektor and</u> Lippmann (1991)	Fairview Lake, NJ	46 campers without asthma ages 8-14 yr, 2,390	-2.2 (-3.0, -1.3) ^b	-51.6 (-72.8, -30.4) ^b
<u>Burnett et al.</u> (1990)	Lake Couchiching, Ontario, Canada	29 campers without asthma ages 7-15 yr, 2,410	-0.32 (-1.7, 1.1) ^b	-7.6 (-42.1, 26.9) ^b
<u>Raizenne et al.</u> (1989)	Lake Erie, Ontario, Canada	112 campers without asthma mean age 11.6 yr, 2,340	-0.50 (-0.83, -0.16) ^b	-11.6 (-19.4, -3.8) ^b
<u>Avol et al. (1990</u>)	Pine Springs, CA	295 campers without asthma ages 8-17 yr, 2,190	-0.58 (-1.0, -0.12) ^b	-12.8 (-23.0, -2.6) ^b
<u>Higgins et al.</u> (1990)	San Bernardino, CA	43 campers without asthma ages 7-13 yr, 2,060	-1.6 (-2.4, -0.87) ^b	-33.6 (-49.3, -17.9) ^b
<u>Kinney et al.</u> (1996)	Pooled analysis of preceding 6 studies	616 campers without asthma ages 7-17 yr, 2,300	-0.87 (-1.1, -0.63)	-20.0 (-25.5, -14.5) ^b
<u>Berry et al. (1991)</u>	Hamilton, NJ	14 campers without asthma 58% age <14 yr, NA	NA	32.8 (6.9, 58.7)
PEF				(mL/sec)
<u>Spektor et al.</u> (1988a)	Fairview Lake, NJ	91 campers without asthma ages 8-15 yr, 4,360	-1.8 (-3.3, -0.40)	-80.0 (-142.7, -17.3) ^b
<u>Burnett et al.</u> (1990)	Lake Couchiching, Ontario, Canada	29 campers without asthma ages 7-15 yr, 5,480	-1.9 (-3.8, -0.05)	-106.4 (-209.9, -2.9) ^b
<u>Raizenne et al.</u> (1989)	Lake Erie, Ontario, Canada	112 campers without asthma mean age 11.6 yr, 5,510	-0.07 (-0.56, 0.41)	-4.0 (-30.7, 22.7) ^b
<u>Avol et al. (1990</u>)	Pine Springs, CA	295 campers without asthma ages 8-17 yr, 4,520	1.9 (0.71, 3.1)	86.8 (31.9, 142) ^b
<u>Higgins et al.</u> (1990)	San Bernardino, CA	43 campers without asthma ages 7-13 yr, 5,070	-0.87 (-2.1, 0.34)	-44.0 (-105, 17.2) ^b
<u>Kinney et al.</u> (1996)	Pooled analysis of preceding 6 studies	616 campers without asthma ages 7-17 yr, 4,222	0.31 (-0.88, 1.5)	6.8 (-19.1, 32.7) ^b
<u>Thurston et al.</u> (1997)	CT River Valley, CT	166 campers with asthma ages 7-13 yr, 5,333	-2.8 (-4.9, -0.59)	-146.7 (-261.7, -31.7)
<u>Neas et al. (1999</u>)	Philadelphia, PA	156 campers without asthma ages 6-11 yr, 4,717	-0.58 (-1.5, 0.33)	-27.5 (-70.8, 15.8)
Berry et al. (1991)	Hamilton, NJ	14 campers without asthma 58% age <14 yr, NA	NA	-40.4 (-132.1, 51.3)

Table 6-3Additional characteristics and quantitative data for studies
represented in Figure 6-3.

*Includes studies form Figure 6-3.

NA = Data not available.

^aAll results are standardized to a 40-ppb increase in 1-h avg or 1-h max O_3 , except that from <u>Neas et al. (1999</u>), which is standardized to a 30-ppb increase in 12-h avg (9 a.m.-9 p.m.) O_3 .

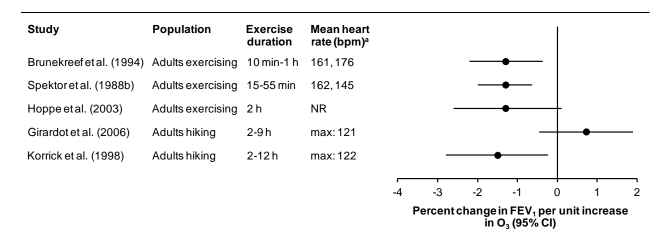
^bEffect estimates were reported in the pooled analysis by Kinney et al. (1996).

1	Similar to controlled human exposure studies, camp studies found interindividual
2	variability in the magnitude of O3-associated changes in lung function. Based on separate
3	regression analyses of data from individual subjects, increases in ambient O ₃
4	concentration were associated with a wide range of changes in lung function across
5	subjects (Berry et al., 1991; Higgins et al., 1990; Spektor et al., 1988a). For example,
6	among children attending camp in Fairview Lake, NJ, 36% of subjects had statistically
7	significant O_3 -associated decreases in FEV ₁ , and the 90th percentile of response was a
8	6.3% decrease in FEV ₁ per a 40-pbb increase in 1-h avg O_3 (Spektor et al., 1988a).
9	In contrast with previous studies, a recent study of children attending six different
	in contrast with providers, a recent starty of enhancing shi anterent
10	summer camps in Belgium did not find an association between ambient O_3 concentration
10 11	
	summer camps in Belgium did not find an association between ambient O ₃ concentration
11	summer camps in Belgium did not find an association between ambient O_3 concentration and lung function (<u>Nickmilder et al., 2007</u>). This study examined similar ambient O_3
11 12	summer camps in Belgium did not find an association between ambient O_3 concentration and lung function (<u>Nickmilder et al., 2007</u>). This study examined similar ambient O_3 concentrations as did previous studies (<u>Table 6-2</u>) but used a less rigorous methodology.
11 12 13	summer camps in Belgium did not find an association between ambient O_3 concentration and lung function (<u>Nickmilder et al., 2007</u>). This study examined similar ambient O_3 concentrations as did previous studies (<u>Table 6-2</u>) but used a less rigorous methodology. Lung function was measured only once in each subject, and mean lung function was

Populations Exercising Outdoors

18outdoors have provided evidence for lung function decrements in healthy adults19associated with increases in ambient O_3 exposure during exercise with durations (10 min20to 12 h) and intensities (heart rates 121-190 beats per min) in the range of those examined21in controlled human exposure studies (Table 6-1). As in the camp studies, lung function22was measured before and after exercise by trained staff. Collectively, studies of adults23found FEV1 decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O_3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3 27concentration also were associated with decreases in lung function in children exercising28outdoors (Table 6-4).	17	As discussed in the 1996 and 2006 O_3 AQCDs, epidemiologic studies of adults exercising
20to 12 h) and intensities (heart rates 121-190 beats per min) in the range of those examined21in controlled human exposure studies (Table 6-1). As in the camp studies, lung function22was measured before and after exercise by trained staff. Collectively, studies of adults23found FEV1 decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O_3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3 27concentration also were associated with decreases in lung function in children exercising	18	outdoors have provided evidence for lung function decrements in healthy adults
21in controlled human exposure studies (Table 6-1). As in the camp studies, lung function22was measured before and after exercise by trained staff. Collectively, studies of adults23found FEV1 decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O_3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3 27concentration also were associated with decreases in lung function in children exercising	19	associated with increases in ambient O3 exposure during exercise with durations (10 min
22was measured before and after exercise by trained staff. Collectively, studies of adults23found FEV1 decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O_3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3 27concentration also were associated with decreases in lung function in children exercising	20	to 12 h) and intensities (heart rates 121-190 beats per min) in the range of those examined
23found FEV_1 decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O_3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3 27concentration also were associated with decreases in lung function in children exercising	21	in controlled human exposure studies (Table 6-1). As in the camp studies, lung function
24Table 6-4). The magnitude of association did not appear to be related to study mean25ambient O3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O327concentration also were associated with decreases in lung function in children exercising	22	was measured before and after exercise by trained staff. Collectively, studies of adults
25ambient O3 concentrations (Table 6-2), exercise duration, or the mean heart rate26measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O327concentration also were associated with decreases in lung function in children exercising	23	found FEV ₁ decrements of 1.3 to 1.5% per unit increase in O_3^{-1} (Figure 6-4 and
 26 measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O₃ 27 concentration also were associated with decreases in lung function in children exercising 	24	Table 6-4). The magnitude of association did not appear to be related to study mean
27 concentration also were associated with decreases in lung function in children exercising	25	ambient O_3 concentrations (<u>Table 6-2</u>), exercise duration, or the mean heart rate
	26	measured during exercise (Figure 6-4 and Table 6-4). Increases in ambient O_3
28 outdoors (<u>Table 6-4</u>).	27	concentration also were associated with decreases in lung function in children exercising
	28	outdoors (<u>Table 6-4</u>).

¹Effect estimates were standardized to a 40-ppb increase in O_3 averaged over 15 min to 1 h and a 30-ppb increase for O_3 averaged over 2 to 12 h.



Note: Studies generally are presented in order of increasing duration of outdoor exercise. Data refer to the maximum or mean measured during exercise or in different groups or conditions as described in Table 6-4.

^abpm = beats per minute. NR = Not reported. Effect estimates are from single-pollutant models and are standardized to a 40-ppb increase for O_3 concentrations averaged over 15 minutes to 1 hour and a 30-ppb increase for O_3 concentrations averaged over 2 to 12 hours.

Figure 6-4 Percent change in FEV₁ in association with ambient ozone concentrations among adults exercising outdoors.

Table 6-4Additional characteristics and quantitative data for studies
represented in Figure 6-4 plus results from studies in children
exercising outdoors.

Study*	Location	Population	Exercise Duration, Mean Heart Rate	O₃ Averaging Time	Parameter	Standardized Percent Change (95% CI) ^a
Studies of a	dults					
<u>Brunekreef et</u> al. (1994)	Netherlands	29 adults exercising, ages 18-37 yr	10 min - 2.4 h, HR: 161 bpm (training), 176 bpm (races)	Exercise duration	FEV₁ PEF	-1.3 (-2.2, -0.37) -2.5 (-3.8, -1.2)
<u>Spektor et al.</u> (1988b)	Tuxedo, NY	30 adults exercising, ages 21-44 yr	15 - 55 min, HR:162 bpm if $\dot{V}_{\rm E}$ >100 L, 145 bpm if $\dot{V}_{\rm E}$ 60-100 L	30-min avg	FEV ₁	-1.31 (-2.0, -0.65)
<u>Hoppe et al.</u> (2003)	Munich, Germany	43 adults and children exercising, ages 13-38 yr	2 h, HR: NR	30-min max (1-4 p.m.)	FEV₁ PEF	-1.3 (-2.6, 0.10) -2.8 (-5.9, 0.31)
<u>Girardot et al.</u> (2006)	Great Smoky Mt, TN	354 adult day hikers, ages 18- 82 yr	2-9 h, max HR:121 bpm	Hike duration	FEV₁ PEF	0.72 (-0.46, 1.90) 3.5 (-0.11, 7.2)
<u>Korrick et al.</u> (1998)	Mt. Washington, NH	530 adult day hikers, ages 18- 64 yr	2-12 h, max HR: 122 bpm	Hike duration	FEV₁ PEF	-1.5 (-2.8, -0.24) -0.54 (-4.0, 2.9)
<u>Selwyn et al.</u> <u>(1985</u>)	Houston, TX	24 adults exercising, ages 29-47 yr	Duration: NR, max HR: 179 bpm in males, 183 bpm in females	15-min max	FEV ₁	-16 mL (-28.8, -3.2) ^b
Studies of cl	nildren not incl	uded in Figure 6-4				
<u>Braun-</u> <u>Fahrlander et</u> al. (1994)	Switzerland	128 children exercising, ages 9-11 yr	10 min, max HR: 180 bpm	30-min avg	PEF	-3.8 (-6.7, -0.96)
<u>Castillejos et</u> <u>al. (1995</u>)	Mexico City, Mexico	40 children exercising, ages 7- 11 yr	2 15 min with 15 min rest periods, max HR: <190 bpm	1-h avg over full exercise-rest period	FEV ₁	-0.48 (-0.72, -0.24)
<u>Hoek et al.</u> (1993)	Wageningen, Netherlands	65 children exercising, ages 7- 12 yr	25 min-1.5 h, HR: NR	1-h max	PEF	-2.2 (-4.9, 0.54)
*Includes studie	es from Figure 6-4	<u>,</u> plus others.				

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HR = heart rate, bpm = beats per minute, \dot{V}_E = minute ventilation, NR = Not reported.

^aEffect estimates are standardized to a 40-ppb increase for O_3 concentrations averaged over 15 minutes to 1 hour and a 30-ppb increase for O_3 concentrations averaged over 2 to 12 hours.

^bResults not included in the figure because data were not provided to calculate percent change in lung function.

1Compared with the studies of individuals exercising outdoors described above, analyses2of day-hikers assessed lung function only on one day per subjects but examined longer3periods of outdoor activity and included much larger sample sizes. Studies of adult day-4hikers had similar design but produced contrasting results (Girardot et al., 2006; Korrick5et al., 1998). Among 530 hikers on Mt. Washington, NH, Korrick et al. (1998) reported

1	posthike declines in FEV ₁ and FVC of 1.5% and 1.3%, respectively, per a 30-ppb
2	increase in 2- to 12-h avg O ₃ . Associations with FEV ₁ /FVC, FEF _{25-75%} , and PEF were
3	weaker. In contrast, among 354 hikers on Great Smoky Mt, TN, Girardot et al. (2006)
4	found that higher O ₃ concentrations were associated with posthike increases in many of
5	the same lung function indices (Figure 6-4 and Table 6-4). These studies were similar in
6	the examination of a mostly white, healthy population and of changes in lung function
7	associated with ambient O ₃ concentrations measured on site during multihour (2-12 h)
8	periods of outdoor exercise. Mean O ₃ concentrations were similar as were the population
9	mean and variability in lung function. However, Girardot et al. (2006) differed from
10	Korrick et al. (1998) in several aspects, including a shorter hike time (maximum: 9 versus
11	12 h), older age of subjects (maximum: 82 versus 64 yr), and measurement of lung
12	function by a larger number of less well-trained technicians. The impact of these
13	differences on the heterogeneity in results between the studies was not examined.
14	Similar to the camp studies, some studies of outdoor exercise examined and found
15	interindividual variability in the magnitude of O3-associated decreases in lung function.
16	In Korrick et al. (1998), although a 30-ppb increase in 2- to 12-h avg ambient O_3
17	concentration was associated with a group mean decrement in $\text{FEF}_{25-75\%}$ of -0.81%
18	(95% CI: -4.9, 3.3), some individuals experienced a $>10\%$ decline. The odds of $>10\%$
19	decline in FEF _{25-75%} increased with increasing ambient O ₃ concentration (OR: 2.3
20	[95% CI: 1.2, 6.7] per 30-ppb increase in 2- to 12-h avg O ₃). Likewise, Hoppe et al.
21	(2003) found that compared with days with 30-min max (1-4 p.m.) ambient O_3
22	concentrations <40 ppb, on days with $\Omega_2 > 50$ ppb, 14% of athletes had at least a 10%

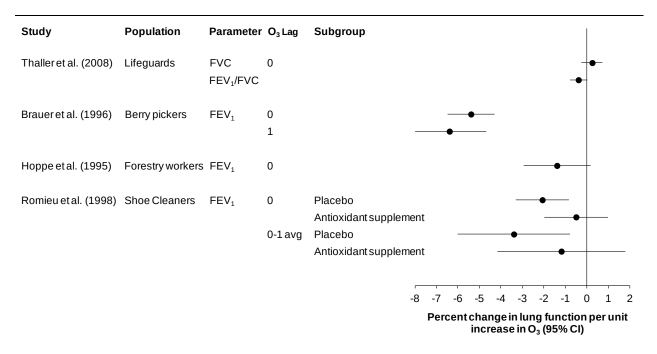
concentrations <40 ppb, on days with O₃ >50 ppb, 14% of athletes had at least a 10%
 decrease in lung function or 20% increase in airway resistance.

Outdoor Workers

24 Consistent findings in outdoor workers add to the evidence that short-term increases in 25 ambient O_3 exposure decrease lung function in healthy adults (Figure 6-5 and Table 6-5). 26 Except for Hoppe et al. (1995), studies used central site ambient O₃ concentrations. 27 However, in outdoor workers, ambient concentrations have been more highly correlated 28 with and similar in magnitude to personal exposures (Section 4.3.3) likely because 29 workers spend long periods of time outdoors (6-14 hours across studies) and the O₃ 30 averaging times examined correspond to periods of outdoor work. For example, in a 31 subset of berry pickers, the correlation and ratio of personal to ambient 24-h avg O₃ 32 concentrations (15 km from work site) were 0.64 and 0.96, respectively (Brauer and 33 Brook, 1997). The 6-h avg personal-ambient ratio in a population of shoe cleaners in 34 Mexico City was 0.56 (O'Neill et al., 2003). Many studies of outdoor workers found that 35 in addition to same-day concentrations, O3 concentrations lagged 1 or 2 days (Chan and

2associated with equal or larger decrements in lung function (Figure 6-5 and Table 6-5).3Similar to other populations with increased outdoor exposure, most of the magnitudes of O ₃ -associated lung function decrements in outdoor workers were small, i.e., <1% to 3.4% per unit increase in O ₃ concentration ¹ . The magnitude of decrease was not found to depend strongly on duration of outdoor work or ambient O ₃ concentration. The largest decrease (6.4% per 40-ppb increase in 1-h max O ₃) was observed in berry pickers in British Columbia who were examined during a period of relatively low ambient O ₃ concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a much smaller O ₃ -associated decrease in FEV1 was found in shoe cleaners in Mexico City who were examined during a period of higher O ₃ concentrations (work shift mean: 13 67.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon IFEV1/FVC per 40-ppb increase in 1-h max O ₃) was observed in lifeguards in Galveston, TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry pickers but characterized by a similar range of ambient O ₃ concentrations. Not all studies provided information on ventilation rate or pulse rate, thus it was not possible to ascertain whether differences in the magnitude of O ₃ -associated lung function decrement across studies were related to differences in the level of exertion of workers.	1	Wu, 2005; Brauer et al., 1996) or averaged over 2 days (Romieu et al., 1998b) were
4 O_3 -associated lung function decrements in outdoor workers were small, i.e., <1% to 3.4%5per unit increase in O_3 concentration ¹ . The magnitude of decrease was not found to6depend strongly on duration of outdoor work or ambient O_3 concentration. The largest7decrease (6.4% per 40-ppb increase in 1-h max O_3) was observed in berry pickers in8British Columbia who were examined during a period of relatively low ambient O_3 9concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O_3 -associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O_3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O_3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O_3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O_3 -associated lung function decrement across	2	associated with equal or larger decrements in lung function (Figure 6-5 and Table 6-5).
5per unit increase in O3 concentration ¹ . The magnitude of decrease was not found to6depend strongly on duration of outdoor work or ambient O3 concentration. The largest7decrease (6.4% per 40-ppb increase in 1-h max O3) was observed in berry pickers in8British Columbia who were examined during a period of relatively low ambient O39concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O3-associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	3	Similar to other populations with increased outdoor exposure, most of the magnitudes of
6depend strongly on duration of outdoor work or ambient O_3 concentration. The largest7decrease (6.4% per 40-ppb increase in 1-h max O_3) was observed in berry pickers in8British Columbia who were examined during a period of relatively low ambient O_3 9concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O_3 -associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O_3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O_3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O_3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O_3 -associated lung function decrement across	4	O_3 -associated lung function decrements in outdoor workers were small, i.e., <1% to 3.4%
7decrease (6.4% per 40-ppb increase in 1-h max O3) was observed in berry pickers in8British Columbia who were examined during a period of relatively low ambient O39concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O3-associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	5	per unit increase in O_3 concentration ¹ . The magnitude of decrease was not found to
8British Columbia who were examined during a period of relatively low ambient O39concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O3-associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	6	depend strongly on duration of outdoor work or ambient O ₃ concentration. The largest
9concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O_3 -associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O_3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O_3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O_3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O_3 -associated lung function decrement across	7	decrease (6.4% per 40-ppb increase in 1-h max O ₃) was observed in berry pickers in
10outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a11much smaller O_3 -associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O_3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O_3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O_3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O_3 -associated lung function decrement across	8	British Columbia who were examined during a period of relatively low ambient O_3
11much smaller O_3 -associated decrease in FEV1 was found in shoe cleaners in Mexico City12who were examined during a period of higher O_3 concentrations (work shift mean:1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O_3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O_3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O_3 -associated lung function decrement across	9	concentrations (work shift mean: 26.0 ppb [SD: 11.8]) but had longer daily periods of
 who were examined during a period of higher O₃ concentrations (work shift mean: 67.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon FEV₁/FVC per 40-ppb increase in 1-h max O₃) was observed in lifeguards in Galveston, TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry pickers but characterized by a similar range of ambient O₃ concentrations. Not all studies provided information on ventilation rate or pulse rate, thus it was not possible to ascertain whether differences in the magnitude of O₃-associated lung function decrement across 	10	outdoor work (8-14 hours) (Brauer et al., 1996) (Figure 6-5 and Table 6-5). However, a
1367.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	11	much smaller O3-associated decrease in FEV1 was found in shoe cleaners in Mexico City
14pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	12	who were examined during a period of higher O ₃ concentrations (work shift mean:
15FEV1/FVC per 40-ppb increase in 1-h max O3) was observed in lifeguards in Galveston,16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	13	67.3 ppb [SD: 24]) but had period of outdoor work that was as long as that of the berry
16TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	14	pickers. The smallest magnitude of decrease (-0.4% [95% CI: -0.8, 0] in afternoon
17pickers but characterized by a similar range of ambient O3 concentrations. Not all studies18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	15	FEV ₁ /FVC per 40-ppb increase in 1-h max O ₃) was observed in lifeguards in Galveston,
18provided information on ventilation rate or pulse rate, thus it was not possible to ascertain19whether differences in the magnitude of O3-associated lung function decrement across	16	TX (Thaller et al., 2008) whose outdoor work periods were shorter than those of the berry
19 whether differences in the magnitude of O_3 -associated lung function decrement across	17	pickers but characterized by a similar range of ambient O ₃ concentrations. Not all studies
	18	provided information on ventilation rate or pulse rate, thus it was not possible to ascertain
20 studies were related to differences in the level of exertion of workers.	19	whether differences in the magnitude of O3-associated lung function decrement across
	20	studies were related to differences in the level of exertion of workers.

 $^{^{1}}$ Effect estimates were standardized to a 40-ppb increase for O₃ averaged over 30 min to 1 h and a 30-ppb increase for O₃ averaged over 8 or 12 h.



Note: Studies generally are presented in order of increasing mean ambient O_3 concentration. Effect estimates are from single-pollutant models and are standardized to a 40-ppb increase for 30-min, 1-h avg, or 1-h max O_3 concentrations.

Figure 6-5 Percent change in FEV₁ or FEV₁/FVC in association with ambient ozone concentrations among outdoor workers.

Study*	Location	Population	Parameter	Duration of Outdoor Work	O₃ Averaging Time	O₃ Lag	Subgroup	Standardized Percent Change (95% CI) ^a
<u>Thaller</u> <u>et al.</u> (2008)	Galveston, TX	142 lifeguards, ages 16-27 yr	FVC	6-8 h	1-h max 12-h avg	0		0.24 (-0.28, 0.72) 0.15 (-0.06, 0.36)
			FEV ₁ /FVC	-	1-h max 12-h avg	-		-0.40 (-0.80, 0) -0.60 (-1.2, 0)
<u>Brauer</u> <u>et al.</u> (1996)	British Columbia, Canada	58 berry pickers, ages 10-69 yr	FEV ₁	8-14 h	1-h max	0 1		-5.4 (-6.5, -4.3) -6.4 (-8.0, -4.7)
<u>Hoppe et</u> <u>al.</u> (1995)	Munich, Germany	41 forestry workers, ages 20-60 yr	FEV ₁	Not reported	30-min max (1 - 4 p.m.)	0		-1.4 (-3.0, 0.16)
<u>Romieu</u> <u>et al.</u> (1998b)	Mexico City, Mexico	47 male shoe cleaners, mean (SD) age: 38.9	FEV ₁	Mean (SD): 9 h (1)	1-h avg before lung function	0	Placebo Antioxidant	-2.1 (-3.3, -0.85) -0.52 (-2.0, 0.97)
<u>(19900</u>)		(3D) age. 38.9 (10) yr			measurement	0-1 avg	Placebo Antioxidant	-3.4 (-6.0, -0.78) -1.2 (-4.2, 1.8)
<u>Chan</u> <u>and Wu</u> (2005) ^b	Taichung City, Taiwan	43 mail carriers. Mean (SD) age: 39 (8) yr	PEF	8 h	1-h max	0 1		-1.3 (-1.7, -0.92) -1.4 (-1.7, -1.2)
(2000)		00 (0) yi			8-h avg (9 a.m 5 p.m.)	0 1		-1.6 (-2.2, -1.1) -1.9 (-2.5, -1.3)

Table 6-5Additional characteristics and quantitative data for studies
represented in Figure 6-5.

*Includes studies from Figure 6-5, plus others.

^aEffect estimates are standardized to a 40-ppb increase for 30-min, 1-h avg, or 1-h max O₃ and a 30-ppb increase for 8-h avg or 12-h avg O₃.

^bPEF results not included in figure.

Associations at Lower Ozone Concentrations

1	In some studies of populations with increased outdoor exposures, O3-associated lung
2	function decrements were observed when maximum or average ambient O ₃
3	concentrations over 30 minutes to 12 hours did not exceed 80 ppb (Chan and Wu, 2005;
4	Korrick et al., 1998; Hoppe et al., 1995; Braun-Fahrlander et al., 1994) (presented within
5	Table 6-2). Korrick et al. (1998) found associations between hike-time average (2-12 h)
6	O_3 concentrations and lung function between concentrations 40 and 74 ppb but not
7	<40 ppb. Several other studies that included higher maximum ambient O ₃ concentrations
8	restricted analyses to observations with 10-min to 1-hour average O3 concentrations
9	<80 ppb (<u>Table 6-6</u>). <u>Higgins et al. (1990</u>) found that O ₃ -associated lung function
10	decrements in children attending camp were limited largely to 1-h avg ambient
11	concentrations >120 ppb; however, many other studies found associations in the lower
12	range of O_3 concentrations (<u>Table 6-6</u>). Among adults exercising outdoors, <u>Spektor et al.</u>

1	(1988b) found that for most lung function parameters, effect estimates in analyses
2	restricted to 30-min max ambient O ₃ concentrations <80 ppb were similar to those
3	obtained for the full range of O_3 concentrations (<u>Table 6-6</u>). In a study of children
4	attending summer camp, similar effects were estimated for the full range of 1-h avg O_3
5	concentrations and those <60 ppb (Spektor et al., 1988a). Brunekreef et al. (1994) found
6	increases in ambient O_3 concentration (10-min to 1-h) during outdoor exercise to be
7	associated with decreases in FEV_1 in analyses restricted to concentrations <61
8	(Table 6-6) and <51 ppb (quantitative results not reported). Whereas Brunekreef et al.
9	(1994) found that effect estimates were near zero with O_3 concentrations <41 ppb
10	(Brunekreef et al., 1994), Brauer et al. (1996) found that associations persisted with
11	1-h max O_3 concentrations <40 ppb (quantitative results not provided).

Table 6-6Associations between ambient ozone concentration and FEV1decrements in different ranges of ambient ozone concentrations.

Study	Location	Population	O ₃ Averaging Time	O₃ Concentration Range	Standardized Percent Change (95% CI) ^a
<u>Brunekreef et al.</u> (1994)	Netherlands	29 adults exercising, ages 18-37 yr	10-min to 1-hour average during exercise	Full range O ₃ <61 ppb	-1.3 (-2.2, -0.37) -2.1 (-4.5, 0.32)
<u>Spektor et al.</u> (1988a)	Fairview Lake, NJ	91 children without asthma at camp, ages 8-15 yr	1-hour average before afternoon FEV ₁ measurement	Full range O ₃ <60 ppb O ₃ <80 ppb	-2.7 (-3.3, -2.0) -2.2 (-3.7, -0.80) -1.4 (-2.5, -0.34)
<u>Spektor et al.</u> (1988b)	Tuxedo, NY	30 adults exercising, ages 21-44 yr	30-min average during exercise	Full range O ₃ <80 ppb	-1.3 (-2.0, -0.64) -1.3 (-2.4, -0.08)
<u>1998</u>)	Mt. Washington, NH	53 adult day hikers, ages 18-64 yr	Hike duration (2-12 h)	Full range O ₃ 40-74 ppb	-1.5 (-2.8, -0.24) -2.6 (-4.9, -0.32)
Higgins et al. (1990)	San Bernardino, CA	43 children without asthma at camp, ages 7-13 yr	1-hour average in the 6-hours before FEV ₁ measurement	>120 ppb <120 ppb	-1.4 (-2.8, 0.03) 0.34 (-1.3, 2.0)

^aResults are presented in order of increasing maximum O_3 concentration included in models. Effect estimates are standardized to a 40-ppb increase for O_3 concentrations averaged over 10 min to 1 h and a 30-ppb increase for O_3 concentrations averaged over 2 to 12 h.

Children with Asthma

12	Increases in ambient O ₃ concentration are associated with lung function decrements in
13	children with asthma in epidemiologic studies conducted across diverse geographical
14	locations and a range of ambient O_3 concentrations (<u>Table 6-7</u>). Whereas most studies of
15	populations with increased outdoor exposures monitored O_3 concentrations at the site of
16	subjects' outdoor activities and used trained staff to measure lung function, studies of

1	children with asthma relied more on O_3 measured at central monitoring sites and lung
2	function measured by subjects. Nonetheless, compared with the camp studies, studies of
3	children with asthma have provided an understanding of the changes in lung function
4	associated with patterns of outdoor activity and ambient O_3 exposure that likely better
5	represent those of children in the general population. Further, these studies have provided
6	more information on factors that potentially may increase the risk of O ₃ -associated
7	respiratory effects and on potential confounding by copollutant exposure or meteorology.

Table 6-7Mean and upper percentile concentrations of ozone in
epidemiologic studies of lung function in children with asthma.

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)	
<u>Jalaludin et al.</u> (2000)	Sydney, Australia	February- December 1994	15-h avg (6 a.m 9 p.m.)	12	Max: 43	
			1-h max	26	91	
<u>Lewis et al.</u> (2005)	Detroit, MI	February 2001- May 2002	24-h avg 8-h max	27.6, 26.5 ^a 40.4, 41.4 ^a	Overall max: 66.3 ^a Overall max: 92.0 ^a	
<u>Just et al. (2002)</u>	Paris, France	April-June 1996	24-h avg	30.0 ^b	Max: 61.7 ^b	
<u>Hoppe et al.</u> (2003)	Munich, Germany	Summers 1992- 1995	30-min max (1-4 p.m.)	High O_3 days: 66.9° Control O_3 days: 32.5°	Max: 91 (high O ₃ days) ^c 39 (control O ₃ days) ^c	
<u>Thurston et al.</u> (1997)	CT River Valley, CT	June 1991-1993	1-h max	83.6 ^c	Max: 160 ^c	
<u>Romieu et al.</u> (2006); (2004b; 2002)	Mexico City, Mexico	October 1998- April 2000	8-h max 1-h max	69 102	Max: 184 Max: 309	
<u>Romieu et al.</u> (1997)	Southern Mexico City, Mexico	April-July 1991; November 1991-February 1992	1-h max	196	Max: 390	
<u>Romieu et al.</u> (1996)	Northern Mexico City, Mexico	April-July 1991; November 1991-February 1992	1-h max	190	Max: 370	
<u>O'Connor et al.</u> (2008)	Boston, MA; Bronx, Manhattan NY; Chicago, IL; Dallas, TX, Seattle, WA; Tucson, AZ (ICAS)	August 1998- July 2001	24-h avg	NR	NR	
<u>Mortimer et al.</u> (2002) (2000)	Bronx, East Harlem, NY; Baltimore, MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St. Louis, MO (NCICAS)	June-August 1993	8-h avg (10 a.m6 p.m.)	48	NR	
<u>Gielen et al.</u> (1997)	Amsterdam, Netherlands	April-July 1995	8-h max	34.2 ^b	Max: 56.5 ^b	

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
Dales et al.	Windsor, ON, Canada	October-	24-h avg	13.0	75th: 26.0
<u>(2009</u>) (<u>2009a</u>)		December 2005	1-h max	27.2	75th: 32.8
Rabinovitch et al. (2004)	Denver, CO	November- March 1999- 2002	1-h max	28.2	75th: 36.0, Max 70.0
Barraza-Villarreal et al. (2008)	Mexico City, Mexico	June 2003-June 2005	8-h moving avg	31.6	Max: 86.3
<u>Wiwatanadate</u> and Trakultivakorn (2010)	Chiang Mai, Thailand	August 2005- June 2006	24-h avg	17.5	90th: 26.82, Max: 34.65
<u>Delfino et al.</u> (2004)	Alpine, CA	September- October 1999; April-June 2000	8-h max	62.9	90th: 83.9, Max: 105.9
Hernández-	Mexico City, Mexico	May-September	24-h avg	26.3	75th: 35.3; Max: 62.8
<u>Cadena et al.</u> (2009)		2005	1-h max	74.5	75th: 92.5; Max: 165

*Note: Studies presented in order of first appearance in the text of this section.

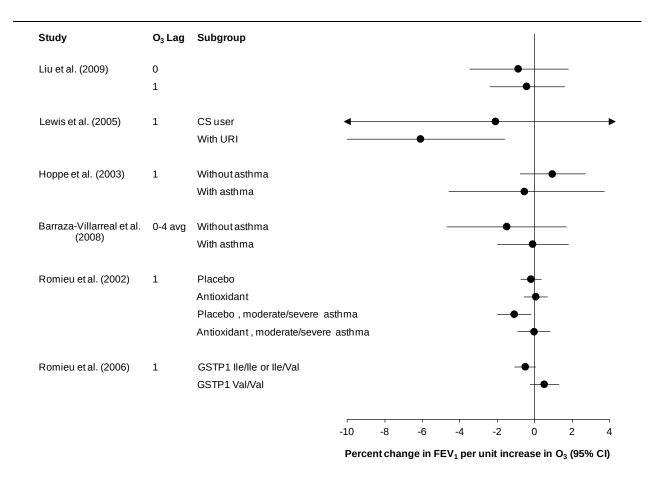
ICAS = Inner City Asthma Study, NR = Not Reported, NCICAS = National Cooperative Inner-City Asthma Study.

^aMeasurements at two sites established by investigators and located within 5 km of most subjects' residences.

 b Concentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

^cMeasured where subjects spent daytime hours.

1	In a majority of studies, including large U.S. multicity studies and several smaller studies
2	conducted in the U.S., Mexico City, and Europe, an increase in ambient O_3 concentration
3	(various averaging times and lags) was associated with a decrement in FEV_1 (Figure 6-6
4	and <u>Table 6-8</u>) or PEF (Figure 6-7 and <u>Table 6-9</u>) in children with asthma. Results were
5	more variable for FEV_1 , which typically was measured on nonconsecutive days, than for
6	PEF, which was measured daily. Further, associations with FEV_1 were found in specific
7	subgroups. Some studies found that increases in ambient O ₃ concentration were
8	associated with greater lung function variability, i.e., a deviation from a baseline level.
9	These results pointed to associations of O ₃ with poorer lung function, as indicated by a
10	decrease from the individual's mean lung function over the study period (Jalaludin et al.,
11	2000), a decrease in lung function over the course of the day (Lewis et al., 2005), or a
12	decrease in the lowest daily measurement (Just et al., 2002). Within many studies,
13	increases in O ₃ concentration were associated concurrently (at the same or similar lag)
14	with decreases in lung function and increases in respiratory symptoms (Just et al., 2002;
15	Mortimer et al., 2002; Gielen et al., 1997; Romieu et al., 1997; Thurston et al., 1997;
16	Romieu et al., 1996) (see Figure 6-11 and Table 6-20 for symptom results).



Note: Results generally are presented in order of increasing mean ambient O_3 concentration. CS = Corticosteroid, URI = Upper respiratory inf ection. Effect estimates are from single-pollutant models and are standardized to a 40-ppb increase for 30-min or 1-h max O_3 concentrations, a 30-ppb increase for 8-h max or 8-h avg O_3 concentrations, and a 20-ppb increase for 24-h avg O_3 concentrations.

Figure 6-6 Percent change in FEV₁ in association with ambient ozone concentrations among children with asthma.

Table 6-8Characteristics and quantitative data for studies represented in
Figure 6-6, of FEV1 or FVC in children with asthma.

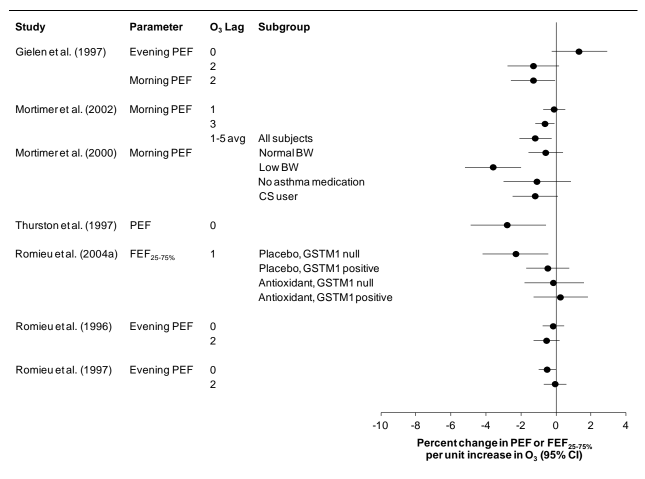
Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Parameter	Subgroup	Standardized Percent Change (95% CI) ^a
<u>Liu et al.</u> (2009a)	Windsor, ON, Canada 182 children with asthma, ages 9-14 yr	24-h avg	0 1	FEV ₁		-0.89 (-3.5, 1.8) -0.44 (-2.4, 1.6)
<u>Lewis et al.</u> (2005)	Detroit, MI 86 children with asthma,	8-h max	1	Lowest daily FEV ₁	CS user With URI	-2.1 (-11.4, 8.3) -6.1 (-10.4, -1.6)
	mean (SD) age 9.1 (1.4) yr		2	-	CS user With URI	-8.0 (-13.5, -2.1) -5.4 (-11.3, 1.0)
<u>Hoppe et</u> <u>al. (2003</u>)	Munich, Germany 43 children, ages 12-23 yr	30-min max (1-4 p.m.)	1	Afternoon FEV ₁	Without asthma With asthma	0.93 (-0.80, 2.7) -0.56 (-4.6, 3.7)
				Afternoon FVC	Without asthma With asthma	-0.09 (-1.7, 1.6) -3.5 (-5.9, -1.0)
<u>Barraza-</u> <u>Villarreal et</u> al. (2008)	Mexico City, Mexico 208 children, ages 6-14 yr	8-h avg	0-4 avg	FEV ₁	50 without asthma 158 with asthma	-1.5 (-4.7, 1.7) -0.12 (-2.0, 1.8)
Romieu et	Mexico City, Mexico	1-h max	1	FEV_1	Placebo	-0.21 (-0.77, 0.36)
<u>al. (2002</u>)	158 children with asthma, ages 6-17 yr				Antioxidant supplement	0.05 (-0.60, 0.69)
					Placebo, moderate/severe asthma	-1.1 (-2.0, -0.19)
					Antioxidant supplement, moderate/severe asthma	-0.04 (-0.92, 0.83)
<u>Romieu et</u> <u>al. (2006</u>)	Mexico City, Mexico 151 children with asthma, mean age 9 yr	1-h max	1	FEV ₁	GSTP1 lle/lle or lle/Val GSTP1 Val/Val	-0.51 (-1.1, 0.05) 0.50 (-0.25, 1.3)
Studies no	t included in Figure 6-6 ^b					
<u>Dales et al.</u> (2009)	Windsor, ON, Canada 182 children with asthma, ages 9-14 yr	1-h max	0	Evening percent predicted FEV ₁		-0.47 (-1.9, 0.95)
Rabinovitch et al. (2004)	Denver, CO 86 children with asthma, ages 6-12 yr	1-h max	0-2 avg	Morning FEV ₁ (mL)		55 (-2.4, 108)
<u>O'Connor</u> <u>et al. (2008</u>)	Boston, MA; Bronx, Manhattan NY; Chicago, IL; Dallas, TX, Seattle, WA; Tucson, AZ	24-h avg	1-5 avg	Change in percent predicted FEV ₁		-0.41 (-1.0, 0.21)
	861 children with asthma, mean (SD) age 7.7 (2.0) yr					

*Includes studies in Figure 6-6, plus others

CS = corticosteroid, URI = Upper respiratory infection.

^aEffect estimates are standardized to a 40-ppb increase for 30-min or 1-h max O_3 , a 30-ppb increase for 8-h max or 8-h avg O_3 , and a 20-ppb increase for 24-h avg O_3 .

^bResults not presented in <u>Figure 6-6</u> because a different form of FEV₁ with a different scale was examined or because sufficient data were not provided to calculate percent change in FEV₁.



Note: Results generally are presented in order of increasing mean ambient O_3 concentration. BW = birth weight, CS = Corticosteroid. Effect estimates are from single pollutant models and are standardized to a 40-ppb increase for 1-h max O_3 concentrations and a 30-ppb increase for 8-h max or 8-h avg O_3 concentrations.

Figure 6-7 Percent change in PEF or FEF_{25-75%} in association with ambient ozone concentrations among children with asthma.

Table 6-9	Characteristics and quantitative data for studies represented in
	Figure 6-7, of PEF or FEF _{25-75%} in children with asthma.

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Parameter	Subgroup	Standardized Percent Change (95% CI) ^a
<u>Gielen et al.</u> (1997)	Amsterdam, Netherlands 61 children with asthma, ages 7- 13 yr	8-h max	0 2 2	Evening PEF Evening PEF Morning PEF		-1.3 (-0.25, 2.9) -1.3 (-2.8, 0.16) -1.3 (-2.6, -0.08)
Mortimer et al. (2002)	Bronx, East Harlem, NY; Baltimore, MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St. Louis, MO 846 children with asthma, ages 4- 9 yr	8-h avg (10 a.m 6 p.m.)	1 3 1-5 avg	Morning PEF	All subjects	-0.12 (-0.76, 0.52) -0.64 (-1.2, -0.10) -1.2 (-2.1, -0.26)
Mortimer et al. (2000)	Bronx, East Harlem, NY; Baltimore, MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St. Louis, MO 846 children with asthma, ages 4- 9 yr	8-h avg (10 a.m6 p.m.)	1-5 avg	Morning PEF	Normal BW Low BW (<5.5 lbs.) No medication CS user	-0.60 (-1.6, 0.39) -3.6 (-5.2, -2.0) -1.1 (-3.0, 0.84) -1.2 (-2.5, 0.11)
<u>Thurston et al.</u> (1997)	CT River Valley, CT 166 children with asthma, ages 7- 13 yr	1-h max	0	Intraday change PEF		-2.8 (-4.9, -0.59)
Romieu et al. (2004b)	Mexico City, Mexico 158 children with asthma, mean age 9 yr	1-h max	1	FEF _{25-75%}	Placebo, GSTM1 null Placebo, GSTM1 positive Antioxidant, GSTM1 null Antioxidant, GSTM1 positive	-2.3 (-4.2, -0.44) -0.48 (-1.7, 0.74) -0.16 (-1.8, 1.6) 0.24 (-1.3, 1.8)
<u>Romieu et al.</u> (1996)	Northern Mexico City, Mexico 71 children with asthma, ages 5- 7 yr	1-h max	0 2	Evening PEF		-0.17 (-0.79, 0.46) -0.55 (-1.3, 0.19)
<u>Romieu et al.</u> (1997)	Southern Mexico City, Mexico 65 children with asthma, ages 5- 13 yr	1-h max	0 2	Evening PEF		-0.52 (-1.0, -0.01) -0.06 (-0.70, 0.58)

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Parameter	Subgroup	Standardized Percent Change (95% CI) ^a
Studies not in	ncluded in Figure 6-7 ^b					
<u>Jalaludin et al.</u> (2000)	Sydney, Australia 20 children with asthma and AHR, mean (SD) age 9.6 (0.9) yr	24-h avg 1-h max	0	Daily deviation from mean PEF		-2.4 (-5.1, 0.28) ^c -1.3 (-2.8, 0.17) ^c
Wiwatanadate and Trakultivakorn (2010)	Chiang Mai, Thailand 31 children with asthma, ages 4- 11 yr	24-h avg	0 5	Daily avg PEF (L/min)		1.0 (-1.6, 3.6) -2.6 (-5.2, 0)
<u>O'Connor et al.</u> (2008)	Boston, MA; Bronx, Manhattan NY; Chicago, IL; Dallas, TX, Seattle, WA; Tucson, AZ 861 Children with asthma, mean (SD) age 7.7 (2.0) yr	24-h avg	1-5 avg	Change in percent predicted PEF		-0.22 (-0.86, 0.43)
<u>Just et al.</u> (2002)	Paris, France 82 children with asthma, mean (SD) age 10.9 (2.5) yr	8-h avg	0-2 avg	Percent variability PEF		15.6 (0, 31.2)

*Includes studies in Figure 6-7, plus others

BW = birth weight, CS = corticosteroid, AHR = Airway hyperresponsiveness.

^aEffect estimates are standardized to a 40-ppb increase for 1-h max O_3 , a 30-ppb increase for 8-h max or avg O_3 , and a 20-ppb increase for 24-h avg O_3 .

^bResults are not presented in <u>Figure 6-7</u> because a different form of PEF with a different scale was examined or because sufficient data were not provided to calculate percent change in PEF.

 $^{\circ}$ Outcome defined as the normalized percent deviation from individual mean PEF during the study period. Quantitative results from generalized estimating equations were provided only for models that included PM₁₀ and NO₂.

1	The most geographically representative data were provided by the large, multi-U.S. city
2	National Cooperative Inner City Asthma Study (NCICAS) (Mortimer et al., 2002; 2000)
3	and Inner-City Asthma Study (ICAS) (O'Connor et al., 2008). Although the two studies
4	differed in the cities, seasons, racial distribution of subjects, and lung function indices
5	examined, results were fairly similar. In ICAS, which included children with asthma and
6	atopy (i.e., allergic sensitization) and year-round examinations of lung function, a 20-ppb
7	increase in the lag 1-5 average of 24-h avg O_3 was associated with a 0.41-point decrease
8	in percent predicted FEV ₁ (95% CI: -1.0, 0.21) and a 0.22-point decrease in percent
9	predicted PEF (95% CI: -0.86, 0.43) (<u>O'Connor et al., 2008</u>).
10	Increases in lag 1-5 avg O_3 (8-h avg, 10 a.m6 p.m.) also were associated with declines
11	in PEF in NCICAS, which included different U.S. cities, summer-only measurements,
12	larger proportions of Black and Hispanic children, and fewer subjects with atopy (79%)
13	(Mortimer et al., 2002). Ozone concentrations lagged 3 to 5 days were associated with
14	larger PEF decrements than were O_3 concentrations lagged 1 to 2 days (Figure 6-7 and
15	Table 6-9). NCICAS additionally identified groups potentially at increased risk of
16	O3-associated PEF decrements, namely, males, children of Hispanic ethnicity, children
17	living in crowded housing, and as indicated in Figure 6-7 and Table 6-9, children with

1	birth weight <5.5 lbs (Mortimer et al., 2000). Somewhat paradoxically, O ₃ was associated
2	with a larger decrease in PEF among subjects taking cromolyn, medication typically used
3	to treat asthma due to allergy, but a smaller decrease among subjects with positive atopy
4	(as determined by skin prick test). NCICAS also indicated robust associations with
5	consideration of other sources of heterogeneity. Except for Baltimore, MD, effect
6	estimates were similar across the study cities (1.1 to 1.7% decrease in PEF per 30-ppb
7	increase in lag 1-5 avg of 8-h avg O_3). Results were similar with O_3 averaged from all
8	available city monitors and concentrations averaged from the three monitors closest to
9	subject ZIP code centroid (1.2% and 1.0%, respectively, per 30-ppb increase in O ₃). At
10	concentrations <80 ppb, a 30-ppb increase in lag 1-5 of 8-h avg O ₃ was associated with a
11	1.4% decrease (95% CI: -2.6, -0.21) in PEF, which was similar to the effect estimated for
12	the full range of O_3 concentrations (Figure 6-7 and Table 6-9). In a study of children with
13	asthma in the Netherlands, Gielen et al. (1997) estimated similar effects for the full range
14	of 8-h max O_3 concentrations and concentrations <51 ppb.
15	Several but not all controlled human exposure studies have reported slightly larger
16	O_3 -induced FEV ₁ decrements in adults with asthma (Section <u>6.2.1.1</u>). However, in the

 O_3 -induced FEV₁ decrements in adults with asthma (Section 6.2.1.1). However, in the 17 few epidemiologic studies that compared children with and without asthma, evidence did 18 not conclusively indicate that children with asthma were at increased risk of 19 O₃-associated lung function decrements. Hoppe et al. (2003) and Jalaludin et al. (2000) 20 generally found larger O₃-associated lung function decrements in children with asthma; 21 whereas Raizenne et al. (1989) did not consistently demonstrate differences between 22 campers with and without asthma. In their study of children in Mexico City, Barraza-23 Villarreal et al. (2008) estimated larger O₃-associated decreases in children without 24 asthma; however, 72% of these children had atopy. These findings indicate that children 25 with atopy, who also have airway inflammation and similar respiratory symptoms, may 26 experience respiratory effects from short-term ambient O₃ exposure.

27 As shown in Figure 6-6 and Figure 6-7 and Table 6-8 and Table 6-9, lung function 28 decrements in children with asthma mostly ranged from <1% to 2% per unit increase in 29 ambient O_3 concentration¹. Larger magnitudes of decrease, were found in children with 30 asthma who were using CS, had a concurrent upper respiratory infection (URI), were 31 GSTM1 null, had airway hyperresponsiveness, or had increased outdoor exposure 32 (Romieu et al., 2006; Lewis et al., 2005; Romieu et al., 2004b; Jalaludin et al., 2000) than 33 among children with asthma overall (Barraza-Villarreal et al., 2008; Lewis et al., 2005; 34 Delfino et al., 2004; Romieu et al., 2002). For example, Jalaludin et al. (2000) estimated a 35 -5.2% deviation from mean FEV₁ per a 20-ppb increase in 24-h avg O₃ concentration 36 among children with asthma and airway hyperresponsiveness and a much smaller -0.71%

¹Effect estimates were standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 8-h max or avg, and 24-h avg O₃.

1	deviation among children with asthma without airway hyperresponsiveness. In a group of
2	86 children with asthma in Detroit, MI, Lewis et al. (2005) reported that associations
3	between ambient O_3 concentration and FEV_1 were confined largely to children with
4	asthma who used CS or had a concurrent URI, 8.0% and 5.4% decreases, respectively, in
5	the mean of lowest daily FEV_1 per 30-ppb increase in 8-h max ambient O_3 concentration.
6	Heterogeneity in response to O ₃ exposure also was demonstrated by observations that
7	some individuals experienced larger O3-associated lung function decrements than the
8	population mean effect estimate. Similar observations were made in controlled human
9	exposure studies (Section 6.2.1.1). Mortimer et al. (2002) found that for a 30-ppb
10	increase in lag 1-5 avg of 8-h avg O ₃ , there was a 30% (95% CI: 4, 61) higher incidence
11	of >10% decline in PEF. Likewise, <u>Hoppe et al. (2003</u>) found that while the percentages
12	of lung function decrements were variable and small, 47% of children with asthma
13	experienced >10% decline in FEV ₁ , FVC, or PEF or 20% increase in airway resistance on
14	days with 30-min (1-4 p.m.) max ambient O ₃ concentrations >50 ppb relative to days
15	with <40 ppb O ₃ .

Effect modification by corticosteroid use

16 In controlled human exposure studies, CS treatment of subjects with asthma generally has 17 not prevented O_3 -induced FEV₁ decrements (Section <u>6.2.1.1</u>). Epidemiologic evidence is 18 equivocal, with findings that use of inhaled CS attenuated (Hernández-Cadena et al., 19 2009), increased (Lewis et al., 2005), and did not affect (Mortimer et al., 2000), ambient 20 O₃-associated lung function decrements. In winter-only studies, consideration of CS use 21 largely did not influence associations between ambient O_3 and various lung function 22 indices (Liu et al., 2009a; Rabinovitch et al., 2004). Similarly equivocal evidence was 23 found for modification of associations with respiratory symptoms (Section 6.2.4.1). The 24 assessment of effect modification by CS use has been hampered by differences in the 25 severity of asthma among CS users and the definition of CS use. Additionally, 26 investigators did not assess adherence to reported CS regimen, and misclassification of 27 CS use may bias findings. For example, Mortimer et al. (2000) classified children by no 28 or any CS use at baseline but did not measure daily use during the study period. Lewis et 29 al. (2005) defined CS use as use for at least 50% of study days and estimated larger 30 O_3 -associated FEV₁ decrements among CS users (Figure 6-6 and Table 6-8) than among 31 CS nonusers (quantitative results not reported). In this study, most children with 32 moderate to severe asthma (91%) were classified as CS users. However, CS users had a 33 higher percent predicted FEV₁. In contrast, <u>Hernández-Cadena et al. (2009</u>) observed 34 larger O_3 -related decrements in FEV₁ among the 60 CS nonusers than among the 25 CS 35 users. A definition for CS use was not provided; however, children with persistent asthma were included among the group of CS nonusers. Thus, across studies, both CS use and nonuse have been used to indicate more severe, uncontrolled asthma.

Effect modification by antioxidant capacity

1

2

3 Ozone is a powerful oxidant whose secondary oxidation products have been described to 4 initiate the key modes of action that mediate decreases in lung function, including the 5 activation of neural reflexes (Section 5.3.2). Additionally, O_3 exposure of humans and 6 animals has induced changes in the levels of antioxidants in the ELF (Section 5.3.3). 7 These observations provide biological plausibility for diminished antioxidant capacity to 8 increase the risk of O₃-associated respiratory effects and for augmented antioxidant 9 capacity to decrease risk. Controlled human exposure studies have demonstrated the 10 protective effects of α -tocopherol (vitamin E) and ascorbate (vitamin C) supplementation 11 on O_3 -induced lung function decrements (Section 6.2.1.1), and epidemiologic studies of 12 children with asthma conducted in Mexico City produced similar findings. Particularly 13 among children with moderate to severe asthma, increases in ambient O_3 concentration 14 were associated with a smaller decrease in FEV_1 in the group supplemented with vitamin 15 C and E as compared with the placebo group (Romieu et al., 2002) (Figure 6-6 and 16 Table 6-8). Romieu et al. (2009) also demonstrated an interaction between dietary 17 antioxidant intake and ambient O₃ concentrations by finding that the main effect of diet 18 was modified by ambient O_3 concentrations. Diets high in antioxidant vitamins and/or 19 omega-3 fatty acids protected against FEV_1 decrements at 8-h max O₃ concentrations 20 \geq 38 ppb. Results for the main effect of O₃ on FEV₁ or effect modification by diet were 21 not presented. 22 Antioxidant capacity also can be characterized by variants in genes encoding xenobiotic 23 metabolizing enzymes with altered enzymatic activity. Ambient O₃-associated FEF_{25-75%} 24 decrements were larger among children with asthma with the GSTM1 null genotype, 25 which is associated with lack of oxidant metabolizing activity (Romieu et al., 2004b). 26 The difference in association between GSTM1 null and positive subjects was minimal in 27 children supplemented with antioxidant vitamins (Figure 6-7 and Table 6-9). Although 28 these findings are biologically plausible given the well-characterized evidence for the 29 secondary oxidation products of O_3 mediating effects, it is important to note that a larger 30 body of controlled human exposure studies has not consistently found larger O₃-induced 31 lung function decrements in GSTM1 null subjects (Section 6.2.1.1). Effect modification 32 by GSTP1 variants is less clear. Romieu et al. (2006) observed larger O₃-associated 33 decreases in FEV₁ in children with asthma with the GSTP1 Ile/Ile or Ile/Val variant, 34 which are associated with relatively higher oxidative metabolism activity (Figure 6-6 and

<u>Table 6-8</u>). An increase in ambient O_3 concentration was associated with an increase in FEV₁ among children with the GSTP1 Val/Val variant, which is associated with reduced

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1oxidative metabolism. Rather than reflecting effect modification by the GSTP1 variant,2these results may reflect effect modification by asthma severity, as 77% of subjects with3the GSTP1 Ile/Ile genotype had moderate to severe asthma. In support of this alternate4hypothesis, another analysis of the same cohort indicated a larger O_3 -associated5decrement in FEV1 among children with moderate to severe asthma than among all6subjects with asthma (Romieu et al., 2002).

Exposure Measurement Error

- 7Across the studies of children with asthma, lung function decrements were associated8with ambient O_3 concentrations assigned to subjects using various exposure assessment9methods. As described in Section 4.3.3, exposure measurement error due to use of10ambient concentrations measured at central sites has varied, depending on the population11and season examined. Because there are a limited number of studies of each method, it is12difficult to conclude that a particular method of exposure assessment produced stronger13results.
- 14 Seasonal differences have been observed in the personal-ambient O₃ relationship 15 (Section 4.3.3); however, in children with asthma, O₃-associated lung function decrements were found in studies conducted in summer months and over multiple 16 17 seasons. Lung function was associated with O₃ measured on site of subjects' daytime 18 hours in summer months (Hoppe et al., 2003; Thurston et al., 1997), factors that have 19 contributed to higher personal-ambient O₃ ratios and correlations. Many year-round 20 studies in Mexico City (Romieu et al., 2006; 2004b; 2002; 1997; 1996) and a study in 21 Detroit, MI (Lewis et al., 2005) found associations with O_3 measured at sites within 5 km 22 of children's home or school. Children with asthma examined by Romieu et al. (2006); 23 (2004b; 2002) had a personal-ambient ratio and correlation for 48- to 72-h avg O₃ 24 concentrations were 0.17 and 0.35, respectively (Ramírez-Aguilar et al., 2008). These 25 findings indicate that the effects of personal O₃ exposure on lung function decrements 26 may have been underestimated in the children in Mexico City. Associations were found 27 with O₃ concentrations averaged across multiple community monitoring sites (O'Connor 28 et al., 2008; Just et al., 2002; Mortimer et al., 2002; Jalaludin et al., 2000) and measured 29 at a single site (Gielen et al., 1997), O_3 measured at multiple sites within a region have 30 shown high temporal correlation (Darrow et al., 2011a; Gent et al., 2003).
- 31Studies of children with asthma restricted to winter months provided little evidence of an32association between various single- and multi-day lags of ambient O3 concentration and33lung function decrements with several studies reporting O3-associated increases in lung34function (Dales et al., 2009; Liu et al., 2009a; Rabinovitch et al., 2004). One explanation35for these results may be lower indoor than outdoor O3 concentrations, variable indoor to

1outdoor ratios, and lower correlations between personal and ambient O3 concentrations in2non-summer months (Section 4.3.2 and Section 4.3.3). As noted for other respiratory3endpoints such as respiratory hospital admissions, ED visits, and mortality, associations4with O3 generally are lower in colder seasons.

Adults with Respiratory Disease

5	Relative to studies in children with asthma, studies of adults with asthma or COPD have
6	been limited in number. Details from these studies regarding location, time period, and
7	ambient O_3 concentrations are presented in <u>Table 6-10</u> . Increases in ambient O_3
8	concentration were not consistently associated with lung function decrements in adults
9	with respiratory disease. Several different exposure assessment methods were used,
10	including monitoring personal exposures (Delfino et al., 1997), monitoring on site of
11	outdoor activity (Girardot et al., 2006; Korrick et al., 1998), and using measurements
12	from one (Peacock et al., 2011; Wiwatanadate and Liwsrisakun, 2011; Thaller et al.,
13	2008; Ross et al., 2002) to several central monitors (Khatri et al., 2009; Lagorio et al.,
14	2006; Park et al., 2005a). There was not a clear indication that differences in exposure
15	assessment methodology contributed to inconsistencies in findings.

Table 6-10 Mean and upper percentile concentrations of ozone in epidemiologic studies of lung function in adults with respiratory disease.

Study*	Location	Study Period	O ₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Delfino et al.</u> (<u>1997</u>)	Alpine, CA	May-July 1994	12-h avg personal (8 a.m8 p.m.)	18	90th: 38 Max: 80
<u>Girardot et al.</u> (2006)	Great Smoky Mountain NP, TN	August-October 2002 June-August 2003	Hike-time avg (2-9 h)	48.1	Max: 74.2
<u>Korrick et al.</u> (1998)	Mt. Washington, NH	Summer 1991, 1992	Hike-time avg (2-12 h)	40	Max: 74
Peacock et al. (2011)	London, England	All-year 1995-1997	8-h max	15.5	Autumn/Winter Max: 32 Spring/Summer Max: 74
Wiwatanadate and Liwsrisakun (2011)	Chiang Mai, Thailand	August 2005-June 2006	24-h avg	17.5	90th: 26.82 Max: 34.65
<u>Thaller et al.</u> (2008); <u>Brooks</u> (2010)	Galveston, TX	Summer 2002-2004	1-h max	35 (median)	Max: 118
Ross et al. (2002)	East Moline, IL	April-October 1994	8-h avg	41.5	Max: 78.3
Khatri et al. (2009)	Atlanta, GA	May-September 2003, 2005, 2006	8-h max	With asthma: 61(median) ^ª No asthma: 56 (median) ^ª	75th (all subjects): 74 ^ª
Lagorio et al. (2006)	Rome, Italy	May-June, November- December 1999	24-h avg	Spring: 36.2 ^b Winter: 8.0 ^b	Overall max: 48.6 ^b
Park et al. (2005a)	Incheon, Korea	March-June 2002	24-h avg	Dust event days: 23.6 Control days: 25.1	NR

* Note: Studies presented in order of first appearance in the text of this section.

NR = Not reported.

^aIndividual-level estimates were calculated based on time spent in the vicinity of various O_3 monitors. ^bConcentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

1	Comparisons of adults with asthma (8-18% of study population) and without asthma did
2	not conclusively demonstrate that adults with asthma had larger ambient O3-associated
3	lung function decrements. Several studies examined on-site or central-site ambient O3
4	concentrations measured while subjects were outdoors, and ambient O3 measured during
5	time spent outdoors has been closer in magnitude and more correlated with personal
6	exposures (Section <u>4.3.3</u>). In a panel study of lifeguards (ages 16-27 years) in Galveston,
7	TX, a larger O_3 -associated decrement in FEV ₁ /FVC was found among the 16 lifeguards
8	with asthma (-1.6% [95% CI: -2.8, -0.4] per 40 ppb increase in 1-h max O ₃) than among
9	the 126 lifeguards without asthma (-0.40% [95% CI: -0.80, 0] per 40-ppb increase in

1	1-h max O_3) Brooks (2010). In Korrick et al. (1998), hikers with a history of asthma or
2	wheeze had larger O_3 -associated lung function decrements (e.g4.4% [95% CI: -7.5,
3	-1.2] in FEV ₁ per 30-ppb increase in 2-12 h avg O_3). In contrast, <u>Girardot et al. (2006</u>)
4	generally did not find O_3 -associated lung function decrements in hikers with or without
5	respiratory disease history. In a cross-sectional study of 38 adults with asthma and
6	13 adults without asthma, <u>Khatri et al. (2009</u>) used central site O_3 measurements but
7	aimed to account for spatial variability by calculating an average of concentrations
8	measured at sites closest to each subject's location during each hour. Investigators
8 9	reported a larger O_3 -associated decrease in percent predicted FEV ₁ /FVC in the 38
10	
	subjects with atopy (with or without asthma) (-12 points [95% CI: -21, -3] per 30-ppb
11	increase in 8-h max O_3) than in subjects with asthma (-4.7 points [95% CI: -11, 2.3]).
12	Among adults with asthma, O_3 was associated with an increase in FEV ₁ .
13	In panel studies that exclusively examined adults with asthma, increases in ambient O_3
14	concentrations, across the multiple lags examined, generally were associated with
15	increases in lung function (Wiwatanadate and Liwsrisakun, 2011; Lagorio et al., 2006;
16	Park et al., 2005a). These studies were conducted in Europe and Asia during periods of
17	low ambient O ₃ concentrations, including one conducted in Korea during a period of dust
18	storms (<u>Park et al., 2005a</u>)
10	
19	Some studies included children and adults with asthma. Among subjects ages 9-46 years
20	(41% adults) in Alpine, CA with low personal 12-h avg O_3 exposures (55% samples
21	below limit of detection) and a majority of sampling hours spent indoors (mean 71%),
22	<u>Delfino et al. (1997</u>) reported that neither 12-h avg personal exposure nor ambient O_3
23	concentration was associated with a decrease in PEF. Ross et al. (2002) examined
24	subjects ages 5-49 years (proportion of adults not reported) in East Moline, IL and found
25	that a 20-ppb increase in lag 0 of 24-h avg O_3 was associated with a 2.6 L/min decrease
26	(95% CI: -4.3, -0.90) in evening PEF. In this population with asthma, an increase in lag 0
27	O_3 also was associated with an increase in symptom score.
28	Controlled human exposure studies have found diminished, statistically nonsignificant
29	O_3 -induced lung function responses in older adults with COPD (Section <u>6.2.1.1</u>).
30	Similarly, epidemiologic studies do not provide strong evidence that short-term increases
31	in ambient O_3 exposure result in lung function decrements in adults with COPD.
32	Inconsistent associations were reported for PEF, FEV ₁ , and FVC in a study that followed
33	94 adults with COPD (ages 40-83 years) in London, England daily over two years
34	(<u>Peacock et al., 2011</u>). For example, a 30-ppb increase in 8-h max O_3 was associated with
35	a 1.7 L/min decrease (95% CI: -3.1, -0.39) in PEF in an analysis of summer 1996 but not
36	summer 1997 (-0.21 L/min [95% CI: -2.4, 2.0]). Further, in this study, an increase in
37	ambient O_3 concentration was associated with lower odds of a large PEF decrement (OR
- 1	anoten of concentration was associated with lower ordes of a hirger Dr decientent (OK

1	for a >20% drop from an individual's median value: 0.89 [95% CI: 0.72, 1.10] per
2	30-ppb increase in lag 1 of 8-h max O ₃) and was not consistently associated with
3	increases in respiratory symptoms (Peacock et al., 2011). Inconsistent associations also
4	were reported in a small panel study of 11 adults with COPD (mean age 67 years) in
5	Rome, Italy (Lagorio et al., 2006).

Populations Not Restricted to Individuals with Asthma

6	Several studies have examined associations between ambient O3 concentrations and lung
7	function in groups that included children with and without asthma; however, a limited
8	number of studies have examined groups of children or adults restricted to healthy
9	individuals. Details from studies not restricted to individuals with asthma regarding
10	location, time period, and ambient O_3 concentrations are presented in <u>Table 6-11</u> .

Table 6-11Mean and upper percentile concentrations of ozone in
epidemiologic studies of lung function in populations not restricted
to individuals with asthma.

Study*	Location	Study Period	O ₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
Avol et al. (1998b)	6 southern CA communities	Spring and summer 1994	24-h avg personal	NR	NR
Hoppe et al. (2003)	Munich, Germany	Summers 1992 ⁻ 1995	30-min max (1-4 p.m.)	High O_3 days: 70.4 ^a Control O_3 days: 29.8 ^a	Max (high O ₃ days): 99 ^a Max (control O ₃ days): 39 ^a
<u>Chen et al. (1999)</u>	3 Taiwan communities	May-January, 1995-1996	1-h max (8 a.m6 p.m.)	NR	Max: 110.3ª
Gold et al. (1999)	Mexico City, Mexico	January- November 1991	24-h avg	52.0 ^a	Max: 103ª
Ward et al. (2002)	Birmingham and Sandwell, England	January-March and May-July 1997	24-h avg	Winter median: 13.0 Summer median: 22.0	Winter Max: 33 Summer Max: 41
<u>Ulmer et al. (1997</u>)	Freudenstadt and Villingen, Germany	March-October 1994	30-min avg	Freudenstadt median: 50.6 Villingen median: 32.1	Freudenstadt 95th: 89.8 Villingen 95th: 70.1
<u>Linn et al. (1996</u>)	Rubidoux, Upland, Torrence, CA	September-June 1992-1994	24-h avg personal 24-h avg ambient	5 23	Max: 16 Max: 53
<u>Scarlett et al.</u> (1996)	Surrey, England	June-July 1994	8-h max	50.7a	Max: 128a
<u>Neuberger et al.</u> (2004)	Vienna, Austria	June-October 1999, January- April 2000	NR	NR	NR
<u>Alexeeff et al.</u> (2008); (2007)	Greater Boston, MA; NAS	January 1995- June 2005	48-h avg	24.4b	NR

Study*	Location	Study Period	O ₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
Steinvil et al. (2009)	Tel Aviv, Israel	September 2002- November 2007	8-h avg (10 a.m6 p.m.)	41.1	75th: 48.7 Max: 72.8
<u>Naeher et al. (1999</u>)	Multiple communities, VA	May-September 1995-1996	8-h max	53.7	Max: 87.6
<u>Son et al. (2010</u>)	Ulsan, Korea	All-year, 2003- 2007	8-h max	35.86 (avg of 13 monitors)	Max: 59.53

* Note: Studies presented in order of first appearance in the text of this section.

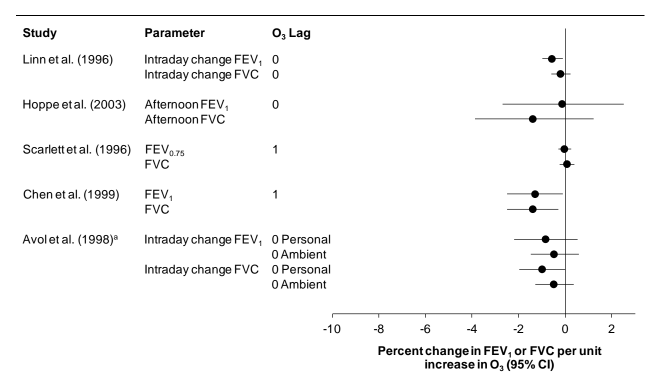
NAS = Normative Aging Study, NR = Not Reported.

^aMeasured at subjects' schools where lung function was measured.

^bMeasured at central monitoring sites established by investigators. Concentrations were averaged across four monitors.

Children

1	Based on studies available at the time of the 2006 O ₃ AQCD, evidence consistently links
2	increases in ambient O_3 concentration with decrements in FEV ₁ and PEF in children
3	(U.S. EPA, 2006b) (Figure 6-8 and Table 6-12). These associations were found with
4	personal O_3 exposures (Avol et al., 1998b), ambient O_3 measured at children's schools
5	where lung function was measured (Hoppe et al., 2003; Chen et al., 1999; Gold et al.,
6	<u>1999</u>), and ambient O_3 measured at sites within the community (Ward et al., 2002; Ulmer
7	et al., 1997; Linn et al., 1996). Among children in California who spent a mean 2-3 hours
8	outdoors per day and whose personal-ambient O_3 correlation was 0.28 across multiple
9	seasons, Avol et al. (1998b) found slightly larger O_3 -associated decrements in FEV ₁ and
10	FVC for 24-h avg personal exposures than for 1-h max ambient measurements
11	(Figure 6-8 and Table 6-12). The effect estimates for personal exposures were similar in
12	magnitude to those found in other studies for ambient O_3 measured at schools (Hoppe et
13	al., 2003; Chen et al., 1999). In another study of children in California, Linn et al. (1996)
14	did not present results for personal O_3 exposures but found FEV_1 decrements in
15	association with increases in ambient O3 concentrations in children who spent 1-2 hours
16	per day outdoors and whose personal-ambient correlations were 0.61. Because of
17	between-study heterogeneity in populations and ambient O3 concentrations examined, it
18	is difficult to assess how the method of exposure assessment may have influenced
19	findings.



Note: The 95% CI was constructed using a standard error that was estimated from the p-value. Results generally are presented in order of increasing mean ambient O_3 concentration. Effect estimates are from single-pollutant models and are standardized to a 40-, 30-, and 20-ppb increase for a 1-hour (or 30-min) max, 8-h max, and 24-h avg O_3 concentrations, respectively.

Figure 6-8 Percent change in FEV₁ or FVC in association with ambient ozone concentrations in studies of children in the general population.

Table 6-12Characteristics and quantitative data for studies represented in
Figure 6-8, of lung function in children.

Study*	Location/ Population	O₃ Averaging Time	O₃ Lag	Parameter	Standardized Percent Change (95% CI) ^a
<u>Linn et al. (1996</u>)	3 southern CA communities 269 children, 4th and 5th grades	24-h avg	0	Intraday change FEV ₁ Intraday change FVC	-0.58 (-1.0, -0.13) -0.21 (-0.62, 0.20)
<u>Hoppe et al.</u> (2003)	Munich, Germany 44 children, ages 6-8 yr	30-min max (1 - 4 p.m.)	0	Afternoon FEV ₁ Afternoon FVC	-0.14 (-2.7, 2.5) -1.4 (-3.9, -1.2)
<u>Scarlett et al.</u> (1996)	Surrey, England 154 children, ages 7-11 yr	8-h max	1	FEV _{0.75} FVC	-0.04 (-0.32, 0.23) 0.07 (-0.25, 0.39)
<u>Chen et al. (1999</u>)	3 Taiwan communities 941 children, mean (SD) age 9.8 (1.6) yr	1-h max	1	FEV ₁ FVC	-1.5 (-2.8, -0.12) -1.6 (-2.9, -0.33)
Avol et al. (1998b)	3 southern CA communities 195 children, ages 10-12 yr	24-h avg personal 1-h max ambient 24-h avg personal 1-h max ambient	0	Intraday change FEV ₁ Intraday change FEV ₁ Intraday change FVC Intraday change FVC	-0.85 (-2.2, 0.53) ^b -0.49 (-1.5, 0.57) ^b -1.0 (-2.0, 0) ^b -0.50 (-1.3, 0.35) ^b
Studies of child	ren not included in Figure 6-8 ^c				
<u>Ulmer et al. (1997</u>)	Freudenstadt and Villingen, Germany 135 children, ages 8-10 yr	30-min max	1	FEV ₁ (mL)	-59 (-103, 14) ^b
Ward et al. (2002)	Birmingham and Sandwell, England 162 children, age 9 yr	24-h avg	0 2	Daily deviation from mean PEF (L/min)	-3.2 (-8.3, 2.0) ^d -6.7 (-12, -1.4) ^d
<u>Gold et al. (1999</u>)	Mexico City, Mexico 40 children, ages 8-11 yr	24-h avg	0 1-10 avg	Intraday change PEF (% change)	-0.47 (-1.1, 0.11) -3.4 (-5.4, -1.5)

^{*}Includes studies in Figure 6-8 plus others.

^aEffect estimates are standardized to a 40-, 30-, and 20-ppb increase for 1-h (or 30-min) max, 8-h max, and 24-h avg O_3 , respectively.

^bThe 95% CI was constructed using a standard error that was estimated from the p-value.

^cResults are not presented in Figure 6-8 because sufficient data were not provided to calculate percent change in FEV₁ or PEF was analyzed.

^dEffect estimates are from analyses restricted to summer months.

1	In the limited number of studies that examined only healthy children, increases in
2	ambient O_3 concentration were associated with decreases (Hoppe et al., 2003) or no
3	change in lung function (Neuberger et al., 2004). Several studies that included small
4	proportions (4-10%) of children with history of respiratory disease or symptoms found
5	associations between increases in ambient O_3 concentration and lung function decrements
6	(Chen et al., 1999; Ulmer et al., 1997; Scarlett et al., 1996). Based on analysis of
7	interaction terms for O_3 concentration and asthma/wheeze history, <u>Avol et al. (1998b</u>)

1	and Ward et al. (2002) did not find differences in O_3 -associated lung function decrements
2	between children with history of asthma or wheeze and healthy children. Combined,
3	these lines of evidence indicate that the ambient O3-associated lung function decrements
4	in children were not solely due to effects in children with asthma, and that increases in
5	ambient O ₃ exposure may decrease lung function in healthy children.
6	Among the studies of children, the magnitudes of decrease in lung function per unit
7	increase in ambient O_3 concentration ¹ ranged from <1 to 4%, a range similar to that
8	estimated in children with asthma. Comparable data were not adequately available to
9	assess whether mean lung function differed between groups of children with asthma and
10	healthy children. In contrast with children with asthma, O3-associated decreases in lung
11	function were not consistently accompanied by O3-associated increases in respiratory
12	symptoms in children in the general population. For example, Gold et al. (1999) found
13	O ₃ -associated decreases in PEF and increases in phlegm; however, the increase in phlegm
14	was associated with lag 1 O3 concentrations whereas the PEF decrement was found with
15	single-day lags 2 to 4 of O ₃ . Also, O ₃ was weakly associated with cough and shortness of
16	breath among children in England (Ward et al., 2002) and was associated with a decrease
17	in respiratory symptom score among children in California (Linn et al., 1996).

Adults

18	Compared with children, in a more limited body of studies, O ₃ was less consistently
19	associated with lung function decrements in populations of adults not restricted to healthy
20	subjects (Table 6-13). In studies that included only healthy adults, increases in ambient
21	O_3 concentration were associated with decreases (Naeher et al., 1999) and increases in
22	lung function (Steinvil et al., 2009). Contrasting results also were found in studies of
23	older adults (Alexeeff et al., 2008; Alexeeff et al., 2007; Hoppe et al., 2003).

¹Effect estimates were standardized to a 40-, 30-, and 20-ppb increase for 1-h max (or 30-min max), 8-h max, and 24-h avg O₃.

Study ^a	Location/Population	O₃ Averaging Time	O₃ Lag	Parameter	O ₃ Assessment Method/Subgroup	Standardized Effect Estimate (95% CI) ^b
<u>Son et al.</u> (2010)	Ulsan, Korea 2,102 children and adults, ages 7-97 yr	8-h max	0-2 avg	Change in percent predicted FEV ₁	All monitor avg Nearest monitor IDW Kriging	-1.4 (-2.7, -0.08) -0.76 (-1.8, 0.25) -1.1 (-2.2, 0.05) -1.4 (-2.6, -0.11)
<u>Steinvil et</u> al. (2009)	Tel Aviv, Israel 2,380 healthy adults, mean age 43 yr, 75th percentile: 52 yr	8-h avg (10 a.m - 6 p.m.)	0 0-6 avg	FEV ₁ (mL)		60 (0, 120) 141 (33, 234)
<u>Naeher et</u> <u>al. (1999</u>)	Multiple communities, VA 473 healthy women, ages 19 - 43 yr	24-h avg	0 0-2 avg	Evening PEF (L/min)		-1.7 (-3.4, 0.03) -3.0 (-4.4, -1.7)
<u>Hoppe et</u> <u>al. (2003</u>)	Munich, Germany 61 older adults, ages 69-95 yr	30-min max (1-4 p.m.)	0 1	% change in afternoon FEV ₁		0.75 (-2.1, 3.7) 1.2 (-1.3, 3.6)
Alexeeff et al. (2008)	Greater Boston, MA 1,015 older adults, mean (SD) age: 68.8 (7.2) yr at baseline	24-h avg	0-1 avg	% change in FEV ₁	GSTP1 Ile/Ile GSTP1 Ile/Val or Val/Val	-1.0 (-2.2, 0.20) -2.3 (-3.5, -1.0)
Alexeeff et al. (2007)	Greater Boston, MA 904 older adults, mean (SD) age: 68.8 (7.3) yr at baseline	24-h avg	0-1 avg	% change in FEV ₁	BMI <30 BMI ≥ 30 No AHR AHR BMI ≥ 30 and AHR	-1.5 (-2.5, -0.51) -3.5 (-5.1, -1.9) -1.7 (-2.7, -0.73) -4.0 (-6.2, -1.8) -5.3 (-8.2, -2.3)

Table 6-13 Associations between ambient ozone concentration and lung function in studies of adults.

^aResults generally are presented in order of increasing mean ambient O₃ concentration.

IDW = Inverse distance weighting, BMI = Body mass index, AHR = airway hyperresponsiveness. ^bEffect estimates are standardized to a 40-ppb increase for 30-min max O_3 , 30-ppb increase for 8-h max or 8-h avg O_3 , and 20-ppb increase for 24-h avg O₃.

 increases in ambient O₃ concentrations assigned to subjects using various methods with potentially varying degrees of measurement error. These methods included the average of multiple intra-city monitors, nearest monitor, estimates from spatial interpolation (Son et al., 2010), average of monitors in multiple towns (Alexceff et al., 2008; 2007), and one site for multiple towns (Nacher et al., 1999). In a large cross-sectional study, conducted in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of association with lung function among ambient O₃ concentrations averaged across 13 city monitors, concentrations from the nearest monitor, inverse distance-weighted 	1	Despite mixed results overall, lung function decrements in adults were associated with
 multiple intra-city monitors, nearest monitor, estimates from spatial interpolation (Son et al., 2010), average of monitors in multiple towns (Alexceff et al., 2008; 2007), and one site for multiple towns (Naeher et al., 1999). In a large cross-sectional study, conducted in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of association with lung function among ambient O₃ concentrations averaged across 13 city 	2	increases in ambient O_3 concentrations assigned to subjects using various methods with
5al., 2010), average of monitors in multiple towns (Alexeeff et al., 2008; 2007), and one6site for multiple towns (Naeher et al., 1999). In a large cross-sectional study, conducted7in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in8Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of9association with lung function among ambient O3 concentrations averaged across 13 city	3	potentially varying degrees of measurement error. These methods included the average of
6site for multiple towns (Nacher et al., 1999). In a large cross-sectional study, conducted7in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in8Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of9association with lung function among ambient O3 concentrations averaged across 13 city	4	multiple intra-city monitors, nearest monitor, estimates from spatial interpolation (Son et
 in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in Ulsan, Korea, <u>Son et al. (2010</u>) did not find a consistent difference in the magnitude of association with lung function among ambient O₃ concentrations averaged across 13 city 	5	al., 2010), average of monitors in multiple towns (Alexeeff et al., 2008; 2007), and one
8 Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of 9 association with lung function among ambient O_3 concentrations averaged across 13 city	6	site for multiple towns (Naeher et al., 1999). In a large cross-sectional study, conducted
9 association with lung function among ambient O_3 concentrations averaged across 13 city	7	in 2,102 children and adults (mean age: 45 years) living near a petrochemical plant in
	8	Ulsan, Korea, Son et al. (2010) did not find a consistent difference in the magnitude of
10 monitors, concentrations from the nearest monitor, inverse distance-weighted	9	association with lung function among ambient O_3 concentrations averaged across 13 city
	10	monitors, concentrations from the nearest monitor, inverse distance-weighted

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4 the study population mean percent predicted FEV_1 was 82.85%, indicating a large	ed
	ed,
5 Description of a big of a still and a desire discuss a between the Destriction Destriction of the discussion of the di	
5 proportion of subjects with underlying airway obstruction. Results from this study were	;
6 not adjusted for meteorological factors and thus, confounding cannot be ruled out.	
7 Importantly, the similarities among exposure assessment methods in <u>Son et al. (2010</u>)	
8 may apply mostly to populations living within the same region of a city. The majority of	of
9 women examined by <u>Naeher et al. (1999</u>) lived >60 miles from the single available	
10 central site monitor. However, in the nonurban (southwest Virginia) study area, O ₃	
11 concentrations may be more spatially homogeneous (Section $4.6.2.1$), and the	
12 concentrations measured at the single site may capture temporal variability in ambient	
13 exposures.	

14 The inconsistent findings for older adults parallel observations from controlled human 15 exposure studies (Section 6.2.1.1). In a study that followed adults ages 69-95 years over a 16 summer in Germany, Hoppe et al. (2003) did not find decreases in lung function in 17 association with ambient O₃ measured at subjects' retirement home. However, recently, 18 the Normative Aging Study found decrements in FEV_1 and FVC in a group of older men 19 (mean [SD] age = 68.9 [7.2] years) in association with ambient O₃ concentrations 20 averaged from four town-specific monitors (Alexeeff et al., 2008), which may less well 21 represent spatial heterogeneity in ambient O_3 exposures. Among all subjects, who were 22 examined once every three years for ten years, associations were found with several lags 23 of 24-h avg O₃ concentration, i.e., 1- to 7-day avg (<u>Alexeeff et al., 2008</u>). Additionally, 24 larger effects were estimated in adults with airway hyperresponsiveness, higher BMI (\geq 25 30), and GSTP1 Ile/Val or Val/Val genetic variants (Val/Val variant produces enzyme 26 with reduced oxidative metabolism activity) (Alexeeff et al., 2008; Alexeeff et al., 2007) 27 (Table 6-13). Larger O_3 -related decrements in FEV₁ and FVC also were observed in 28 subjects with long GT dinucleotide repeats in the promoter region of the gene for the 29 antioxidant enzyme heme oxygenase-1 (Alexeeff et al., 2008), which has been associated 30 with reduced inducibility (Hiltermann et al., 1998). In this cohort, O₃ also was associated 31 with decreases in lung function in adults without airway hyperresponsiveness and those 32 with BMI < 30, indicating effects of O₃ on lung function in healthy older adults. However, 33 the findings may be generalizable only to this study population of older, predominately 34 white men.

Confounding in epidemiologic studies of lung function

35The 1996 O3 AQCD noted uncertainty regarding confounding by temperature and pollen36(U.S. EPA, 1996a); however, collective evidence does not indicate that these factors fully

1	account for the associations observed between increases in ambient O ₃ concentration and
2	lung function decrements. Across the populations examined, most studies that found
3	ambient O3-associated lung function decrements, whether conducted in multiple seasons
4	or only in summer, included temperature in statistical analyses. Some summer camp
5	studies conducted detailed analysis of temperature. In most of these studies, temperature
6	and O_3 were measured at the camps. In two Northeast U.S. studies, an increase in
7	temperature was associated with an increase in lung function (Thurston et al., 1997;
8	Berry et al., 1991). This positive association likely accounted for the nearly 2-fold greater
9	decrease in O_3 -associated PEF found by Thurston et al. (1997) with temperature in the
10	model than with O ₃ alone. In another Northeast U.S. camp study, <u>Spektor et al. (1988a</u>)
11	estimated similar effects for O_3 in a model with and without a temperature-humidity
12	index. In the re analysis of six camp studies, investigators did not include temperature in
13	models because temperature within the normal ambient range had not been shown to
14	affect O3-induced lung function responses in controlled human exposure studies (Kinney
15	<u>et al., 1996</u>).
16	Pollen was evaluated in far fewer studies. Camp studies that examined pollen found that
17	pollen independently was not associated with lung function decrements (Thurston et al.,
18	1997; Avol et al., 1990). Many studies of children with asthma with follow-up over
19	multiple seasons found O3-associated decrements in lung function in models that adjusted
20	for pollen counts (Just et al., 2002; Ross et al., 2002; Jalaludin et al., 2000; Gielen et al.,
21	1997). In these studies, large proportions of subjects had atopy (22-98%), with some
22	studies examining large proportions of subjects specifically with pollen allergy and thus
23	would be more responsive to pollen exposure (Ross et al., 2002; Gielen et al., 1997).

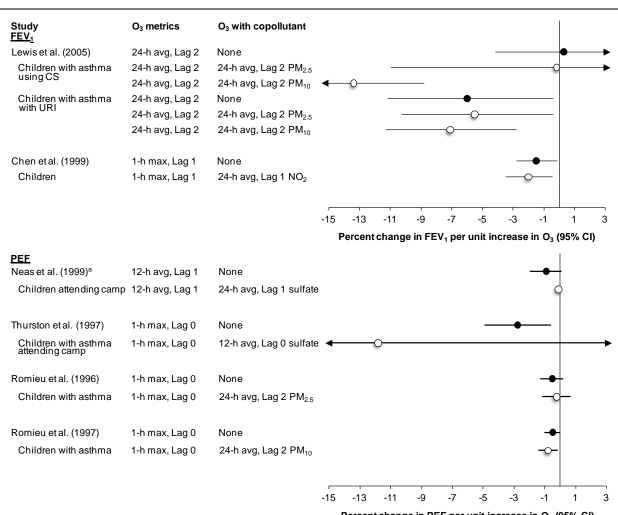
24 A relatively larger number of studies provided information on potential confounding by 25 copollutants such as PM_{2.5}, PM₁₀, NO₂, or SO₂. While studies were varied in how they 26 evaluated confounding, most indicated that O₃-associated lung function decrements were 27 not solely due to copollutant confounding. Some studies of subjects exercising outdoors 28 indicated that ambient concentrations of copollutants such as NO₂, SO₂, or acid aerosol 29 were low and thus, not likely to confound associations observed for O₃ (Hoppe et al., 30 2003; Brunekreef et al., 1994; Hoek et al., 1993). In other studies of children with 31 increased outdoor exposures, O₃ was consistently associated with decreases in lung 32 function, whereas other pollutants such as PM2.5, sulfate, and acid aerosol individually 33 showed variable associations across studies (Thurston et al., 1997; Castillejos et al., 1995; 34 Berry et al., 1991; Avol et al., 1990; Spektor et al., 1988a). Most of these studies 35 measured ambient pollutants on site of subjects' outdoor activity and related lung 36 function changes to the pollutant concentrations measured during outdoor activity. Thus, 37 the degree of exposure measurement error likely is comparable for O_3 and copollutants.

1	Studies that conducted copollutant modeling generally found O ₃ -associated lung function
2	decrements to be robust; most copollutant-adjusted effect estimates fell within the
3	95% CI of the single-pollutant effect estimates (Figure 6-9 and Table 6-14). These studies
4	used central site measurements for both O ₃ and copollutants. There may be residual
5	confounding because of differential exposure measurement error for O ₃ and copollutants
6	due to differing spatial heterogeneity and indoor-outdoor ratios; however, the limited
7	available evidence indicates that personal O ₃ exposures are weakly correlated with
8	personal $PM_{2.5}$ and NO_2 exposures (Section <u>4.3.4.1</u>). Whereas a few studies used the same
9	averaging time for copollutants (Lewis et al., 2005; Jalaludin et al., 2000), more
10	examined 1-h max or 8-h max O ₃ and 24-h avg copollutant concentrations (Son et al.,
11	2010; Chen et al., 1999; Romieu et al., 1997; Romieu et al., 1996). In a Philadelphia-area
12	summer camp study, Neas et al. (1999) was among the few studies to find that the effect
13	estimate for O ₃ was attenuated to near zero in a copollutant model (24-h avg sulfate in
14	this study) (Figure 6-9 and Table 6-14).
15	

15 Ambient O_3 concentrations showed a wide range of correlations with copollutant 16 concentrations (r = -0.31 to 0.74). In Sydney, Australia, Jalaludin et al. (2000) found low 17 correlations of O_3 with PM_{10} (r = 0.13) and NO_2 (r = -0.31), all averaged over 24 hours. 18 In two-pollutant models, PM₁₀ and NO₂ remained associated with increases in PEF, and 19 O₃ remained associated with decreases in PEF in children with asthma. In Detroit, MI, O₃ 20 was moderately correlated with $PM_{2.5}$ (Pearson r = 0.57) and PM_{10} (Pearson r = 0.59), all 21 averaged over 24 hours (Lewis et al., 2005). Adjustment for PM_{10} or $PM_{2.5}$ resulted in a 22 large change in the O_3 -associated FEV₁ decrement in children with asthma, but only in 23 CS users and not in children with concurrent URI (Figure 6-9 and Table 6-14). Studies 24 conducted in Mexico City found small changes in O₃-associated PEF decrements with 25 copollutant adjustment although different averaging times were used for copollutants 26 (Romieu et al., 1997; Romieu et al., 1996) (Figure 6-9 and Table 6-14). In these studies, 27 O_3 was moderately correlated with copollutants such as NO₂ and PM₁₀ (range of Pearson 28 r = 0.38-0.58). Studies conducted in Asia also found that associations between O₃ and 29 lung function were robust to adjustment for weakly- to moderately-correlated 30 copollutants; effect estimates for copollutants generally were attenuated, indicating that 31 O₃ may confound associations of copollutants (Son et al., 2010; Chen et al., 1999). 32 In a summer camp study conducted in Connecticut, Thurston et al. (1997) found ambient

32In a summer camp study conducted in Connecticut, Indiston et al. (1997) found ambient33concentrations of 1-h max O_3 and 12-h avg sulfate to be highly correlated (r = 0.74),34making it difficult to separate their independent effects. With sulfate in the model, a35larger decrease in PEF was estimated for O_3 ; however, the 95% CI was much wider36(Figure 6-9 and Table 6-14). Investigators found that the association for sulfate was due37to one day when the ambient concentrations of both pollutants were at their peak. With38the removal of this peak day, the sulfate effect was attenuated, whereas O_3 effects

1remained robust (Thurston et al., 1997). Among children with asthma in Thailand, the2O3-associated decrease in PEF was robust to adjustment of SO2; however, different lags3were examined for O3 (lag 5) and SO2 (lag 4) (Wiwatanadate and Trakultivakorn, 2010).



Percent change in PEF per unit increase in O₃ (95% CI)

Note: Results are presented first for FEV₁ then for PEF and then in order of increasing mean ambient O_3 concentration. ^aInformation was not available to calculate 95% CI of the copollutant model. CS = corticosteroid, URI = Upper respiratory infection. Effect estimates are standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 12-h avg, and 24-h avg O_3 , respectively. Black circles represent O_3 effect estimates from single pollutant models, and open circles represent O_3 effect estimates from copollutant models.

Figure 6-9 Comparison of ozone-associated changes in lung function in single- and co-pollutant models.

Study*	Location/Population	Parameter	O ₃ -associated Percent Change in Single-Pollutant Model (95% Cl) ^a	O₃-associated Percent Change in Copollutant Model (95% Cl) ^a
PEF				
<u>Neas et al.</u> (1999)	Philadelphia, PA 156 Children at summer camp, ages 6 - 11 yr	Morning PEF	For 12-h avg, Lag 1 -0.94 (-2.0, 0.08)	With 24-h avg, Lag 1 sulfate -0.10 ^b
<u>Thurston et al.</u> (1997)	CT River Valley 166 Children with asthma at summer camp, ages 7-13 yr	Intraday change PEF	For 1-h max, Lag 0 -2.8 (-4.9, -0.59)	With 12-h avg, Lag 0 sulfate -11.8 (-31.6, 8.1)
<u>Romieu et al.</u> (1996)	Mexico City, Mexico 71 children with asthma, ages 5-7 yr	Evening PEF	For 1-h max, Lag 2 -0.55 (-1.3, 0.19)	With 24-h avg, Lag 2 PM _{2.5} -0.24 (-1.2, 0.68)
<u>Romieu et al.</u> (1997)	Mexico City, Mexico 65 children with asthma, ages 5-13 yr	Evening PEF	For 1-h max, Lag 0 -0.52 (-1.0, -0.01)	With 24-h avg, Lag 0 PM ₁₀ -0.79 (-1.4, -0.16)
FEV ₁				
<u>Lewis et al.</u> (2005)	Detroit, MI Children with asthma using CS 393 person-days	Lowest daily FEV ₁	For 24-h avg, Lag 2 0.29 (-4.2, 5.0)	With 24-h avg, Lag 2 PM _{2.5} -0.18 (-11.0, 11.9) With 24-h avg, Lag 2 PM ₁₀ -13.4 (-17.8, -8.8)
	Children with asthma with URI 231 person-days Overall mean (SD) age 9.1 (1.4 yr)	_	For 24-h avg, Lag 2 -6.0 (-11.2, -0.41)	With 24-h avg, Lag 2 PM _{2.5} -5.5 (-10.3, -0.42) With 24-h avg, Lag 2 PM ₁₀ -7.1 (-11.3, -2.8)
<u>Chen et al.</u> (1999)	3 Taiwan communities 941 children, mean (SD) age 9.8 (1.6) yr	FEV ₁	For 1-h max, Lag 1 -1.5 (-2.8, -0.12)	With 24-h avg, Lag 1 NO ₂ -2.0 (-3.5, -0.43)

Table 6-14Additional characteristics and quantitative data for studies
represented in Figure 6-9.

Study*	Location/Population	Parameter	O ₃ -associated Percent Change in Single-Pollutant Model (95% Cl) ^a	O ₃ -associated Percent Change in Copollutant Model (95% Cl) ^a
Results not in	cluded in Figure 6-9 ^c			
<u>Jalaludin et al.</u> (2000)	Sydney, Australia 125 children with asthma or wheeze, mean (SD) age 9.6 (1.0) yr	Daily deviation from mean PEF	For 24-h avg, Lag 0 -1.8 (-3.5, -0.19)	With 24-h avg, Lag 0 PM ₁₀ , -1.8 (-3.5, -0.19) With 24-h avg, Lag 0 NO ₂ -1.8 (-3.4, -0.11)
<u>Wiwatanadate</u> and Trakultivakorn (2010)	Chiang Mai, Thailand 31 children with asthma, ages 4-11 yr	Evening PEF (L/min)	For 24-h avg, Lag 5 -2.6 (-5.2, 0)	With 24-h avg, Lag 4 SO ₂ -3.2 (-6.2, -0.2)
<u>Son et al.</u> (2010)	Ulsan, Korea 2,102 children and adults, ages 7-97 yr	Change in percent predicted FEV ₁	For 8-h max, Lag 0-2 avg (kriging) -1.4 (-2.6, -0.11)	With 24-h avg, Lag 2 PM ₁₀ (kriging) -1.8 (-3.4, -0.25)

*Includes studies in Figure 6-9 plus others.

CS = Corticosteroid, URI = Upper respiratory infection.

^aEffect estimates are standardized to a 40-ppb increase for 1-h max O_3 , 30-ppb increase for 8-h max or 12-h avg O_3 , and 20-ppb increase for 24-h avg O_3 .

^bInformation was not available to calculate 95% CI.

^cResults are not presented in Figure 6-9 because sufficient data were not provided to calculate percent change in lung function.

1Some studies did not provide quantitative results but reported that O3-associated lung2function decrements remained statistically significant in models that included3copollutants such as PM10, NO2, sulfate, nitrate, or ammonium (Romieu et al., 1998b;4Brauer et al., 1996; Linn et al., 1996; Spektor et al., 1988b).

Several studies estimated robust O₃-associated lung function decrements in multipollutant
models that most often included O₃, NO₂, and either PM_{2.5} or PM₁₀ (O'Connor et al.,
2008; Thaller et al., 2008; Chan and Wu, 2005; Romieu et al., 2002; Korrick et al., 1998;
Higgins et al., 1990). However, the independent effects of O₃ are more difficult to assess
in relation to incremental changes in more than one copollutant.

Summary of Epidemiologic Studies of Lung Function

10The cumulative body of epidemiologic evidence indicates that short-term increases in11ambient O3 concentration are associated with decrements in lung function in children12with asthma (Figure 6-6 and Figure 6-7 and Table 6-8 and Table 6-9) and without13asthma. In contrast with results from controlled human exposure studies, within-study14epidemiologic comparisons did not consistently indicate larger ambient O3-associated15lung function decrements in groups with asthma (children or adults) than in groups16without asthma. Notably, most epidemiologic studies were not designed to assess

1	between-group differences. Based on comparisons between studies, differences were
2	noted between children with and without asthma in so far as in studies of children with
3	asthma, an increase in ambient O ₃ concentration was associated concurrently with lung
4	function decrements and increases in respiratory symptoms (Just et al., 2002; Mortimer et
5	al., 2002; Ross et al., 2002; Gielen et al., 1997; Romieu et al., 1997; Thurston et al.,
6	<u>1997; Romieu et al., 1996</u>). In studies of children in the general population, O ₃ -associated
7	decreases in lung function were not accompanied by O3-associated increases in
8	respiratory symptoms (Ward et al., 2002; Gold et al., 1999; Linn et al., 1996).
9	Across studies of children, there was no clear indication that a particular exposure
10	assessment method using central site measurements produced stronger findings, despite
11	potential differences in exposure measurement error. In children, lung function was
12	associated with ambient O ₃ concentrations measured on site of children's daytime hours
13	(Hoppe et al., 2003; Thurston et al., 1997), at children's schools (Chen et al., 1999; Gold
14	et al., 1999), at the closest site (Romieu et al., 2006; Lewis et al., 2005; Romieu et al.,
15	2004b; 2002; 1997; 1996), at multiple community sites then averaged (O'Connor et al.,
16	2008; Just et al., 2002; Mortimer et al., 2002; Jalaludin et al., 2000), and at a single site
17	(Ward et al., 2002; Gielen et al., 1997; Ulmer et al., 1997; Linn et al., 1996). Among
18	children in California, \ found slightly larger O3-associated lung function decrements for

20 As noted in the 1996 and 2006 O₃ AQCDs, evidence clearly demonstrates ambient 21 O₃-associated lung function decrements in children and adults engaged in outdoor 22 recreation, exercise, or work. Moreover, several results indicated associations with 10-23 min to 12-h avg O₃ concentrations <80 ppb. These studies are noteworthy for their 24 measurement of ambient O_3 on site of and at the time of subjects' outdoor activity, 25 factors that have contributed to higher O₃ personal exposure-ambient concentration 26 correlations and ratios (Section 4.3.3). These epidemiologic results are well supported by 27 observations from controlled human exposure studies of lung function decrements 28 induced by O_3 exposure during exercise (Section 6.2.1.1). Although investigation was 29 relatively limited, increases in ambient O₃ concentration were not consistently associated 30 with lung function decrements in adults with respiratory disease, healthy adults, or older 31 adults.

24-h avg personal exposures than for 1-h max ambient concentrations.

32Across the diverse populations examined, most effect estimates ranged from a <1 to 2%</th>33decrease in lung function per unit increase in O_3 concentration¹. Heterogeneity in34 O_3 -associated respiratory effects within populations was indicated by observations of35larger decreases (3-8%) in children with asthma with CS use or concurrent URI (Lewis et36al., 2005) and older adults with airway hyperresponsiveness and/or BMI >30 (Alexeeff et

19

¹ Effect estimates were standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 8-h max, and 24-h avg O₃.

1	al 2007) Among shildren in Marias City, high distant antioxident inteles attenueted
1 2	al., 2007). Among children in Mexico City, high dietary antioxidant intake attenuated
2 3	O ₃ -associated lung function decrements (<u>Romieu et al., 2004b</u> ; <u>2002</u>), similar to results
	from controlled human exposure studies. Each of these potential effect modifiers was
4	examined in one to two populations; thus, firm conclusions about their influences are not
5	warranted. Adding to the evidence for heterogeneity in response, <u>Hoppe et al. (2003</u>) and
6	<u>Mortimer et al. (2002)</u> found that increases in ambient O_3 concentration were associated
7	with increased incidence of $>10\%$ decline in lung function in children with asthma.
8	Collectively, epidemiologic studies examined and found lung function decrements in
9	association with single-day O ₃ concentrations lagged from 0 to 7 days and concentrations
10	averaged over 2-10 days. More studies found associations with O3 concentrations lagged
11	0 or 1 day (Son et al., 2010; Alexeeff et al., 2008; Lewis et al., 2005; Ross et al., 2002;
12	Jalaludin et al., 2000; Chen et al., 1999; Romieu et al., 1997; Brauer et al., 1996; Romieu
13	et al., 1996; Spektor et al., 1988b) than those lagged 5-7 days (Wiwatanadate and
14	Trakultivakorn, 2010; Hernández-Cadena et al., 2009; Steinvil et al., 2009). Associations
15	with multiday average concentrations (Son et al., 2010; Liu et al., 2009a; Barraza-
16	Villarreal et al., 2008; O'Connor et al., 2008; Alexeeff et al., 2007; Mortimer et al., 2002;
17	Ward et al., 2002; Gold et al., 1999; Naeher et al., 1999; Neas et al., 1999) indicate that
18	elevated exposures over several days may be important. Within studies, O_3
19	concentrations for multiple lag periods were associated with lung function decrements,
20	possibly indicating that multiple modes of action may be involved in the responses.
21	Activation of bronchial C-fibers (Section $5.3.2$) may lead to decreases in lung function as
22	an immediate response to O_3 exposure, and increased airway hyperresponsiveness to
23	antigens resulting from sensitization of airways by O_3 (Section 5.3.5) may mediate lung
24	function responses associated with lagged or multiday O_3 exposures (Peden, 2011).
25	
25	For single- and multi-day average O_3 concentrations, lung function decrements were
26	associated with 1-h max, 8-h max, and 24-h avg O ₃ , with no strong difference in the
27	consistency or magnitude of association among the averaging times. For example, among
28	studies that examined multiple averaging times, <u>Spektor and Lippmann (1991</u>) found a
29	larger magnitude of association for 1-h max O ₃ than for 24-h avg O ₃ . However, other
30	studies found larger magnitudes of association for longer averaging times [8-h max in
31	<u>Chan and Wu (2005)</u> and 12-h avg in <u>Thaller et al. (2008</u>)] than for 1-h max O ₃ . Other
32	studies found no clear difference among O_3 averaging times (<u>Jalaludin et al., 2000</u> ; <u>Chen</u>
33	et al., 1999; Scarlett et al., 1996; Berry et al., 1991).
34	Several studies found that associations with lung function decrements persisted at lower
35	ambient O_3 concentrations. For O_3 concentrations averaged up to 1 hour during outdoor
36	recreation or exercise, associations were found in analyses restricted to O ₃ concentrations
37	<80 ppb (Spektor et al., 1988a; Spektor et al., 1988b), 60 ppb (Brunekreef et al., 1994;

1	Spektor et al., 1988a), and 50 ppb (Brunekreef et al., 1994). Among outdoor workers,
2	Brauer et al. (1996) found a robust association using daily 1-h max O ₃ concentrations
3	<40 ppb. Ulmer et al. (1997) found a robust association in schoolchildren using 30-min
4	max O_3 concentrations <60 ppb. For 8-hour avg O_3 concentrations, associations with lung
5	function decrements in children with asthma were found to persist at concentrations
6	<80 ppb in a U.S. multicity study (for lag 1-5 avg) (Mortimer et al., 2002) and <51 ppb in
7	a study conducted in the Netherlands (for lag 2) (Gielen et al., 1997).

8 Evidence did not demonstrate strong confounding by meteorological factors and 9 copollutant exposures. Most O3 effect estimates for lung function were robust to 10 adjustment for temperature, humidity, and copollutants such as PM_{2.5}, PM₁₀, NO₂, or SO₂. 11 Although examined in few epidemiologic studies, O_3 was associated with decreases in 12 lung function with adjustment for pollen or acid aerosols. The consistency of association 13 in the collective body of evidence with and without adjustment for ambient copollutant 14 concentrations and meteorological factors combined with evidence from controlled 15 human exposure studies for the direct effects of O3 exposure provide strong support for 16 the independent effects of short-term ambient O₃ exposure on lung function decrements.

6.2.1.3 Toxicology: Lung Function

17	The 2006 O ₃ AQCD concluded that pulmonary function decrements occur in a number of
18	species with acute exposures (≤ 1 week), ranging from 0.25 to 0.4 ppm O ₃ (U.S. EPA,
19	<u>2006b</u>). Early work has demonstrated that during acute exposure of ~ 0.2 ppm O ₃ in rats,
20	the most commonly observed alterations are increased frequency of breathing and
21	decreased tidal volume (i.e., rapid, shallow breathing). Decreased lung volumes are
22	observed in rats with acute exposures to 0.5 ppm O ₃ . At concentrations of \geq 1 ppm,
23	breathing mechanics (compliance and resistance) are also affected. Exposures of 6 h/day
24	for 5 days create a pattern of attenuation of pulmonary function decrements in both rats
25	and humans without concurrent attenuation of lung injury and morphological changes,
26	indicating that the attenuation did not result in protection against all the effects of O_3
27	(Tepper et al., 1989). A number of studies examining the effects of O_3 on pulmonary
28	function in rats, mice, and dogs are described in Table 6-13 on page 6-91 (U.S. EPA,
29	<u>1996m</u>) of the 1996 O ₃ AQCD, and Table AX5-11 on page AX5-34 (U.S. EPA, 2006f) of
30	the 2006 O ₃ AQCD (U.S. EPA, 2006b, 1996a). Lung imaging studies using
31	hyperpolarized ³ He provide evidence of ventilation abnormalities in rats following
32	exposure to 0.5 ppm O_3 (Crémillieux et al., 2008). Rats were exposed to 0.5 ppm O_3 for 2
33	or 6 days, either continuously (22 h/day) or alternatingly (12 h/day). Dynamic imaging of
34	lung filling (2 mL/sec) revealed delayed and incomplete filling of lung segments and
35	lobes. Abnormalities were mainly found in the upper regions of the lungs and proposed to

be due to the spatial distribution of O₃ exposure within the lung. Although the small
 number of animals used in the study (n = 3 to 7/group) makes definitive conclusions
 difficult, the authors suggest that the delayed filling of lung lobes or segments is likely a
 result of an increase in airway resistance brought about by narrowing of the peripheral
 small airways.

6.2.2 Airway Hyperresponsiveness

6 Airway hyperresponsiveness refers to a condition in which the conducting airways 7 undergo enhanced bronchoconstriction in response to a variety of stimuli. Airway 8 responsiveness is typically quantified by measuring changes in pulmonary function 9 (e.g., FEV_1 or specific airway resistance [sRaw]) following the inhalation of an 10 aerosolized specific (allergen) or nonspecific (e.g., methacholine) bronchoconstricting 11 agent or another stimulus such as exercise or cold air. Asthmatics are generally more 12 sensitive to bronchoconstricting agents than nonasthmatics, and the use of an airway 13 challenge to inhaled bronchoconstricting agents is a diagnostic test in asthma. Standards 14 for airway responsiveness testing have been developed for the clinical laboratory (ATS, 15 2000a), although variation in methodology for administering the bronchoconstricting 16 agent may affect the results (Cockcroft et al., 2005). There is a wide range of airway 17 responsiveness in nonasthmatic people, and responsiveness is influenced by a wide range 18 of factors, including cigarette smoke, pollutant exposures, respiratory infections, 19 occupational exposures, and respiratory irritants. Airways hyperresponsiveness in 20 response to O_3 exposure has not been examined widely in epidemiologic studies; such 21 evidence is derived primarily from controlled human exposure and toxicological studies.

6.2.2.1 Controlled Human Exposures

22 Beyond its direct effect on lung function, O₃ exposure causes an increase in airway 23 responsiveness in human subjects. Increased airway responsiveness is an important 24 consequence of exposure to ambient O_3 , because the airways are then predisposed to 25 narrowing upon inhalation of a variety of ambient stimuli. 26 Increases in airway responsiveness have been reported for exposures to 80 ppb O₃ and 27 above. Horstman et al. (1990) evaluated airway responsiveness to methacholine in young 28 healthy adults (22 M) exposed to 80, 100, and 120 ppb O₃ (6.6 hours, quasi continuous 29 moderate exercise, 39 L/min). Dose-dependent decreases of 33, 47, and 55% in the 30 cumulative dose of methacholine required to produce a 100% increase in sRaw after 31 exposure to O₃ at 80, 100, and 120 ppb, respectively, were reported. Molfino et al. (1991)

1	reported increased allergen-specific airway responsiveness in mild asthmatics exposed to
2	120 ppb O_3 (1 hour resting exposure). Due to safety concerns, however, the exposures in
3	the Molfino et al. (1991) study were not randomized with FA conducted first and O_3
4	exposure second. Attempts to reproduce the findings of Molfino et al. (1991) using a
5	randomized exposure design have not found statistically significant changes in airway
6	responsiveness at such low levels of O_3 exposure. At a considerably higher exposure to
7	250 ppb O_3 (3 h, light-to-moderate intermittent exercise, 30 L/min), Jorres et al. (1996)
8	found significant increases in specific and non-specific airway responsiveness of mild
9	asthmatics 3 hours following O_3 exposure. Kehrl et al. (1999) found increased reactivity
10	to house dust mite antigen in mild atopic asthmatics 16-18 hours after exposure to
11	160 ppb O_3 (7.6 hours, light quasi continuous exercise, 25 L/min). Holz et al. (2002)
12	demonstrated that repeated daily exposure to lower concentrations of 125 ppb O_3 (3 hours
13	for four consecutive days; intermittent exercise, 30 L/min) causes an increased response
14	to allergen challenge at 20 hours postexposure in allergic airway disease.
15	Ozone exposure of asthmatic subjects, who characteristically have increased airway
16	responsiveness at baseline relative to healthy controls (by nearly two orders of
17	magnitude), can cause further increases in responsiveness (Kreit et al., 1989). Similar
18	relative changes in airway responsiveness are seen in asthmatics and healthy control
19	subject exposed to O ₃ despite their markedly different baseline airway responsiveness.
20	Several studies (Kehrl et al., 1999; Jorres et al., 1996; Molfino et al., 1991) have
21	suggested an increase in specific (i.e., allergen-induced) airway reactivity. An important
22	aspect of increased airway responsiveness after O ₃ exposure is that this may provide
23	biological plausibility for associations observed between increases in ambient O ₃
24	concentrations and increased respiratory symptoms in children with asthma
25	(Section 6.2.4.1) and increased hospital admissions and ED visits for asthma
26	(Section $6.2.7$).
27	Changes in airway responsiveness after O_3 exposure appear to resolve more slowly than
27 28	
	changes in FEV ₁ or respiratory symptoms (Folinsbee and Hazucha, 2000). Studies
29	suggest that O_3 -induced increases in airway responsiveness usually resolve 18 to 24 hours
30	after exposure, but may persist in some individuals for longer periods (Folinsbee and
31	<u>Hazucha, 1989</u>). Furthermore, in studies of repeated exposure to O_3 , changes in airway
32	responsiveness tend to be somewhat less susceptible to attenuation with consecutive
33	exposures than changes in FEV ₁ (Gong et al., 1997a; Folinsbee et al., 1994; Kulle et al.,
34	<u>1982; Dimeo et al., 1981</u>). Increases in airway responsiveness do not appear to be
35	strongly associated with decrements in lung function or increases in symptoms (Aris et
36	al., 1995). Recently, Que et al. (2011) assessed methacholine responsiveness in healthy
37	young adults (83M, 55 F) one day after exposure to 220 ppb O_3 and FA for 2.25 hours
38	(alternating 15 min periods of rest and brisk treadmill walking). Increases in airways

1	responsiveness at 1 day post- O_3 exposure were not correlated with FEV ₁ responses
2	immediately following the O ₃ exposure nor with changes in epithelial permeability
3	assessed 1 day post-O ₃ exposure.

6.2.2.2 Toxicology: Airway Hyperresponsiveness

4	In addition to human subjects, a number of species, including nonhuman primates, dogs,
5	cats, rabbits, and rodents, have been used to examine the effect of O ₃ exposure on airway
6	hyperresponsiveness (see Table 6-14 on page 6-93 (U.S. EPA, 1996n) of the 1996 O_3
7	AQCD and Table AX5-12 on page AX5-36 (U.S. EPA, 2006g) of the 2006 O ₃ AQCD).
8	With a few exceptions, commonly used animal models have been guinea pigs, rats, or
9	mice acutely exposed to O ₃ concentrations of 1 to 3 ppm to induce airway
10	hyperresponsiveness. These animal models are helpful for determining underlying
11	mechanisms of general airway hyperresponsiveness and are relevant for understanding
12	airway responses in humans. Although 1-3 ppm may seem like a high exposure
13	concentration, based on ¹⁸ O ₃ (oxygen-18-labeled O ₃) in the BALF of humans and rats, an
14	exposure of 0.4 ppm O ₃ in exercising humans appears roughly equivalent to an exposure
15	of 2 ppm in resting rats (<u>Hatch et al., 1994</u>).
16	A limited number of studies have observed airway hyperresponsiveness in rodents and
17	guinea pigs after exposure to less than 0.3 ppm O_3 . As previously reported in the 2006 O_3
18	AQCD, one study demonstrated that a very low concentration of O ₃ (0.05 ppm for 4 h)
19	induced airway hyperresponsiveness in some of the nine strains of rats tested (Depuydt et
20	<u>al., 1999</u>). This effect occurred at a concentration of O_3 that was much lower than has
21	been reported to induce airway hyperresponsiveness in any other species. Similar to the
22	effects of O ₃ on other endpoints, these observations suggest a genetic component plays an
23	important role in O ₃ -induced airway hyperresponsiveness in this species and warrants
24	verification in other species. More recently, Chhabra et al. (2010) demonstrated that
25	exposure of ovalbumin (OVA)-sensitized guinea pigs to 0.12 ppm for 2 h/day for
26	4 weeks produced specific airway hyperresponsiveness to an inhaled OVA challenge.
27	Interestingly, in this study, dietary supplementation of the guinea pigs with vitamins C
28	and E ameliorated a portion of the airway hyperresponsiveness as well as indices of
29	inflammation and oxidative stress. Larsen and colleagues conducted an O3 C-R study in
30	mice sensitized by 10 daily inhalation treatments with an OVA aerosol (Larsen et al.,
31	2010). Although airway responsiveness to methacholine was increased in non-sensitized
32	animals exposed to a single 3-hour exposure to 0.5, but not 0.1 or 0.25 ppm O_3 , airway
33	hyperresponsiveness was observed after exposure to 0.1 and 0.25 ppm O_3 in OVA-
34	sensitized mice.

1	In order to evaluate the ability of O_3 to enhance specific and non-specific airway
2	responsiveness, it is important to take into account the phenomenon of attenuation in the
3	effects of O ₃ . Several studies have clearly demonstrated that some effects caused by acute
4	exposure are absent after repeated or prolonged exposures to O ₃ . The ability of the
5	pulmonary system to adapt to repeated insults to O_3 is complex, however, and
6	experimental findings for attenuation to O3-induced airway hyperresponsiveness are
7	inconsistent. Airway hyperresponsiveness was observed in mice after a 3-hour exposure
8	but not in mice exposed continuously for 72 hours to 0.3 ppm (Johnston et al., 2005b).
9	However, the Chhabra study demonstrated O3-induced airway hyperresponsiveness in
10	guinea pigs exposed for 2 h/day for 10 days (Chhabra et al., 2010). Besides the obvious
11	species disparity, these studies differ in that the mice were exposed continuously for
12	72 hours, whereas the guinea pigs were exposed intermittently over 10 days, suggesting
13	that attenuation might be lost with periods of rest in between O_3 exposures.
14	Overall, numerous toxicological studies have demonstrated that O ₃ -induced airway
15	hyperresponsiveness occurs in guinea pigs, rats, and mice after either acute or repeated
16	exposure to relevant concentrations of O_3 . The mechanisms by which O_3 enhances the
17	airway responsiveness to either specific (e.g., OVA) or non-specific (e.g., methacholine)
18	bronchoprovocation are not clear, but appear to be associated with complex cellular and

biochemical changes in the conducting airways. A number of potential mediators and cells may play a role in O_3 -induced airway hyperresponsiveness; mechanistic studies are discussed in greater detail in Section <u>5.3</u>.

6.2.3 Pulmonary Inflammation, Injury and Oxidative Stress

22	In addition to physiological pulmonary responses, respiratory symptoms, and airway
23	hyperresponsiveness, O ₃ exposure has been shown to result in increased epithelial
24	permeability and respiratory tract inflammation. In general, inflammation can be
25	considered as the host response to injury and the induction of inflammation as evidence
26	that injury has occurred. Inflammation induced by exposure of humans to O_3 can have
27	several potential outcomes: (1) inflammation induced by a single exposure (or several
28	exposures over the course of a summer) can resolve entirely; (2) continued acute
29	inflammation can evolve into a chronic inflammatory state; (3) continued inflammation
30	can alter the structure and function of other pulmonary tissue, leading to diseases such as
31	fibrosis; (4) inflammation can alter the body's host defense response to inhaled
32	microorganisms, particularly in potentially at-risk populations such as the very young and
33	old; and (5) inflammation can alter the lung's response to other agents such as allergens
34	or toxins. Except for outcome (1), the possible chronic responses have only been directly
35	observed in animals exposed to O_3 . It is also possible that the profile of response can be

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altered in persons with preexisting pulmonary disease (e.g., asthma, COPD) or smokers. Oxidative stress has been shown to play a key role in initiating and sustaining O_3 -induced inflammation. Secondary oxidation products formed as a result of reactions between O_3 and components of the ELF can increase the expression of cytokines, chemokines, and adhesion molecules and enhance airway epithelium permeability (Section <u>5.3.3</u>. and Section <u>5.3.4</u>).

6.2.3.1 Controlled Human Exposures

- 7 As reported in studies reviewed in the 1996 and 2006 O₃ AQCDs, acute O₃ exposure 8 initiates an acute inflammatory response throughout the respiratory tract that has been 9 observed to persist for at least 18-24 hours postexposure. A meta-analysis of 21 studies 10 (Mudway and Kelly, 2004a) for varied experimental protocols (80-600 ppb O₃; 11 1-6.6 hours duration; light to heavy exercise; bronchoscopy at 0-24 hours post-O₃ 12 exposure) showed that neutrophils (PMN) influx in healthy subjects was linearly 13 associated (p < 0.01) with total O₃ dose (i.e., the product of O₃ concentration, exposure 14 duration, and \dot{V}_{E}). As with FEV₁ responses to O₃, within individual inflammatory 15 responses to O_3 are generally reproducible and correlated between repeat exposures (Holz 16 et al., 1999). Some individuals also appear to be intrinsically more susceptible to 17 increased inflammatory responses to O_3 exposure (Holz et al., 2005).
- 18 The presence of PMNs in the lung has long been accepted as a hallmark of inflammation 19 and is an important indicator that O₃ causes inflammation in the lungs. Neutrophilic 20 inflammation of tissues indicates activation of the innate immune system and requires a 21 complex series of events that are normally followed by processes that clear the evidence 22 of acute inflammation. Inflammatory effects have been assessed in vivo by lavage 23 (proximal airway and bronchoalveolar), bronchial biopsy, and more recently, induced 24 sputum. A single acute exposure (1-4 hours) of humans to moderate concentrations of O₃ 25 (200-600 ppb) while exercising at moderate to heavy intensities results in a number of 26 cellular and biochemical changes in the lung, including an inflammatory response 27 characterized by increased numbers of PMNs, increased permeability of the epithelial 28 lining of the respiratory tract, cell damage, and production of proinflammatory cytokines 29 and prostaglandins (U.S. EPA, 2006b). These changes also occur in humans exposed to 30 80 and 100 ppb O₃ for 6-8 hours (Alexis et al., 2010; Peden et al., 1997; Devlin et al., 31 1991). Significant (p = 0.002) increases in sputum PMN (16-18 hours postexposure) 32 relative to FA responses have been recently reported for 60 ppb O₃ which is the lowest 33 exposure concentration that has been investigated in young healthy adults (Kim et al., 34 2011). Soluble mediators of inflammation such as the cytokines (e.g., IL-6, IL-8) and 35 arachidonic acid metabolites (e.g., prostaglandin [PG]E₂, PGF_{2a}, thromboxane, and

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1	leukotrienes [LTs] such as LTB ₄) have been measured in the BALF of humans exposed
2	to O ₃ . In addition to their role in inflammation, many of these compounds have
3	bronchoconstrictive properties and may be involved in increased airway responsiveness
4	following O_3 exposure. The possible relationship between repetitive bouts of acute
5	inflammation in humans caused by O_3 and the development of chronic respiratory disease
6	is unknown.

Asthma

7 Inflammatory responses to O_3 exposure have also been studied in asthmatic subjects. 8 Asthmatics exposed to 200 ppb O_3 for 4-6 hours with exercise show significantly more 9 neutrophils in BALF (18 hours postexposure) than similarly exposed healthy individuals 10 (Scannell et al., 1996; Basha et al., 1994). In allergic asthmatics who tested positive for 11 Dermatophagoides farinae antigen, there was an eosinophilic inflammation (2-fold 12 increase), as well as neutrophilic inflammation (3-fold increase) 18 hours after exposure 13 to 160 ppb O₃ for 7.6 hours with exercise (Peden et al., 1997). In a study of subjects with 14 intermittent asthma exposed to 400 ppb O_3 for 2 hours, increases in eosinophil cationic 15 protein, neutrophil elastase and IL-8 were found to be significantly increased 16 hours 16 postexposure and comparable in induced sputum and BALF (Hiltermann et al., 1999). At 17 18 hours post- O_3 exposure (200 ppb, 4 hours with exercise) and corrected for FA 18 responses, Scannell et al. (1996) found significantly increased neutrophils in 18 19 asthmatics (12%) compared to 20 healthy subjects (4.5%). This difference in 20 inflammatory response was observed despite no group differences in spirometric 21 responses to O_3 . Scannell et al. (1996) also reported that IL-8 tends to be higher in the 22 BALF of asthmatics compared to nonasthmatics following O₃ exposure, suggesting a 23 possible mediator for the significantly increased neutrophilic inflammation in those 24 subjects. Bosson et al. (2003) found significantly greater epithelial expression of IL-5, 25 IL-8, granulocyte-macrophage colony-stimulating factor and epithelial cell-derived 26 neutrophil-activating peptide-78 in asthmatics compared to healthy subjects following 27 exposure to 200 ppb O_3 for 2 h. In contrast, Stenfors et al. (2002) did not detect a 28 difference in the O₃-induced increases in neutrophil numbers between 15 mild asthmatic 29 and 15 healthy subjects by bronchial wash at the 6 hours postexposure time point. 30 However, the asthmatics were on average 5 years older than the healthy subjects in this 31 study, and it is not yet known how age affects inflammatory responses. It is also possible 32 that the time course of neutrophil influx differs between healthy and asthmatic 33 individuals. Differences between asthmatics and healthy subjects in O₃-mediated 34 activation of innate and adaptive immune responses have been observed in two studies 35 (Hernandez et al., 2010; Bosson et al., 2003), as discussed in Section 6.2.5.4 and 36 Section 5.4.2.2.

- 1 Vagaggini et al. (2002) investigated the effect of prior allergen challenge on responses in 2 mild asthmatics exposed for 2 hours to 270 ppb O₃ or filtered air. At 6 hours 3 postexposure, eosinophil numbers in induced sputum were found to be significantly 4 greater after O_3 than after air exposures. Studies such as this suggest that the time course 5 of eosinophil and neutrophil influx following O_3 exposure can occur at levels detectable 6 within the airway lumen by as early as 6 h. They also suggest that the previous or 7 concurrent activation of proinflammatory pathways within the airway epithelium may 8 enhance the inflammatory effects of O₃. For example, in an in vitro study of primary 9 human nasal epithelial cells and BEAS-2B cell line, cytokine production induced by 10 rhinovirus infection was enhanced synergistically by concurrent exposure to O₃ at 11 200 ppb for 3 hours (Spannhake et al., 2002).
- 12 A few studies have evaluated the effects of corticosteroid usage on the response of 13 asthmatics to O_3 . Vagaggini et al. (2007) evaluated whether corticosteroid usage would 14 prevent O₃-induced lung function decrements and inflammatory responses in a group of 15 subjects with mild persistent asthma (n = 9; 25 ± 7 years). In this study, asthmatics were 16 randomly exposed on four occasions to 270 ppb O₃ or FA for 2 hours with moderate 17 exercise. Exposures were preceded by four days of treatment with prednisone or placebo. 18 Pretreatment with corticosteroids prevented an inflammatory response in induced sputum 19 at 6 hours postexposure. FEV_1 responses were, however, not prevented by corticosteroid 20 treatment and were roughly equivalent to those observed following placebo. Vagaggini et 21 al. (2001) also found budesonide to decrease airway neutrophil influx in asthmatics 22 following O_3 exposure. In contrast, inhalation of corticosteroid budesonide failed to 23 prevent or attenuate O₃-induced responses in healthy subjects as assessed by 24 measurements of lung function, bronchial reactivity and airway inflammation 25 (Nightingale et al., 2000). High doses of inhaled fluticasone and oral prednisolone have 26 each been reported to reduce inflammatory responses to O_3 in healthy individuals (Holz 27 et al., 2005).
- 28 Stenfors et al. (2010) exposed persistent asthmatics (n = 13; aged 33 years) receiving 29 chronic inhaled corticosteroid therapy to 200 ppb O_3 for 2 hours with moderate exercise. 30 At 18 hours postexposure, there was a significant O₃-induced increase in 31 bronchioalveolar lavage (BAL) neutrophils, but not eosinophils. Bronchial biopsy also 32 showed a significant O₃-induced increase in mast cells. This study suggests that the 33 protective effect of acute corticosteroid therapy against inflammatory responses to O_3 in 34 asthmatics demonstrated by Vagaggini et al. (2007) may be lost with continued treatment 35 regimes.

Associations between Inflammation and $\ensuremath{\mathsf{FEV}}\xspace_1$ responses

1	Studies reviewed in the 2006 O_3 AQCD reported that inflammatory responses do not
1 2	appear to be correlated with lung function responses in either asthmatic or healthy
2 3	subjects. In healthy adults (14 M, 6 F) and asthmatic (12 M, 6 F) volunteers exposed to
4	200 ppb O ₃ (4 hours with moderate quasi continuous exercise, $\dot{V}_E = 44$ L/min), percent
5	PMN and total protein in BAL fluids were significantly increased in the asthmatics
6	relative to the healthy controls. Spirometric measures of lung function were significantly
7	decreased following the O_3 exposure in both groups, but were not significantly different
8	between the asthmatic and healthy subjects. Effects of O_3 on PMN and total protein were
9	not correlated with changes in FEV_1 or FVC (<u>Balmes et al., 1997</u> ; <u>Balmes et al., 1996</u>).
10	<u>Devlin et al. (1991</u>) exposed healthy adults (18 M) to 80 and 100 ppb O_3 (6.6-hours with
11	moderate quasi continuous exercise, 40 L/min). In BAL fluid collected 18 hours after
12	exposure to 100 ppb O ₃ , significant increases in PMNs, protein, PGE2, fibronectin, IL-6,
13	lactate dehydrogenase, and α -1 antitrypsin compared to FA. Similar but smaller increases
14	in all mediators were found after exposure to 80 ppb O ₃ except for protein and
15	fibronectin. Changes in BAL markers were not correlated with changes in FEV_1 . Holz et
16	<u>al. (1999</u>) examined inflammatory responses in healthy ($n = 21$) and asthmatic ($n = 15$)
17	subjects exposed to 125 and 250 ppb O ₃ (3 h, light intermittent exercise, 26 L/min).
18	Significantly increased percent PMN in sputum due to O ₃ exposure was observed in both
19	asthmatics and healthy subjects following the 250 ppb exposure. At the lower 125 ppb
20	exposure, only the asthmatic group experienced statistically significant increases in the
21	percent PMN. Significant decrements in FEV_1 were only found following exposure to
22	250 ppb; these changes in FEV_1 did not differ significantly between the asthmatic and
23	healthy groups and were not correlated with changes in PMN levels. Peden et al. (1997)
24	also found no correlation between PMN and FEV ₁ responses in 8 individuals with asthma
25	exposed to 160 ppb O ₃ for 7.6 hours with light-to-moderate exercise ($\dot{V}_E = 25$ L/min).
26	However, a marginally significant correlation ($r = -0.69$, two-tailed $p = 0.08$, $n = 7$) was
27	observed between increases in the percentage of eosinophils and FEV ₁ responses
28	following O ₃ exposure.
29	In contrast to these earlier findings, <u>Vagaggini et al. (2010</u>) recently reported a significant
30	(r = 0.61, p = 0.015) correlation between changes in FEV ₁ and changes in sputum
31	neutrophils in mild-to-moderate asthmatics (n = 23; 33 ± 11 years) exposed to 300 ppb
32	O ₃ for 2 hours with moderate exercise. Eight subjects were categorized as "responders"
33	based on $>10\%$ FEV ₁ decrements. There were no baseline differences between
34	responders and nonresponders. However, at 6 hours post-O ₃ exposure, sputum
35	neutrophils were significantly increased by 15% relative to FA in responders. The
36	neutrophil increase in responders was also significantly greater than the 0.2% increase in
37	nonresponders. Interestingly, the nonresponders in the <u>Vagaggini et al. (2010</u>) study

experienced a significant O_3 -induced 11.3% increase in sputum eosinophils, while responders had an nonsignificant 2.6% decrease.

Time Course of the Inflammatory Response

3 The time course of the inflammatory response to O_3 in humans has not been fully 4 characterized. Different markers exhibit peak responses at different times. Studies in 5 which lavages were performed 1 hour after O_3 exposure (1 hours at 400 ppb or 4 hours at 6 200 ppb) have demonstrated that the inflammatory responses are quickly initiated (Torres 7 et al., 1997; Devlin et al., 1996; Schelegle et al., 1991). Inflammatory mediators and 8 cytokines such as IL-8, IL-6, and PGE₂ are greater at 1 hours than at 18 hours post- O_3 9 exposure (Torres et al., 1997; Devlin et al., 1996). However, IL-8 still remained elevated 10 at 18 hours post-O₃ exposure (4 hours at 200 ppb O₃ versus FA) in healthy subjects 11 (Balmes et al., 1996). Schelegle et al. (1991) found increased PMNs in the "proximal 12 airway" lavage at 1, 6, and 24 hours after O_3 exposure (4 hours at 200 ppb O_3), with a 13 peak response at 6 hours. However, at 18-24 hours after O₃ exposure, PMNs remain 14 elevated relative to 1 hour postexposure (Torres et al., 1997; Schelegle et al., 1991).

Genetic Polymorphisms

15	Alexis et al. (2010) recently reported that a 6.6-hour exposure with moderate exercise to
16	80 ppb O3 caused increased sputum neutrophil levels at 18 hours postexposure in young
17	healthy adults (n = 15; 24 \pm 1 years). In a prior study, <u>Alexis et al. (2009</u>) found genotype
18	effects on inflammatory responses to O ₃ , but not lung function responses following a
19	2-hour exposure to 400 ppb O ₃ . At 4 hours post-O ₃ exposure, both GSTM1 genotypes
20	had significant increases in sputum neutrophils with a tendency for a greater increase in
21	GSTM1-sufficient than null individuals. At 24 hours postexposure, neutrophils had
22	returned to baseline levels in the GSTM1-sufficient individuals. In the GSTM1-null
23	subjects, however, neutrophil levels increased further from 4 hours to 24 hours and were
24	significantly greater than both baseline levels and 24 hours levels in GSTM1-sufficient
25	individuals. <u>Alexis et al. (2009</u>) found that GSTM1-sufficient individuals ($n = 19$;
26	24 ± 3 years) had a decrease in macrophage levels at 4-24 hours postexposure to 400 ppb
27	O_3 for 2 hours with exercise. These studies also provide evidence for activation of innate
28	immunity and antigen presentation, as discussed in Section 5.3.6. Effects of the exposure
29	apart from O_3 cannot be ruled out in the <u>Alexis et al. (2010</u>); (2009) studies, however,
30	since no FA exposure was conducted.
31	Vagaggini et al. (2010) examined FEV ₁ and sputum neutrophils in mild-to-moderate
32	asthmatics (n = 23; 33 ± 11 years) exposed to 300 ppb O ₃ for 2 hours with moderate
33	exercise. Six of the subjects were NQO1 wild type and GSTM1 null, but this genotype

was not found to be associated with O₃-induced changes in lung function or inflammatory

34

1

2

1	responses to O ₃ . <u>Kim et al. (2011</u>) showed a significant ($p = 0.002$) increase in sputum
2	neutrophil levels following a 6.6-hour exposure to 60 ppb O_3 relative to FA in young
3	healthy adults (13 F, 11 M; 25.0 ± 0.5 years). There was no significant effect of GSTM1
4	genotype (half GSTM1-null) on the inflammatory responses observed in these
5	individuals. Previously, inflammatory responses had only been evaluated down to a level
6	of 80 ppb O ₃ .

Repeated Exposures

7 Markers from BALF following both 2 hours (Devlin et al., 1997) and 4 hours (Jorres et 8 al., 2000; Christian et al., 1998) repeated O_3 exposures (up to 5 days) indicate that there is 9 ongoing cellular damage irrespective of the attenuation of some cellular inflammatory 10 responses of the airways, pulmonary function, and symptom responses. Devlin et al. 11 (1997) found that several indicators of inflammation (e.g., PMN, IL-6, PGE₂, fibronectin) 12 were attenuated after 5 days of exposure (i.e., values were not different from FA). 13 However, other markers (LDH, IL-8, total protein, epithelial cells) did not show 14 attenuation, suggesting that tissue damage probably continues to occur during repeated 15 exposure. Some cellular responses did not return to baseline levels for more than 10-16 20 days following O₃ exposure. Christian et al. (1998) showed decreased numbers of 17 neutrophils and a decrease in IL-6 levels in healthy adults after 4 days of exposure versus 18 the single exposure to 200 ppb O_3 for 4 h. Jorres et al. (2000) also found both functional 19 and BALF cellular responses to O_3 were abolished at 24 hours postexposure following 20 the fourth exposure day. However, levels of total protein, IL-6, IL-8, reduced glutathione 21 and ortho-tyrosine were still increased significantly. In addition, visual scores 22 (bronchoscopy) for bronchitis and erythema and the numbers of neutrophils in bronchial 23 mucosal biopsies were increased. Results indicate that, despite an attenuation of some 24 markers of inflammation in BALF and pulmonary function decrements, inflammation 25 within the airways persists following repeated exposure to O_3 . The continued presence of 26 cellular injury markers indicates a persistent effect that may not necessarily be recognized 27 due to the attenuation of spirometric and symptom responses.

Epithelial Permeability

28	A number of studies show that O ₃ exposures increase epithelial cell permeability through
29	direct (technetium-99m labeled diethylene triamine pentaacetic acid, 99mTc-DTPA,
30	clearance) and indirect (e.g., increased BALF albumin, protein) techniques. Kehrl et al.
31	(1987) showed increased ^{99m} Tc-DTPA clearance in healthy young adults (age 20-30 yrs)
32	at 75 minutes postexposure to 400 ppb O_3 for 2 hours. Also in healthy young adults (age
33	26±2 yrs), Foster and Stetkiewicz (1996) have shown that increased ^{99m} Tc-DTPA
34	clearance persists for at least 18-20 hours post- O_3 exposure (130 minutes to average O_3
35	concentration of 240 ppb), and the effect is greater at the lung apices than at the base. In a

1	older group of healthy adults (mean age = 35 yrs), Morrison et al. (2006) observed
2	99m Tc-DTPA clearance at 1 hours and 6 hours postexposure to O ₃ (100 and 400 ppb; 1
3	hour; moderate intermittent exercise, $\dot{V}_E = 40 \text{ L/min}$ to be similar and not statistically
4	different from ^{99m} Tc-DTPA clearance at 1 hours postexposure to FA (1 h; $\dot{V}_{E} = 40$
5	L/min).
6	Increased BALF protein, suggesting O ₃ -induced changes in epithelial permeability, have
7	also been reported at 1 hour and 18 hours postexposure (Devlin et al., 1997; Balmes et
8	al., 1996). Meta-analysis of results from 21 publications (Mudway and Kelly, 2004a) for
9	varied experimental protocols (80-600 ppb O ₃ ; 1-6.6 hours duration; light to heavy
10	exercise; bronchoscopy at 0-24 hours post-O ₃ exposure), showed that increased BALF
11	protein is associated with total inhaled O_3 dose (i.e., the product of O_3 concentration,
12	exposure duration, and \dot{V}_E).
13	It has been postulated that changes in permeability associated with acute inflammation
14	may provide increased access of inhaled antigens, particles, and other inhaled substances
15	deposited on lung surfaces to the smooth muscle, interstitial cells, and the blood. Hence,
16	increases in epithelial permeability following O3 exposure might lead to increases in
17	airway responsiveness to specific and nonspecific agents. Que et al. (2011) investigated
18	this hypothesis in healthy young adults (83M, 55 F) exposed to 220 ppb O_3 for 2.25 hours
19	(alternating 15 min periods of rest and brisk treadmill walking). As has been observed by
20	others for FEV ₁ responses, within individual changes in permeability were correlated
21	between sequential O3 exposures. This indicates intrinsic differences in susceptibility to
22	epithelial damage from O ₃ exposure among individuals. Increases in epithelial
23	permeability at 1 day post-O ₃ exposure were not correlated with FEV_1 responses
24	immediately following O_3 exposure nor with changes in airway responsiveness to
25	methacholine assessed 1 day post- O_3 exposure. The authors concluded that changes in
26	FEV ₁ , permeability, and airway responsiveness following O ₃ exposure were relatively
27	constant over time in young healthy adults; although, these responses appear to be
28	mediated by differing physiologic pathways.

6.2.3.2 Epidemiology

29In the 2006 O3 AQCD, epidemiologic evidence of associations between short-term30increases in ambient O3 concentration (30-min or 1-h max) and changes in pulmonary31inflammation was limited to a few observations of increases in nasal lavage levels of32inflammatory cell counts, eosinophilic cationic protein, and myeloperoxidases (U.S.33EPA, 2006b). In recent years, there has been a large increase in the number of studies34assessing ambient O3-related changes in pulmonary inflammation and oxidative stress,

- 1types of biological samples collected (i.e., lower airway), and types of indicators2examined. Most studies collected samples every 1 to 3 weeks resulting in a total of 3 to 83samples per subject. These recent studies form a larger base to establish coherence with4findings from controlled human exposure and animal studies that have measured the5same or related biological markers. Additionally, these studies provide further biological6plausibility for the associations observed between ambient O3 concentrations and7respiratory symptoms and asthma exacerbations.
- 8 Despite the strengths of studies of inflammation, it is important to note that research in 9 this field continues to develop, and several uncertainties are recognized that may limit 10 inferences of the effects of ambient O₃ exposure. Current areas of development include 11 examination of the clinical relevance of the observed magnitudes of changes in biological 12 markers of pulmonary inflammation (Murugan et al., 2009; Duramad et al., 2007), 13 characterization of the time course of changes between biomarker levels and other 14 endpoints of respiratory morbidity, development of standardized methodologies for 15 collection, improvement of the sensitivity and specificity of assay methods, and 16 characterization of subject factors (e.g., asthma severity and recent medication use) that 17 contribute to inter-individual variability. These sources of uncertainty may contribute to 18 differences in findings among studies.
- 19 Although most of the biomarkers examined in epidemiologic studies were not specific to 20 the lung, most studies collected exhaled breath, exhaled breath condensate (EBC), nasal 21 lavage fluid, or induced sputum with the aim of monitoring inflammatory responses in 22 airways, as opposed to monitoring systemic responses in blood. The biomarker most 23 frequently measured was exhaled nitric oxide (eNO), likely related to its ease of 24 collection in the field and automated measurement. Other biological markers were 25 examined in EBC, induced sputum, and nasal lavage fluid, which are hypothesized to 26 represent the fluid lining the lower or upper airways and contain cytokines, cells, and/or 27 markers of oxidative stress that mediate inflammatory responses (Balbi et al., 2007; 28 Howarth et al., 2005; Hunt, 2002). Table 6-15 presents the locations, time periods, and 29 ambient O_3 concentrations for studies examining associations with biological markers of 30 pulmonary inflammation and oxidative stress. Many studies found that short-term 31 increases in ambient O_3 concentration were associated with increases in pulmonary 32 inflammation and oxidative stress, in particular, studies of children with asthma 33 conducted in Mexico City (Figure 6-10 and Table 6-16 and
- 34 Table 6-17).

Table 6-15	Mean and upper percentile ozone concentrations in studies of
	biological markers of pulmonary inflammation and oxidative stress.

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Barraza-</u> <u>Villarreal et al.</u> (2008)	Mexico City, Mexico	June 2003- June 2005	8-h moving avg	31.6	Max: 86.3
Berhane et al. (2011)	13 Southern California Communities	September 2004- June 2005	8-h avg (10 a.m6 p.m.)	NR	NR
<u>Liu et al.</u> (2009a)	Windsor, ON, Canada	October- December 2005	24-h avg	13.0	95th: 26.5
<u>Khatri et al.</u> (2009)	Atlanta, GA	May- September 2003, 2005, 2006	8-h max	With asthma: 61 (median) ^a No asthma: 56 (median) ^a	75th (all subjects): 74 ^a
<u>Qian et al.</u> (2009)	Boston, MA; New York, NY; Denver, CO; Philadelphia, PA; San Francisco, CA; Madison, WI (SOCS)	February 1997-January 1999	8-h max	33.6	75th: 44.4, Max: 91.5
<u>Romieu et al.</u> (2008)	Mexico City, Mexico	January- October 2004	8-h max	31.1	75th: 38.3, Max: 60.7
<u>Sienra-Monge</u> et al. (2004)	Mexico City, Mexico	All-year 1999- 2000	8-h max	66.2	Max: 142.5
Ferdinands et al. (2008)	Suburb of Atlanta, GA	August 2004	1-h max	61 (median)	75th: 67
<u>Chimenti et al.</u> (2009)	Sicily, Italy	November, February, July, year NR	8-h avg (7 a.m. – 3 p.m.)	November: 32.7 (pre- race), 35.1 (race) ^b	NR
				February: 37.0 (pre-race), 30.8 (race) ^b	
				July: 51.2 (pre-race), 46.1 (race) ^b	
<u>Nickmilder et</u> al. (2007)	Southern Belgium	July-August 2002	1-h max	NR	Max (across 6 camps): 24.5-112.7 ^b
			8-h max	NR	Max (across 6 camps): 18.9-81.1 ^b
<u>Delfino et al.</u> (2010a)	Los Angeles, CA	Warm and cold season 2005-2007	24-h avg	Warm season: 32.1 (median)°	Max: 76.4 [°] Max: 44.9 [°]
				Cool season: 19.1 (median) ^c	WIGA. 77.3
Adamkiewicz	Steubenville, OH	September-	24-h avg	15.3	75th: 20.2, Max: 32.2
<u>et al. (2004</u>)		December 2000	1-h avg ^d	19.8	75th: 27.5, Max: 61.6

* Note: Studies presented in order of first appearance in the text of this section.

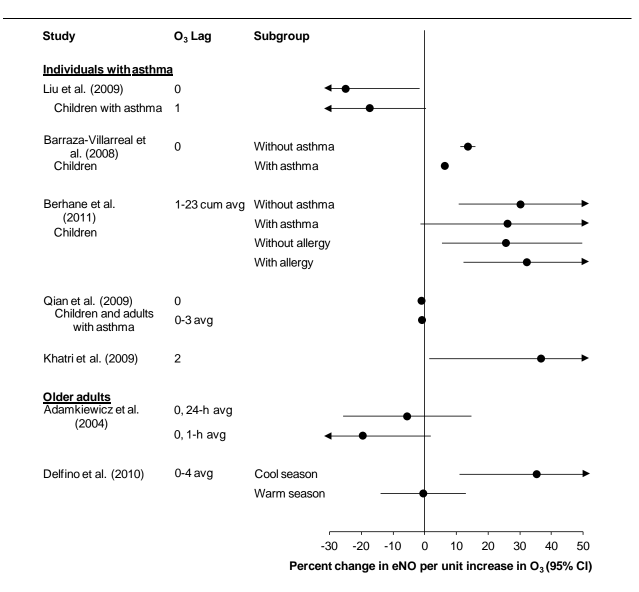
NR = Not Reported, SOCS = Salmeterol Off Corticosteroids Study.

^aIndividual-level estimates were calculated based on time spent in the vicinity of various O₃ monitors.

^bConcentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

^cMeasurements outside subject's residence (retirement home).

^dAverage O_3 concentration in the 1 hour preceding eNO collection.



Note: Results are presented first for children with asthma then for adults with asthma and older adults. Effect estimates are from single-pollutant models and are standardized to a 40-ppb increase for 1-h avg O_3 concentrations, a 30-ppb increase for 8-h max or 8-h avg O_3 concentrations, and a 20-ppb increase for 24-h avg O_3 concentrations.

Figure 6-10 Percent change in exhaled nitric oxide (eNO) in association with ambient ozone concentrations in populations with and without asthma.

Table 6-16	Additional characteristics and quantitative data for studies
	represented in Figure 6-10.

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Subgroup	Standardized % Change (95% CI) ^a
Studies in individ	uals with asthma				
<u>Liu et al. (2009a</u>)	Windsor, ON, Canada 182 children with asthma, ages 9- 14 yr	24-h avg	0 1		-25.1 (-42.9, -1.7) -17.5 (-32.1, -0.24)
Barraza-Villarreal et al. (2008)	Mexico City, Mexico 208 children, ages 6-14 yr	8-h max	0	Without asthma With asthma	13.5 (11.2, 15.8) 6.2 (6.0, 6.5)
Berhane et al. (2011)	13 Southern California communities 2,240 children, ages 6-10 yr	8-h avg (10 a.m6 p.m.)	1-23 cummulative avg	Without asthma With asthma Without allergy With allergy	30.1 (10.6, 53.2) 26.0 (-1.4, 60.9) 25.5 (5.3, 49.6) 32.1 (12.0, 55.9)
Qian et al. (2009)	Boston, MA; New York, NY; Denver, CO; Philadelphia, PA; San Francisco, CA; Madison, WI 119 children and adults with asthma, ages 12-65 yr	8-h max	0 0-3 avg		-1.2 (-1.7, -0.64) -1.0 (-1.8, -0.26)
Khatri et al. (2009)	Atlanta, GA 38 adults with asthma, ages 31- 50 yr	8-h max	2		36.6 (1.2, 71.9)
Studies in older a	dults				
Adamkiewicz et al. (2004)	Steubenville, Ohio 29 older adults, ages 53-90 yr	24-h avg 1-h avg ^⁵	0		-5.7 (-25.9, 14.5) -19.7 (-41.3, 1.9)
Delfino et al. (2010a)	Los Angeles, CA 60 older adults, ages ≥ 65 yr	24-h avg	0-4 avg	Cool season Warm season	35.2 (10.9, 59.5) -0.60 (-14.0, 12.8)

*Includes studies in Figure 6-10.

^aEffect estimates are standardized to a 40-ppb, 30-ppb, and 20-ppb increase for 1-h avg, 8-h max or avg, and 24-h avg O₃, respectively.

^bAverage O_3 concentration in the 1 hour preceding eNO collection.

Table 6-17Associations between short-term ambient ozone exposure and
biological markers of pulmonary inflammation and oxidative stress.

Study	Location/Population	O₃ Averaging Time	O₃ Lag	Biological Marker	Subgroup	Standardized Effect Estimate (95% CI) ^a
<u>Liu et al.</u> (2009a)	Windsor, ON, Canada 182 children with asthma, ages 9 - 14 yr	24-h avg	0	EBC 8-isoprostane (% change) EBC TBARS (% change)		16.2 (-14.9, 56.8) 11.5 (-27.0, 70.1)
<u>Romieu et al.</u> (2008)	Mexico City, Mexico 107 children with asthma, mean (SD) age 9.5 (2.1) yr	8-h max	0	EBC Malondialdehyde ^b		1.9 (1.1, 3.5)
<u>Barraza-</u> <u>Villarreal et al.</u> (2008)	Mexico City, Mexico 208 children, ages 6-14 yr	8-h max	0	Nasal lavage IL-8 (pg/mL)	Without asthma With asthma	1.6 (1.4, 1.8) 1.6 (1.4, 1.9)
				EBC pH	Without asthma With asthma	-0.10 (-0.27, 0.08) [°] -0.10 (-0.20, 0.01) [°]
Sienra-Monge et al. (2004)	Mexico City, Mexico 117 children with asthma, mean age 9 yr	8-h max	0-2 avg	Nasal lavage IL-8 ^b	Placebo Antioxidant	2.2 (1.1, 4.7) 1.0 (0.44, 2.3)
				Nasal lavage IL-6 ^b	Placebo Antioxidant	2.7 (1.4, 5.1) 1.1 (0.53, 2.2)
				Nasal lavage Uric acid ^b	Placebo Antioxidant	0.75 (0.44, 1.3) 1.3 (0.68, 2.4)
				Nasal lavage Glutathione ^b	Placebo Antioxidant	0.79 (0.63, 0.98) 0.80 (0.66, 0.96)
<u>Khatri et al.</u> (2009)	Atlanta, GA 38 adults with asthma, ages 31 - 50 yr	8-h max	2	Blood eosinophils (% change)		2.4 (0.62, 4.2)
Ferdinands et al. (2008)	Atlanta, GA 16 children exercising outdoors, ages 14 - 17 yr	1-h max	0	EBC pH (normalized score)		0.80 (-0.20, 1.8) ^c

Results generally are presented in order of increasing mean ambient O_3 concentration. EBC = exhaled breath condensate, TBARS = thiobarbituric acid reactive substances, IL-8 = interleukin 8, IL-6 = interleukin 6, Antioxidant = group supplemented with vitamins C and E.

^aEffect estimates are standardized to a 40-, 30- and 20-ppb increase for 1-h max, 8-h max, and 24-h avg O₃, respectively.

^bEffect estimates represent the ratio of the geometric means of biological marker per unit increase in O_3 concentration. A ratio <1 indicates a decrease in marker, and a ratio >1 indicates an increase in marker for an increase in O_3 .

^cModel analyzed log-transformed O₃. Decreases and increases in pH indicate increases and decreases in pulmonary inflammation, respectively.

Populations with Asthma

Exhaled Nitric Oxide

1	Neither NO nor eNO has been examined in controlled human exposure or toxicological
2	studies of O ₃ exposure. However, several lines of evidence support its analysis as an
3	indicator of pulmonary inflammation. Inducible NO synthase can be activated by
4	pro-inflammatory cytokines, and NO can be produced by cells such as neutrophils,
5	eosinophils, and epithelial cells in the lung during an inflammatory response (Barnes and
6	Liew, 1995). Further, eNO commonly is higher in individuals with asthma and increases
7	during acute exacerbations (Jones et al., 2001; Kharitonov and Barnes, 2000).
8	As indicated in Figure 6-10 and Table 6-16, short-term increases in ambient O_3
9	concentration (8-h max or avg) were associated with increases in eNO in children with
10	asthma. These studies used different methods to assign exposures using central site O_3
11	measurements: the site closest (within 5 km) to home or school (Barraza-Villarreal et al.,
12	2008) and a single site per community (Berhane et al., 2011). Because information on
13	spatial homogeneity of ambient O_3 concentrations and time spent outdoors was not
14	provided, it is not possible to assess whether these two methods differed in personal-
15	ambient O_3 ratios and correlations. <u>Liu et al. (2009a</u>) (described in Section <u>6.2.1.2</u>)
16	reported O ₃ -associated decreases in eNO; however, this study was restricted to winter.
17	Results for EBC markers of oxidative stress and lung function collectively provided weak
18	evidence of O_3 -associated respiratory effects in this study. As described in Section <u>4.3.3</u> ,
19	in non-summer months, indoor to outdoor O_3 ratios are lower as are personal-ambient
20	ratios, making it more difficult to detect associations with ambient O ₃ concentrations.
21	In contrast with controlled human exposure studies (Section $6.2.3.1$), epidemiologic
22	studies did not find larger O ₃ -associated increases in pulmonary inflammation in groups
23	with asthma than in groups without asthma (Figure 6-10 and Table 6-16). Among
24	children in Southern California, <u>Berhane et al. (2011</u>) estimated similar associations for a
25	1-23 day cumulative average of 8-h avg (10 a.m6 p.m.) O_3 in children with and without
26	asthma. Among children in Mexico City, Barraza-Villarreal et al. (2008) found a larger
27	association (for lag 0 of 8-max O ₃) in children without asthma, most of whom had atopy.
28	Studies that included adults with asthma produced contrasting results (Khatri et al., 2009;
29	<u>Qian et al., 2009</u>). The multicity salmeterol (β -2 agonist) trial (Boston, MA; New York,
30	NY; Denver, CO; Philadelphia, PA; San Francisco, CA; and Madison, WI) involved
31	serial collection of eNO from 119 subjects with asthma, 87% of whom were 20-65 years
32	of age (Qian et al., 2009). Ambient O_3 concentrations were averaged from all sites within
33	20 miles of subjects' zipcode centroids, which in a repeated measures study, may capture
34	the temporal variation in O ₃ reasonably well (<u>Darrow et al., 2011a; Gent et al., 2003</u>).

1 Among all subjects, increases in 8-h max O_3 at multiple lags (0 to 3 single-day and 2 0-4 avg) were associated with decreases in eNO. Results did not vary among the 3 salmeterol-, CS-, and placebo-treated groups, indicating that the counterintuitive findings 4 for O_3 were not only due to the reduction of inflammation by medication. Oian et al. 5 (2009) suggested that at higher concentrations, O₃ may transform NO in airways to 6 reactive nitrogen species. However, this hypothesis was not supported by results from 7 Khatri et al. (2009), who in Atlanta, GA examined overall higher 8-h max O_3 ambient 8 concentrations than did Qian et al. (2009) and found that an increase in O₃ was associated 9 with an increase in eNO in adults with asthma (36.6% [95% CI: 1.2, 71.9] per 30-ppb 10 increase in lag 2 of 8-h max O₃). Although Khatri et al. (2009) was cross-sectional and 11 did not adjust for any meteorological factors, it may have better characterized O₃ 12 exposures because subjects were examined during warm months, and an 8-h max O₃ 13 concentration was calculated for each subject using measurements at the site closest to 14 his/her location each hour.

Other biological markers of pulmonary inflammation and oxidative stress

- 15Short-term increases in ambient O3 concentration were associated with changes in the16levels of pro-inflammatory cytokines and cells, indicators of oxidative stress, and17antioxidants (
- 18 Table 6-17). Importantly, any particular biomarker was examined in only one to two 19 studies, and the evidence in individuals with asthma is derived primarily from studies 20 conducted in Mexico City (Barraza-Villarreal et al., 2008; Romieu et al., 2008; Sienra-21 Monge et al., 2004). These studies measured ambient O_3 concentrations at sites within 5 22 km of subjects' schools or homes. In a Mexico City cohort of children with asthma, 23 school ambient O_3 concentrations averaged over 48 to 72 hours had a ratio and 24 correlation with personal exposures (48- to 72-h avg) of 0.17 and 0.35, respectively 25 (Ramírez-Aguilar et al., 2008). These observations suggest that the effects of personal O_3 26 exposure on inflammation may have been underestimated in the Mexico City studies. 27 Despite the limited evidence, the epidemiologic findings are well supported by controlled 28 human exposure and toxicological studies that measured the same or related endpoints.
- 29Several of the modes of action of O_3 are mediated by reactive oxygen species (ROS)30produced in the airways by O_3 (Section 5.3.3). These ROS are important mediators of31inflammation as they regulate cytokine expression and inflammatory cell activity in32airways (Heidenfelder et al., 2009). Controlled human exposure and toxicological studies,33frequently have found O_3 -induced increases in oxidative stress as shown by increases in34prostaglandins (Section 5.3.3 and Section 6.2.3.1), which are produced by the35peroxidation of cell membrane phospholipids (Morrow et al., 1990). Romieu et al. (2008)

1	analyzed EBC malondialdehyde (MDA), a thiobarbituric acid reactive substance, which
2	like prostaglandins, is derived from lipid peroxidation (Janero, 1990). For a 30-ppb
3	increase in lag 0 of 8-h max O_3 , the ratio of the geometric means of MDA was 1.3
4	(95% CI: 1.0, 1.7). Similar results were reported for lags 1 and 0-1 avg O_3 . A limitation
5	of the study was that 25% of EBC samples had nondetectable levels of MDA, and the
6	random assignment of concentrations between 0 and 4.1 nmol to these samples may have
7	contributed random measurement error to the estimated O_3 effects. Because MDA
8	represents less than 1% of lipid peroxides and is present at low concentrations, its
9	biological relevance has been questioned. However, <u>Romieu et al. (2008</u>) pointed to their
10	observations of statistically significant associations of EBC MDA levels with nasal
11	lavage IL-8 levels to demonstrate its relationship with pulmonary inflammation.
12	Uric acid and glutathione are ROS scavengers that are present in the airway ELF. While
13	the roles of these markers in the inflammatory cascade of asthma are not well defined,
14	they have been observed to be consumed in response to short-term O_3 exposure as part of
15	an antioxidant response in controlled human exposure and animal studies (Section 5.3.3).
16	Results from an epidemiologic study also indicate that a similar antioxidant response may
17	be induced by increases in ambient O_3 exposure.2004) Sienra-Monge et al. (2004) found
18	O_3 -associated decreases in nasal lavage levels of uric acid and glutathione in children
19	with asthma not supplemented with antioxidant vitamins (
20 21	Table 6-17). The magnitudes of decrease were similar for O_3 concentrations lagged 2 or 3 days and averaged over 3 days.
22	Both controlled human exposure and toxicological studies have found O_3 -induced
23	increases in the cytokines IL-6 and IL-8 (Section 5.3.3, Section 6.2.3.1, and
24	Section 6.2.3.3), which are involved in initiating an influx of neutrophils, a hallmark of
25	O_3 -induced inflammation (Section 6.2.3.1). Epidemiologic studies conducted in Mexico
26	City had similar findings. Sienra-Monge et al. (2004) found that 8-h max O_3 was
27	associated with increases in nasal lavage levels of IL-6 and IL-8 (placebo group), with
28	larger effects estimated for lag 0-2 avg than for lag 2 or 3 O_3 (
29 30 31 32	Table 6-17). In another cohort of children with asthma, a 30-ppb increase in lag 0 of 8-h max O ₃ was associated with a 1.61 pg/mL increase (95% CI: 1.4, 1.8) in nasal lavage levels of IL-8 (<u>Barraza-Villarreal et al., 2008</u>). This study also reported a small O ₃ -associated decrease in EBC pH (
33 34 35 36	Table 6-17). EBC pH, which is thought to reflect the proton-buffering capacity of ammonium in airways, decreases upon asthma exacerbation, and is negatively correlated with airway levels of pro-inflammatory cytokines (<u>Carpagnano et al., 2005</u> ; <u>Kostikas et al., 2002</u> ; <u>Hunt et al., 2000</u>).

1 2 3 4 5 6 7	Albeit with limited investigation, controlled human exposure studies have found O_3 -induced increases in eosinophils in adults with asthma (Section <u>6.2.3.1</u>). Eosinophils are believed to be the main effector cells that initiate and sustain inflammation in asthma and allergy (<u>Schmekel et al., 2001</u>). Consistent with these findings, in a cross-sectional study of adults with asthma in Atlanta, GA, a 30-ppb increase in lag 2 of 8-h max O_3 was associated with a 2.4% increase (95% CI: 0.62, 4.2) in blood eosinophils (<u>Khatri et al., 2009</u>). Potential confounding by weather was not evaluated in models.
8 9 10 11 12 13 14 15 16 17	The prominent influences demonstrated for ROS and antioxidants in mediating the respiratory effects of O_3 provide biological plausibility for effect modification by antioxidant capacity. Effect modification by antioxidant capacity has been described for O_3 -associated lung function in controlled human exposure and epidemiologic studies (Section <u>6.2.1.1</u> and Section <u>6.2.1.2</u>). An epidemiologic study conducted in Mexico City also found that vitamin C and E supplements, which potentially increase antioxidant capacity, attenuated O_3 -associated inflammation and oxidative stress. Among children with asthma supplemented daily with vitamin C and E, the ratios of the geometric means of nasal lavage IL-6 and IL-8 per 30-ppb increases in lag 0-2 avg of 8-h max O_3 were 1.0, reflecting no change with increases in O_3 concentration (
18 19 20 21 22 23 24 25 26	Table 6-17) (Sienra-Monge et al., 2004). The results did not clearly delineate interactions among O ₃ concentrations, endogenous antioxidants, and dietary antioxidants (Table 6-17). Ozone was associated with increases in uric acid in the antioxidant group but decreases in the placebo group across the O ₃ lags examined. Associations with glutathione were similar in the two groups. In another cohort, 8-h max O ₃ concentrations \geq 38 ppb enhanced the effects of diets high in antioxidant vitamins and/or omega-3 fatty acids on protecting against O ₃ -related increases in nasal lavage IL-8 (Romieu et al., 2009). Information on the main effects of O ₃ or effect modification by diet was not presented.
27 28 29 30 31 32 33 34 35 36	The levels of several biological markers such as eNO, EBC pH, and MDA consistently differ between groups with and without asthma and change during an asthma exacerbation (Corradi et al., 2003; Hunt et al., 2000); however, the magnitudes of change associated with these overt effects are not well defined. Ozone-associated increases in interleukins and indicators of oxidative stress were small: 1-2% increase for each 30-ppb increase in 8-h max O ₃ concentration (Table 6-17). Ozone-associated increases in eNO were larger: 6-36% increase per 30-ppb increase in 8-h max ambient O ₃ concentration (Berhane et al., 2011; Delfino et al., 2010a; Khatri et al., 2009; Barraza-Villarreal et al., 2008). Some studies in populations with asthma found O ₃ -associated increases in pulmonary inflammation concurrently (at the

1same lag) with O3-associated in respiratory symptoms. For example, among adults with2asthma in Atlanta, an increase in ambient O3 concentration was associated with increases3in eNO, blood eosinophils, and a decrease in quality of life score, which incorporates4indices for symptoms and activity limitations (Khatri et al., 2009). Also, among children5with asthma in Mexico City, O3 was associated with increases in eNO and nasal lavage6IL-8 and concurrently assessed cough but not wheeze (Barraza-Villarreal et al., 2008).

Children without Asthma

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- In the limited investigation, short-term increases in ambient O_3 concentration (8-h max or avg) were associated with increases in pulmonary inflammation in children without asthma (Berhane et al., 2011; Barraza-Villarreal et al., 2008) (Figure 6-10 and Table 6-16 and
- 11 Table 6-17). The study of children in Mexico City found a larger O₃-associated increase 12 in eNO in the children without asthma than with asthma (13.5% versus 6.2% increase per 13 30-ppb increase in lag 0 of 8-h max O₃) (Barraza-Villarreal et al., 2008). Ozone was 14 associated with similar magnitudes of change in IL-8 and EBC pH in children with and 15 without asthma. A distinguishing feature of this study was that 72% of children without asthma had allergies. A study conducted in 13 Southern California communities also 16 17 found that increases in ambient O₃ concentration (8-h avg, 10 a.m.-6 p.m.) were 18 associated with increases in eNO in children with respiratory allergy (Berhane et al., 19 <u>2011</u>). Coherence for these epidemiologic findings is provided by observations of 20 O_3 -induced allergic inflammation in animal models of allergy (Section <u>6.2.3.3</u> and 21 Section 6.2.6).
- 22 Berhane et al. (2011) found O₃-associated increases in eNO in children without asthma 23 and children without respiratory allergy, providing evidence for effects on pulmonary 24 inflammation in healthy children. This study provided detailed information on differences 25 in association among various lags of 8-h avg (10 a.m.-6 p.m.) O₃. Ozone concentrations 26 averaged over the several hours preceding eNO collection were not significantly 27 associated with eNO. Consistent with other studies examining pulmonary inflammation 28 and oxidative stress, Berhane et al. (2011) found that relatively short lags of O_3 , i.e., 1 to 29 5 days, were associated with increases in eNO. However, among several types of lag-30 based models, including unconstrained lag models, polynomial distributed lag models, 31 spline-based distributed lag models, and cumulative lag models, a 23-day cumulative lag 32 of O₃ best fit the data. Among the studies evaluated in this ISA, Berhane et al. (2011) was 33 unique in evaluating and finding larger respiratory effects for multi-week (e.g., 13-34 30 days) average O_3 concentrations. The mechanism for the effects of O_3 peaking with a

23-day cumulative lag of exposure has not been delineated. Further, with examination of such long lag periods, there is greater potential for residual confounding by weather.

Populations with Increased Outdoor Exposures

3 With limited investigation, increases in ambient O_3 concentration were not consistently 4 associated with pulmonary inflammation in populations engaged in outdoor activity or 5 exercise. Common limitations of these studies were the small numbers of subjects and 6 lack of consideration for potential confounding factors. A study in 16 adolescent long-7 distance runners near Atlanta, GA was noted for the daily collection of EBC and the 8 likely greater extent to which ambient O_3 concentrations represented ambient exposures 9 because of the analysis of O₃ concentrations measured during outdoor running at a site 10 less than 1 mile from the exercise track (Ferdinands et al., 2008). Increases in 1-h max O_3 11 (lags 0 to 2) were associated with increases in EBC pH, indicating O₃-associated 12 decreases in pulmonary inflammation. Among 9 adult male runners in Sicily, Italy 13 examined 3 days before and 20 hours after 3 races in fall, winter, and summer, weekly 14 average O₃ concentrations (8-h avg, 7 a.m.-3 p.m.) were positively correlated with 15 apoptosis of neutrophils (Spearman's r = 0.70, p < 0.005) and bronchial epithelial cell 16 differential counts (Spearman's r = 0.47, p < 0.05) but not with neutrophil or macrophage 17 cell counts or levels of the pro-inflammatory cytokines TNF- α and IL-8 (Chimenti et al., 18 2009). Associations with O_3 concentrations measured during the races (mean 35 to 89 19 minutes) were not examined. This study provides evidence for new endpoints; however, 20 the implications of findings are limited due to the lack of a rigorous statistical analysis.

21 In a cross-sectional study of children at camps in south Belgium, although lung function 22 was not associated with O_3 measured at camps during outdoor activity, an association 23 was found for eNO (Nickmilder et al., 2007). Children at camps with lag 0 1-h max O_3 24 concentrations >85.2 ppb had greater increases in intraday eNO compared with children 25 at camps with O_3 concentrations <51 ppb. A benchmark dose analysis indicated that the 26 threshold for an O₃-associated increase of 4.3 ppb eNO (their definition of increased 27 pulmonary inflammation) was 68.6 ppb for 1-h max O_3 and 56.3 ppb for 8-h max O_3 . 28 While these results provide additional evidence for O_3 -associated increases in pulmonary 29 inflammation in healthy children, they should be interpreted with caution since they were 30 unadjusted for any potential confounding factors and based on camp-level comparisons.

Older Adults

31The panel studies examining O3-associated changes in eNO in older adults produced32contrasting findings (Figure 6-10 and Table 6-16). The studies differed with respect to33geographic location, inclusion of healthy subjects, exposure assessment method, and lags

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1	of O_3 examined. Delfino et al. (2010a) followed 60 older adults with coronary artery
2	disease in the Los Angeles, CA area for 6 weeks each during a warm and cool season; the
3	specific months were not specified. Ambient O3 was measured at subjects' retirement
4	homes, possibly reducing some exposure measurement error due to spatial variability.
5	Multiday averages of O_3 (3- to 9-day) were associated with increases in eNO, with effect
6	estimates increasing with increasing number of averaging days. In contrast with most
7	other studies, an association was found in the cool season but not warm season (increase
8	in eNO per 20-ppb increase in lag 0-4 avg of 24-h avg O ₃ : 4.1 ppb [95% CI: 1.3, 6.9] in
9	cool season, -0.01 ppb [95% CI: -2.3, 2.1] in warm season). Despite these unusual
10	findings for the cool season, they were similar to findings from another study of
11	Los Angeles area adults with asthma, which indicated an O3-associated decrease in
12	indoor activity during the fall season (Eiswerth et al., 2005).

- 13In a cool season (September-December) study conducted in older adults (ages 54-1491 years) in Steubenville, OH, Adamkiewicz et al. (2004) found that increases in O315(1-h avg and 24-h avg before eNO collection) were associated with decreases in eNO,16reflecting decreases in pulmonary inflammation (Figure 6-10 and
- 17Table 6-17). The study included healthy adults and those with asthma or COPD. A study18in a subset of these adults illustrated why it is difficult to detect effects with central site19 O_3 concentrations in the cool season by showing that subjects spent $\geq 90\%$ of time20indoors and >77% at home and had a mean 24-h avg O_3 personal-ambient ratio of 0.2721(Sarnat et al., 2006a).

Confounding in Epidemiologic Studies of Pulmonary Inflammation and Oxidative Stress

- 22 Except where noted in the preceding text; epidemiologic studies of pulmonary 23 inflammation and oxidative stress accounted for potential confounding by meteorological 24 factors. Increases in ambient O₃ concentration were associated with pulmonary 25 inflammation or oxidative stress in models that adjusted for temperature and/or humidity 26 (Delfino et al., 2010a; Barraza-Villarreal et al., 2008; Romieu et al., 2008). Final results 27 from Sienra-Monge et al. (2004) and Berhane et al. (2011) were not adjusted for 28 temperature because associations were not altered by adjustment for temperature. Most 29 studies conducted over multiple seasons adjusted for season or time trend.
- 30In evidence limited to a small number of studies conducted in Mexico City, O_3 -associated31pulmonary inflammation and oxidative stress were not found to be confounded by $PM_{2.5}$ 32or PM_{10} . These studies, which analyzed 8-hour averages for both O_3 and PM, found33robust associations for O_3 (Barraza-Villarreal et al., 2008; Romieu et al., 2008; Sienra-34Monge et al., 2004). Ozone and PM, both measured at central sites located within 5 km of

1	subjects' schools or homes, were moderately correlated ($r = 0.46-0.54$). Weak
2	correlations have been found between personal exposures of O_3 and $PM_{2.5}$
3	(Section <u>4.3.4.1</u>). Only <u>Romieu et al. (2008</u>) provided quantitative results. Lag 0 of
4	8-h max O_3 was associated with the same magnitude of increase in MDA without and
5	with adjustment for lag 0 of 8-h max PM _{2.5} (ratio of geometric means for a 30-ppb
6	increase: 1.3 [95% CI: 1.0, 1.7]). In comparison, the O ₃ -adjusted effect estimate for PM _{2.5}
7	was cut in half.

Summary of Epidemiologic Studies of Pulmonary Inflammation and Oxidative Stress

- 8 Many epidemiologic studies provided evidence that short-term increases in ambient O_3 9 exposure increase pulmonary inflammation and oxidative stress in children with asthma, with evidence primarily provided by studies conducted in Mexico City. By also finding 10 11 that associations were attenuated with higher antioxidant intake, these studies indicated 12 that inhaled O₃ may be an important source of ROS in airways and/or may increase 13 pulmonary inflammation via oxidative stress-mediated mechanisms. Studies found 14 O₃-associated increases in pulmonary inflammation in children with allergy (Berhane et 15 al., 2011; Barraza-Villarreal et al., 2008). The limited available evidence in children and 16 adults with increased outdoor exposures and older adults was inconclusive. Temperature 17 and humidity were not found to confound O₃ associations. Copollutant models were 18 analyzed in a few studies conducted in Mexico City; O₃ effect estimates were robust to 19 adjustment for moderately correlated (r = 0.46-0.54) $PM_{2.5}$ or PM_{10} (Barraza-Villarreal et 20 al., 2008; Romieu et al., 2008; Sienra-Monge et al., 2004).
- 21 Ozone-associated increases in pulmonary inflammation and oxidative stress were found 22 in studies that used varied exposure assessment methods: measurement on site of 23 subjects' outdoor activity (Nickmilder et al., 2007), average of concentrations measured 24 at the closest site each hour Khatri et al. (2009), measurement at a site within 5 km of 25 subjects' schools or homes (Barraza-Villarreal et al., 2008; Romieu et al., 2008; Sienra-26 Monge et al., 2004), and measurement at single site per town (Berhane et al., 2011). 27 While these methods may differ in the degree of exposure measurement error, in the 28 limited body of evidence, there was not a clear indication that the method of exposure 29 assessment influenced the strength or magnitude of associations.
- 30Most studies examined and found associations with 8-h max or daytime 8-h avg O331concentrations, although associations were found for 1-h max (Nickmilder et al., 2007)32and 24-h avg O3 (Delfino et al., 2010a). Collectively, studies examined single-day O333concentrations lagged from 0 to 5 days and concentrations averaged over 2 to 9 days. Lag340 of 8-h max O3 was most frequently examined and consistently associated with

1	pulmonary inflammation and oxidative stress. However, in the few studies that examined
2	multiple O ₃ lags, multiday average 8-h max or 8-h avg concentrations were associated
3	with larger increases in pulmonary inflammation and oxidative stress (Berhane et al.,
4	2011; Delfino et al., 2010a; Sienra-Monge et al., 2004). These findings for multiday
5	average O_3 concentrations are supported by controlled human exposure (Section <u>6.2.3.1</u>)
6	and animal studies (Section $6.2.3.3$) that similarly have found that some markers of
7	pulmonary inflammation remain elevated with O ₃ exposures repeated over multiple days.
8	Several epidemiologic studies concurrently examined associations of ambient O ₃
9	concentrations with biological markers of pulmonary inflammation and lung function or
10	respiratory symptoms. Whether evaluated at the same or different lags of O ₃ , associations
11	generally were stronger for biological markers of airway inflammation than for lung
12	function (Khatri et al., 2009; Barraza-Villarreal et al., 2008; Nickmilder et al., 2007).
13	Controlled human exposure studies also have demonstrated a lack of correlation between
14	inflammatory and spirometric responses induced by O_3 exposure (Section <u>6.2.3.1</u>).
15	Evidence has suggested that O3-related respiratory morbidity may occur via multiple
16	mechanisms with varying time courses of action, and the examination of a limited
17	number of O ₃ lags in these aforementioned studies may explain some of the
18	inconsistencies in associations of O3 with measures of pulmonary inflammation and lung
19	function. In contrast, based on examination in a few studies, increases in ambient O_3
20	concentration were associated concurrently (at the same lag) with increases in pulmonary
21	inflammation and increases in respiratory symptoms or activity limitations in the same
22	population of individuals with asthma (Khatri et al., 2009; Barraza-Villarreal et al.,
23	<u>2008</u>).

6.2.3.3 Toxicology: Inflammation and Injury

24	The 2006 O ₃ AQCD states that the "extensive human clinical and animal toxicological
25	evidence, together with the limited available epidemiologic evidence, is clearly indicative
26	of a causal role for O_3 in inflammatory responses in the airways" (U.S. EPA, 2006b).
27	Airway ciliated epithelial cells and Type 1 cells are the most O3-sensitive cells and are
28	initial targets of O_3 . These cells are damaged by O_3 and produce a number of
29	pro-inflammatory mediators (e.g., interleukins [IL-6, IL-8], PGE ₂) capable of initiating a
30	cascade of events leading to PMN influx into the lung, activation of alveolar
31	macrophages, inflammation, and increased permeability across the epithelial barrier. One
32	critical aspect of inflammation is the potential for metaplasia and alterations in
33	pulmonary morphology. Studies have observed increased thickness of the alveolar septa,
34	presumably due to increased cellularity after acute exposure to O ₃ . Epithelial hyperplasia
35	starts early in exposure and increases in magnitude for several weeks, after which it

- 1 plateaus until exposure ceases. When exposure persists for a month and longer, excess 2 collagen and interstitial fibrosis are observed. This response, discussed in Chapter 7, 3 continues to increase in magnitude throughout exposure and can even continue to 4 increase after exposure ends (Last et al., 1984). Previously reviewed toxicological studies 5 of the ability of O_3 to cause inflammation, injury, and morphological changes are 6 described in Table 6-5 on page 6-25 (U.S. EPA, 1996f) and Table 6-10 (U.S. EPA, 7 1996k) and Table 6-11 (U.S. EPA, 1996l) beginning on page 6-61 of the 1996 O₃ AQCD, 8 and Tables AX5-8 (U.S. EPA, 2006d) and AX5-9 (U.S. EPA, 2006e), beginning on page 9 AX5-17 of the 2006 O₃ AQCD. Numerous recent in vitro and in vivo studies add to this 10 very large body of evidence for O_3 -induced inflammation and injury, and provide new 11 information regarding the underlying mechanisms (see Section 5.3).
- 12 A number of species, including dogs, rabbits, guinea pigs, rats, and mice have been used 13 as models to study the pulmonary effects of O_3 , but the similarity of non-human primates 14 to humans makes them an attractive model in which to study the pulmonary response to 15 O₃. As reviewed in the 1996 and 2006 O₃ AQCDs, several pulmonary effects, including 16 inflammation, changes in morphometry, and airway hyperresponsiveness, have been 17 observed in macaque and rhesus monkeys after acute exposure to O_3 (Table 6-18 presents 18 a highlight of these studies). Increases in inflammatory cells were observed after a single 19 8-h exposure of adult rhesus monkeys to 1 ppm O₃ (Hyde et al., 1992). Inflammation was 20 linked to morphometric changes, such as increases in necrotic cells, smooth muscle, 21 fibroblasts, and nonciliated bronchiolar cells, which were observed in the trachea, 22 bronchi, or respiratory bronchioles. Effects have also been observed after short-term 23 repeated exposure to O_3 at concentrations that are more relevant to ambient O_3 24 concentrations. Morphometry changes in the lung, nose, and vocal cords were observed 25 after exposure to 0.15 ppm O_3 for 8-h/day for 6 days (Harkema et al., 1993; Dimitriadis, 26 1992; Harkema et al., 1987a). Since 2006, however, only one study has been published 27 regarding acute exposure of non-human primates to O_3 (a number of recent chronic 28 studies in non-human primates are described in Chapter 7). In this study, a single 6-hour 29 exposure of adult male cynomolgus monkeys to 1 ppm O₃ induced significant increases 30 in inflammatory and injury markers, including BAL neutrophils, total protein, alkaline 31 phosphatase, IL-6, IL-8, and G-CSF (Hicks et al., 2010a). Gene expression analysis 32 confirmed the increases in the pro-inflammatory cytokine IL-8, which had been 33 previously described in O_3 exposed rhesus monkeys (Chang et al., 1998). The 34 anti-inflammatory cytokine IL-10 was also elevated, but the fold changes in IL-10 and 35 G-CSF were relatively low and highly variable. The single exposure also caused necrosis 36 and sloughing of the epithelial lining of the most distal portions of the terminal 37 bronchioles and the respiratory bronchioles. Bronchiolitis, alveolitis, parenchymal and 38 centriacinar inflammation were also observed. A second exposure protocol (two 39 exposures with a 2-week inter-exposure period) resulted in similar inflammatory

responses, with the exception of total protein and alkaline phosphatase levels which were attenuated, indicating that attenuation of some but not all lavage parameters occurred upon repeated exposure of non-human primates to O_3 (Hicks et al., 2010a). This variability in attenuation is similar to the findings of earlier reports in rodents (Wiester et al., 1996c) and non-human primates (Tyler et al., 1988).

6 Table 6-18 describes key morphometric studies conducted in non-human primates 7 exposed to O_3 . Morphologic observations made by <u>Dungworth (1976)</u>; (<u>1975</u>) indicate 8 that the rat and Bonnet monkey (Macaca radiata) are approximately equal in 9 susceptibility to short-term effects of O₃. Mild but discernible lesions were caused in both 10 species by exposure to 0.2 ppm O₃ for 8 h/day for 7 days. The authors stated that 11 detectable morphological effects in the rat occurred at levels as low as 0.1 ppm O_3 . In 12 both species, the lesion occurred at the junction of the small airways and the gaseous 13 exchange region. In rats, the prominent features were accumulation of macrophages, 14 replacement of necrotic Type 1 epithelial cells with Type 2 cells, and damage to ciliated 15 and nonciliated Clara cells. The principal site of damage was the alveolar duct. In 16 monkeys, the prominent O_3 -induced injury was limited to the small airways. At 0.2 ppm 17 O_3 , the lesion was observed at the proximal portion of the respiratory bronchioles. As 18 concentrations of O_3 were increased up to 0.8 ppm, the severity of the lesion increased, 19 and the damage extended distally to involve the proximal portions of the alveolar duct. 20 Mellick et al. (1977) found similar but more pronounced effects when rhesus monkeys (3 21 to 5 years of age) were exposed to 0.5 and 0.8 ppm O_3 , 8 hours/day for 7 days. In these 22 experiments, the respiratory bronchioles were the most severely damaged, and more 23 distal parenchymal regions were unaffected. Major effects were hyperplasia and 24 hypertrophy of the nonciliated bronchiolar epithelial cells and the accumulation of 25 macrophages intraluminally. In mice, continuous exposure to 0.5 ppm O_3 caused nodular 26 hyperplasia of Clara cells after 7 days of exposure. Similar findings were reported by 27 Schwartz (1976) and Schwartz et al. (1976), who exposed rats to 0.2, 0.5 or 0.8 ppm O_3 28 for 8 or 24 hours/day for 1 week. Changes observed within the proximal alveoli included 29 infiltration of inflammatory cells and swelling and necrosis of Type 1 cells. In the 30 terminal bronchiole, the changes reported were shortened cilia, clustering of basal bodies 31 in ciliated cells suggesting ciliogenesis, and reduction in height or loss of cytoplasmic 32 luminal projection of the Clara cells. Effects were seen at O_3 concentrations as low as 33 0.2 ppm. A dose-dependent pulmonary response to the three levels of O_3 was evident. No 34 differences were observed in morphologic characteristics of the lesions between rats 35 exposed continuously and those exposed intermittently for 8 hours/day.

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Reference	O ₃ concentration (ppm)	Exposure duration	Species, Sex, Age	Observation
<u>Harkema et al.</u> (1993)	0.15	8 h/day for 6 days	<i>Macaca radiata</i> (bonnet macaques) 2-6 years old	Several fold increase in thickness of surface epithelium in respiratory bronchioles; increase in interstitial mass with increase in proportion of cuboidal cells.
<u>Harkema et al.</u> (1987a); (1987b)	0.15	8 h/day for 6 days	<i>Macaca radiata</i> , M, F 2-6 years old	Ciliated cell necrosis, shortened cilia, and increased mucous cells in the respiratory epithelium of nose after 0.15 ppm; changes in nonciliated cells, intraepithelial leukocytes, and mucous cells in the transitional epithelium
Dungworth (1976)	0.2 0.5 0.8	8 h/day for 7 days for monkey and rat; continuous at 0.5 ppm for 7 days for mouse	Adult Rhesus and bonnet monkeys; S-D rats; Mice	In both rats and monkeys mild but discernible lesions were observed at 0.2 ppm; similar severity between species but different site of lesions – respiratory bronchioles for monkey and damage to ciliated, Clara, and alveolar epithelial cells for rat; Clara cell hyperplasia in mice
<u>Leonard et al.</u> (1991)	0.25	8 h/day for 7 days	Macaca radiata age not specified	The O_3 exposure level is not clear – the abstract states 0.64 ppm, but the text mentions only 0.25 ppm. Morphometric changes in vocal cord mucosa: disruption and hyperplasia of stratified squamous epithelium; epithelial and connective tissue thickness increased
<u>Chang et al.</u> (1998)	0.96	8 h	Rhesus, M age not specified	Increase in IL-8 in airway epithelium correlated with PMN influx
<u>Hyde et al. (1992</u>)	0.96	8 h	Rhesus, M 2 - 8.5 years old	Increased PMNs; morphometric changes in trachea, conducting airways, respiratory bronchioles including increased smooth muscle in bronchi and RB.
<u>Hicks et al.</u> (2010b)	1.0	6 h	Cynomolgus, M 5-7 kg (Adult)	Increase in PMNs and IL-8 in lavage fluid

Table 6-18Morphometric observations in non-human primates after acute
ozone exposure.

1	Exposure of adult BALB/c mice to 0.1 ppm O ₃ for 4 hours increased BAL levels of
2	keratinocyte chemoattractant (KC; IL-8 homologue) (~ fold), IL-6 (~12-fold), and TNF- α
3	(~ 2-fold) (<u>Damera et al., 2010</u>). Additionally, O_3 increased BAL neutrophils by 21%
4	without changes in other cell types. A trend of increased neutrophils with increased O ₃
5	concentration (0.12-2 ppm) was observed in BALB/c mice exposed for 3 hours (Jang et
6	al., 2005). Although alterations in the epithelium of the airways were not evident in 129J
7	mice after 4 hours of exposure to 0.2 ppm O_3 (Plopper et al., 2006), detachment of the
8	bronchiolar epithelium was observed in SD rats after 5 days or 60 days of exposure to
9	0.25 ppm O_3 (Oyarzún et al., 2005). Subacute (65 hours) exposure to 0.3 ppm O_3 induced
10	pulmonary inflammation, cytokine induction, and enhanced vascular permeability in wild
11	type mice of a mixed background (129/Ola and C57BL/6) and these effects were
12	exacerbated in metallothionein I/II knockout mice (Inoue et al., 2008). Three hours or
13	72 hours of exposure to 0.3 ppm O_3 resulted in similar levels of IL-6 expression in the
14	lungs of C57BL/6 mice (Johnston et al., 2005b), along with increases in BAL protein,

1	sTNFR1, and sTNFR2. Increased neutrophils were observed only after the 72-hour
2	exposure, and neither exposure resulted in detectable levels of IL-6 or KC protein. Levels
3	of BAL protein, sTNFR1, and sTNFR2 were higher in the 72-hour exposure group than
4	in the 3-hour exposure group. In another study, the same subacute (72 hours) exposure
5	protocol elicited increases in BALF protein, IP-10, sTNFR1, macrophages, neutrophils,
6	and IL-6, IL-1 α , and IL-1 β expression (Johnston et al., 2007). Yoon et al. (2007) exposed
7	C57BL/6J mice continuously to 0.3 ppm O_3 for 6, 24, 48, or 72 hours, and observed
8	elevated levels of KC, MIP-2, metalloproteinases, and inflammatory cells in the lungs at
9	various time points. A similar exposure protocol using C3H/HeJ and C3H/OuJ mice
10	demonstrated elevations in protein, PMNs, and KC, which were predominantly TLR 4
11	pathway dependent based on their prominence in the TLR 4 sufficient C3H/OuJ strain
12	Bauer et al. (2011). C3H/OuJ mice also had elevated levels of the heat-shock protein
13	HSP70, and further experiments in HSP70 deficient mice indicated a role for this
14	particular pathway in O_3 -related injury, discussed in more detail in Chapter 5.
	F
15	As reviewed in the 2006 O_3 AQCD, the time course for changes in BAL depends on the
16	parameters being studied. Similarly, after exposing adult C57BL mice to 0.5 ppm O_3 for
17	3 hours, <u>Han et al. (2008</u>) observed early (5 hours postexposure) increases in BAL TNF- α
18	and IL-1 β , which diminished by 24 hours postexposure. Total BAL protein was elevated
19	at 24 hours, but there were only minimal or negligible changes in LDH, total cells, or
20	PMNs. Ozone increased BAL mucin levels (with statistical significance by 24 hours
21	postexposure), and significantly elevated surfactant protein D at both time points. Prior
22	intratracheal (IT) exposure to multiwalled carbon nanotubes enhanced most of these
23	effects, but the majority of responses to the combined exposure were not greater than
24	those to nanotubes alone. Ozone exposure did not induce markers of oxidative stress in
25	lung tissue, BAL, or serum. Consistent with this study, Aibo et al. (2010) did not detect
26	changes in BAL inflammatory cell numbers in the same mouse strain after a 6-hour
27	exposure to 0.25 or 0.5 ppm. The majority of inflammatory cytokines (pulmonary or
28	circulating) were not significantly changed (as assessed 9 hours post-O ₃ exposure).
29	Exposure of C57BL/6 mice to 1 ppm for 3 hours increased BAL total cells, neutrophils,
30	and KC; these responses were greatest at 24 hours postexposure. F2-isoprostane
31	(8-isoprostane), a marker of oxidative stress, was also elevated by O_3 , peaking at
32	48 hours postexposure (<u>Voynow et al., 2009</u>).
33	Atopic asthma appears to be a risk factor for more severe O_3 induced airway
34	inflammation in humans (Balmes et al., 1997; Scannell et al., 1996), and allergic animal
35	models are often used to investigate the effects of O_3 on this potentially at-risk
36	population. Farraj et al. (2010) exposed allergen-sensitized adult male BALB/c mice to
37	0.5 ppm O_3 for 5 hours once per week for 4 weeks. Ovalbumin-sensitized mice exposed
38	to O_3 had significantly increased BAL eosinophils by 85% and neutrophils by 103%
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1	relative to OVA sensitized mice exposed to air, but these changes were not evident upon
2	histopathological evaluation of the lung, and no O ₃ induced lesions were evident in the
3	nasal passages. Ozone increased BAL levels of N-acetyl-glucosaminidase (NAG; a
4	marker of injury) and protein. DEP co-exposure (2.0 mg/m ³ , nose only) inhibited these
5	responses. These pro-inflammatory effects in an allergic mouse model have also been
6	observed in rats. Wagner et al. (2007) exposed the relatively O ₃ -resistant Brown Norway
7	rat strain to 1 ppm O ₃ after sensitizing and challenging with OVA. Rats were exposed for
8	2 days, and airway inflammation was assessed one day later. Filtered air for controls
9	contained less than 0.02 ppm O ₃ . Histopathology indicated O ₃ induced site-specific lung
10	lesions in the centriacinar regions, characterized by wall thickening partly due to
11	inflammatory cells influx. BAL neutrophils were elevated by O_3 in allergic rats, and
12	modestly increased in non-allergic animals (not significant). A slight (but not significant)
13	increase in macrophages was observed, but eosinophil numbers were not affected by O ₃ .
14	Soluble mediators of inflammation (Cys-LT, MCP-1, and IL-6) were elevated by O_3 in
15	allergic animals but not non-allergic rats. Treatment with γT , which neutralizes oxidized
16	lipid radicals and protects lipids and proteins from nitrosative damage, did not alter the
17	morphologic character or severity of the centriacinar lesions caused by O ₃ , nor did it
18	reduce neutrophil influx. It did, however, significantly reduce O ₃ -induced soluble
19	inflammatory mediators in allergic rats. The effects of O ₃ in animal models of allergic
20	asthma are discussed in Section $6.2.6$.
21	In summary, a large number of toxicology studies have demonstrated that acute exposure
22	

21In summary, a large number of toxicology studies have demonstrated that acute exposure22to O_3 produces injury and inflammation in the mammalian lung, supporting the23observations in controlled human exposure studies (Section 6.2.3.1). These acute24changes, both in inflammation and morphology, provide a limited amount of evidence for25long term sequelae of exposure to O_3 . Related alterations resulting from long term26exposure, such as fibrotic changes, are discussed in Chapter 7.

6.2.4 Respiratory Symptoms and Medication Use

27	Controlled human exposure and toxicological studies have described modes of action
28	through which short-term O ₃ exposure may increase respiratory symptoms by
29	demonstrating O_3 -induced airway hyperresponsiveness (Section <u>6.2.2</u>) and pulmonary
30	inflammation (Section $6.2.3.1$ and Section $6.2.3.3$). Epidemiologic studies have not
31	widely examined associations between ambient O3 concentrations and airway
32	hyperresponsiveness but have found O3-associated increases in pulmonary inflammation
33	and oxidative stress (Section $6.3.2.2$). In addition to lung function decrements, controlled
34	human exposure studies clearly indicate O3-induced increases in respiratory symptoms
35	including pain on deep inspiration, shortness of breath, and cough. This evidence is

- 1 detailed in Section 6.2.1.1; however, salient observations include an increase in 2 respiratory symptoms with increasing concentration and duration of O_3 exposure and 3 activity level of exposed subjects (McDonnell et al., 1999b). Further, increases in total 4 subjective respiratory symptoms have been reported following 5.6 and 6.6 hours of 5 exposure to 60 ppb O_3 relative to baseline (Adams, 2006a). At 70 ppb, Schelegle et al. 6 (2009) observed a statistically significant O₃-induced FEV₁ decrement of 6.1% at 6.6 7 hours and a significant increase in total subjective symptoms at 5.6 and 6.6 hours. The 8 findings for O₃-induced respiratory symptoms in controlled human exposure studies and 9 the evidence integrated across disciplines describing underlying modes of action provide 10 biological plausibility for epidemiologic associations observed between short-term increases in ambient O₃ concentration and increases in respiratory symptoms. 11
- 12 In epidemiologic studies, respiratory symptom data typically are collected by having 13 subjects (or their parents) record symptoms and medication use in a diary without direct 14 supervision by study staff. Several limitations of symptom reports are well recognized: 15 recall error if not recorded daily, differences among subjects in the interpretation of 16 symptoms, differential reporting by subjects with and without asthma, and occurrence in 17 a smaller percentage of the population compared with changes in lung function and 18 biological markers of pulmonary inflammation. Nonetheless, symptom diaries remain a 19 convenient tool to collect individual-level data from a large number of subjects and allow 20 modeling of associations between daily changes in O_3 concentration and daily changes in 21 respiratory morbidity. Importantly, most of the limitations described above are sources of 22 random measurement error that can bias effect estimates to the null or increase the 23 uncertainty around effect estimates. Furthermore, because respiratory symptoms are 24 associated with limitations in activity and function and are the primary reason for using 25 medication and seeking medical care, the evidence is directly coherent with the consistent 26 associations observed between increases in ambient O₃ concentration and increases in 27 asthma ED visits (Section 6.2.7.3).

28 Most studies were conducted in individuals with asthma, and as was concluded in the 29 2006 O₃ AOCD (U.S. EPA, 2006b, 1996a), the collective body of epidemiologic 30 evidence indicates that short-term increases in ambient O₃ concentrations are associated 31 with increases in respiratory symptoms in children with asthma. Studies also found 32 O₃-associated increases in the use of asthma medication in children. In a smaller body of 33 studies, increases in ambient O₃ concentration were associated with increases in 34 respiratory symptoms in adults with asthma. Ozone-associated increases in respiratory 35 symptoms in healthy populations were not as clearly indicated.

6.2.4.1 Children with Asthma

Respiratory Symptoms

1	<u>Table 6-19</u> presents the locations, time periods, and ambient O_3 concentrations for studies
2	examining respiratory symptoms and medication use in children with asthma. The
3	evidence supporting associations between short-term increases in ambient O ₃
4	concentration and increases in respiratory symptoms in children with asthma is derived
5	mostly from examination of 1-h max, 8-h max, or 8-h avg O_3 concentrations and strong
6	findings from a large body of single-region or single-city studies (Figure 6-11 and
7	Table 6-20). The few available U.S. multicity studies produced less consistent
8	associations.
9	Similar to lung function, associations with respiratory symptoms in children with asthma
10	were found with ambient O ₃ concentrations assigned to subjects using various methods
11	with potentially different degrees of exposure measurement error. As was discussed for
12	lung function, methods included measurement of O3 on site of and at the time of outdoor
13	activity (Thurston et al., 1997), which is associated with higher ambient-personal O_3
14	correlations and ratios (Section $4.3.3$); O ₃ concentrations measured at sites within 5 km of
15	subjects' home or school (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997;
16	<u>1996</u>); O_3 measured at a single city site (<u>Gielen et al., 1997</u>); and O_3 concentrations
17	averaged across multiple sites (Gent et al., 2003; Mortimer et al., 2002). In analyses with
18	O_3 averaged across multiple sites, which were restricted to warm seasons, O_3
19	concentrations within the region were temporally correlated as indicated by high
20	statewide correlations [median $r = 0.83$ in <u>Gent et al. (2003</u>)] or similar odds ratios for O_3
21	averaged across all within-city monitors and that averaged from the three closest sites
22	(Mortimer et al., 2002). In these panel studies, the averaged ambient concentrations may
23	have well represented the temporal variability in subjects' ambient O ₃ exposures.

Table 6-19Mean and upper percentile ozone concentrations in epidemiologic
studies of respiratory symptoms, medication use, and activity
levels in children with asthma.

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Thurston et</u> <u>al. (1997</u>)	CT River Valley, CT	June 1991- 1993	1-h max	83.6ª	Max: 160 ^ª
<u>Escamilla-</u> <u>Nuñez et al.</u> (2008)	Mexico City, Mexico	July-March 2003-2005	1-h max	86.5	NR
<u>2006</u>)	Mexico City, Mexico	October 1998-	8-h max	69	Max: 184
		April 2000	1-h max	102	Max: 309
<u>1997</u>)	Southern Mexico City, Mexico	April-July 1991; November 1991-February 1992	1-h max	196	Max: 390
<u>Romieu et al.</u> (1996)	Northern Mexico City, Mexico	April-July 1991; November 1991-February 1992	1-h max	190	Max: 370
<u>Gent et al.</u> (2003)	CT, southern MA	April- September 2001	8-h rolling avg 1-h max	51.3, 50.0 (median) 58.6, 55.5 (median)	Max: 99.6 Max: 125.5
<u>Mortimer et</u> <u>al. (2002);</u> (<u>2000</u>)	Bronx, East Harlem, NY; Baltimore, MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St. Louis, MO; (NCICAS)	June-August 1993	8-h avg (10 a.m 6 p.m.)	48	NR
<u>Gielen et al.</u> (1997)	Amsterdam, Netherlands	April-July 1995	8-h max	34.2 ^b	Max: 56.5 [♭]
Delfino et al.	Los Angeles, CA	November	8-h max	17.1	90th: 26.1, Max: 37
<u>(2003</u>)		1999-January 2000	1-h max	25.4	90th: 38.0, Max: 52
Rabinovitch et al. (2004)	Denver, CO	November- March 1999- 2002	1-h max	28.2	75th: 60, Max: 70.0
Schildcrout et al. (2006)	Albuquerque, NM; Baltimore, MD; Boston, MA; Denver, CO; San Diego, CA; Seattle, WA; St. Louis, MO; Toronto, ON, Canada (CAMP)	May- September 1994-1995	1-h max	Range in medians across cities: 43.0- 65.8	Range in 90th across cities: 61.5-94.7
<u>Jalaludin et</u> <u>al. (2004</u>)	Sydney, Australia	February- December 1994	15-h avg (6 a.m9 p.m.)	12	Max: 43

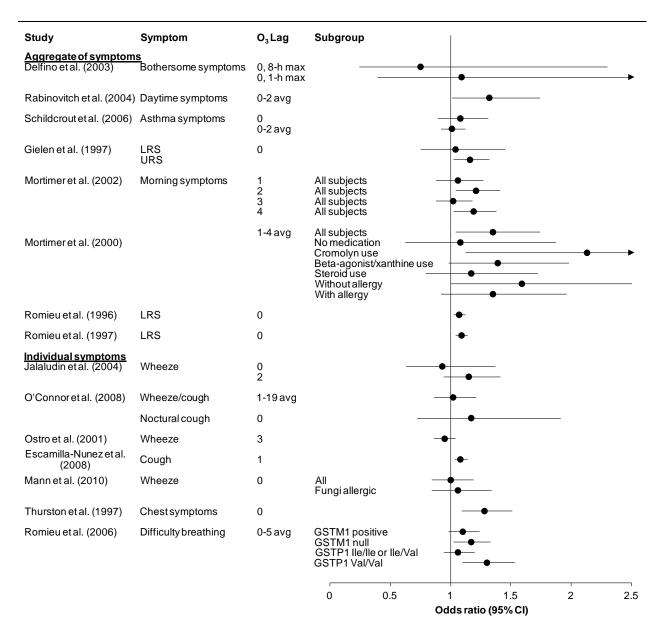
Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>O'Connor et</u> <u>al. (2008</u>)	Boston, MA; Bronx, Manhattan NY; Chicago, IL; Dallas, TX, Seattle, WA; Tucson, AZ (ICAS)	August 1998- July 2001	24-h avg	NR	NR
Ostro et al.	Los Angeles, CA	August-	1-h max	Los Angeles: 59.5	Max: 130
<u>(2001</u>)		October 1993		Pasadena: 95.8	Max: 220
<u>Mann et al.</u> (2010)	Fresno/Clovis, California	Winter- Summer 2000- 2005	8-h max	49.4 (median)	75th: 69.5, Max: 120.0
<u>Just et al.</u> (2002)	Paris, France	April-June 1996	24-h avg	30.0 ^b	Max: 61.7 ^b

* Note: Studies presented in order of first appearance in the text of this section.

NCICAS = National Cooperative Inner-City Asthma Study, NR = Not Reported, ICAS = Inner City Asthma Study, CAMP = Childhood Asthma Management Program.

^aMeasured on site of subjects' outdoor activity.

^bConcentrations converted from µg/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).



Note: Results are presented first for aggregate indices of symptoms then for individual symptoms. Within each category, results generally are organized in order of increasing mean ambient O_3 concentration. LRS = lower respiratory symptoms, URS = upper respiratory symptoms. Odds ratios are from single-pollutant models and are standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 8-h max (or 8-h avg or 15-h avg), and 24-h avg O_3 concentrations, respectively.

Figure 6-11 Associations between ambient ozone concentrations and respiratory symptoms in children with asthma.

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Symptom	Subgroup	Standardized OR (95% CI) ^a
Studies exa	mining aggregates of symptoms	5				
Delfino et al.	Los Angeles, CA	8-h max	0	Bothersome		0.75 (0.24, 2.30)
<u>(2003</u>)	22 children with asthma, ages 10- 16 yr	1-h max		symptoms		1.09 (0.39, 3.03)
Rabinovitch	Denver, CO	1-h max	0-2	Daytime		1.32 (1.01, 1.74)
<u>et al. (2004</u>)	86 children with asthma, ages 6- 12 yr		avg	symptoms		
Schildcrout et	Albuquerque, NM; Baltimore, MD;	1-h max	0	Asthma		1.08 (0.89, 1.31)
<u>al. (2006</u>)	Boston, MA; Denver, CO; San Diego, CA; Seattle, WA; St. Louis, MO; Toronto, ON, Canada		0-2 avg	symptoms		1.01 (0.92, 1.12)
	990 children with asthma, ages 5- 12 yr					
Gielen et al.	Amsterdam, Netherlands	8-h max	0	LRS		1.04 (0.75, 1.45)
<u>(1997</u>)	61 children with asthma, ages 7- 13 yr			URS		1.16 (1.02, 1.32)
(<u>2002</u>);	Bronx, East Harlem, NY; Baltimore,	8-h avg	1	Morning	All subjects	1.06 (0.88, 1.27)
Mortimer et al. (2000)	MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St.	(10 a.m	2	symptoms	All subjects	1.21 (1.04, 1.41)
<u>ai. (2000</u>)	Louis, MO	6 p.m.)	3		All subjects	1.02 (0.88, 1.18)
	846 children with asthma, ages 4-		4		All subjects	1.19 (1.02, 1.38)
	9 yr		1-4		All subjects	1.35 (1.04, 1.74)
			avg		No medication	1.08 (0.62, 1.87
					use	2.13 (1.12, 4.04)
					Cromolyn use	1.39 (0.98, 1.98)
					β- agonist/xanthine	1.17 (0.79, 1.72)
					USE	1.59 (1.00, 2.52)
					Steroid use	1.35 (0.92, 1.96)
					Without allergy	
					With allergy	
<u>Romieu et al.</u> (1996)	northern Mexico City, Mexico	1-h max	0	LRS		1.07 (1.02, 1.12)
(1996)	71 children with asthma, ages 5- 7 yr					
Romieu et al.	southern Mexico City, Mexico	1-h max	0	LRS		1.09 (1.04, 1.14)
<u>(1997</u>)	65 children with asthma, ages 5- 13 yr					
Studies exa	mining individual symptoms					
Jalaludin et	Sydney, Australia	15-h avg	0	Wheeze		0.93 (0.63, 1.37)
<u>al. (2004</u>)	125 children with asthma, mean age 9.6 yr	(6 a.m 9 p.m.)	2			1.15 (0.94, 1.41)
<u>O'Connor et</u> <u>al. (2008</u>)	Boston, MA; Bronx, Manhattan NY; Chicago, IL; Dallas, TX, Seattle, WA; Tucson, AZ	24-h avg	1-19 avg	Wheeze/cough		1.02 (0.86, 1.21)
	861 children with asthma, mean (SD) age 7.7 (2.0) yr					

Table 6-20Additional characteristics and quantitative data for studies
presented in Figure 6-11.

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Symptom	Subgroup	Standardized OR (95% CI) ^a
<u>Just et al.</u> (2002)	Paris, France 82 children with asthma, mean (SD) age 10.9 (2.5) yr	24-h avg	0	Nocturnal cough incidence		1.17 (0.72, 1.91)
<u>Ostro et al.</u> (2001)	Los Angeles, CA 138 children with asthma, ages 6- 13 yr	1-h max	3	Wheeze		0.95 (0.86, 1.04)
<u>Escamilla-</u> <u>Nuñez et al.</u> (2008)	Mexico City, Mexico 147 children with asthma, mean age 9.6 yr	1-h max	1	Wheeze		1.08 (1.03, 1.14)
<u>Mann et al.</u> (2010)	Fresno/Clovia, California 280 children with asthma, ages 6- 11 yr	8-h max	0	Wheeze	All Fungi allergic	1.00 (0.84, 1.19) 1.06 (0.84, 1.34)
<u>Thurston et</u> <u>al. (1997</u>)	CT River Valley, CT 166 children with asthma, ages 7- 13 yr	1-h max	0	Chest symptoms		1.28 (1.09, 1.51)
<u>Romieu et al.</u> (2006)	Mexico City, Mexico 151 children with asthma, mean age 9 yr	1-h max	0-5 avg	Difficulty breathing	GSTM1 positive GSTM1 null GSTP1 lle/lle or lle/Val GSTP1 Val/Val	1.10 (0.98, 1.24) 1.17 (1.02, 1.33) 1.06 (0.94, 1.20) 1.30 (1.10, 1.53)
<u>Gent et al.</u> (2003) ^b	CT, southern MA 130 children with asthma on maintenance medication	1-h max	0	Wheeze	$O_3 < 43.2 \text{ ppb}$ $O_3 43.2-51.5 \text{ ppb}$ $O_3 51.6-58.8 \text{ ppb}$ $O_3 58.9-72.6 \text{ ppb}$ $O_3 \ge 72.7 \text{ ppb}$	1.00 (reference) 1.04 (0.89, 1.21) 1.16 (1.00, 1.35) 1.16 (1.00, 1.35) 1.22 (0.97, 1.53)
				Chest tightness	$O_3 < 43.2 \text{ ppb}$ $O_3 43.2-51.5 \text{ ppb}$ $O_3 51.6-58.8 \text{ ppb}$ $O_3 58.9-72.6 \text{ ppb}$ $O_3 \ge 72.7 \text{ ppb}$	1.00 (reference) 1.11 (0.91, 1.36) 1.01 (0.83, 1.23) 1.16 (0.97, 1.39) 1.31 (0.97, 1.77)

*Includes studies for <u>Figure 6-11</u>, plus others.

LRS = Lower respiratory symptoms, URS = Upper respiratory symptoms.

^aEffect estimates are standardized to a 40, 30, and 20 ppb increase for 1-h max, 8-h max (or 8-h avg or 15-h avg), and 24-h avg O₃, respectively.

^bResults not included in <u>Figure 6-11</u> because results presented per quintile of ambient O₃ concentration.

1	Among U.S. multicity studies of children with asthma, each of which examined a
2	different O3 averaging time, O3 was not consistently associated with increases in
3	respiratory symptoms (O'Connor et al., 2008; Schildcrout et al., 2006; Mortimer et al.,
4	<u>2002</u>). In the NCICAS cohort (described in Section <u>$6.2.1.2$</u>), increases in most evaluated
5	lags of O_3 (1 to 4 and 1-4 avg) were associated with increases in asthma symptoms. A
6	30-ppb increase in lag 1-4 avg, of 8-h avg (10 a.m6 p.m.), O ₃ was associated with an
7	increase in morning asthma symptoms with an OR of 1.35 (95% CI: 1.04, 1.69)
8	(Mortimer et al., 2002). The OR was similar in an analysis restricted to O_3 concentrations
9	<80 ppb. Associations were similarly strong for lags 2 and 4 of O_3 but weaker for lags 1

1	and 3 (Figure 6-11 and Table 6-20). In the ICAS cohort (described in Section $6.2.1.2$),
2	associations of 19-day avg of 24-h avg O_3 with wheeze and nighttime asthma were
3	positive and negative, respectively (O'Connor et al., 2008). NCICAS collected symptom
4	data daily (Mortimer et al., 2002; 2000), whereas in ICAS, every 2 months, parents
5	reported the number of days with symptoms over the previous 2 weeks (O'Connor et al.,
6	2008). Thus, ICAS was precluded from examining associations with single-day O_3
7	concentrations and shorter lag periods.
8	Like NCICAS, the U.S. multicity Childhood Asthma Management Program (CAMP,
9	with cities in common with NCICAS and ICAS, Table 6-19) collected daily symptom
10	data, analyzed data collected between May and September, and evaluated multiple lags of
11	O_3 (<u>Schildcrout et al., 2006</u>). However, associations in CAMP were weaker for all
12	evaluated lags of O ₃ . In meta-analyses that combined city-specific estimates, a 40-ppb
13	increase in lag 0 of 1-h max O ₃ was associated with asthma symptoms with an OR of
14	1.08 (95% CI: 0.89, 1.31). Odds ratios for lags 1 and 2 and the 3-day sum of O_3 were
15	between 1.0 and 1.03. In this study, data available from an average of 12 subjects per day
16	per city were used to produce city-specific ORs. These city-specific ORs then were
17	combined in meta-analyses to produce study-wide ORs. Because O ₃ analyses were
18	restricted to warm seasons, there likely was less power to detect associations with O ₃ than
19	with other pollutants, which were analyzed using year-round data.
20	Several longitudinal studies conducted in different cohorts of children with asthma in
20 21	Several longitudinal studies conducted in different cohorts of children with asthma in Mexico City, Mexico examined and found increases in respiratory symptoms in
21	Mexico City, Mexico examined and found increases in respiratory symptoms in
21 22	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (<u>Escamilla-Nuñez et al., 2008</u> ; <u>Romieu et al.</u> ,
21 22 23	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (<u>Escamilla-Nuñez et al., 2008</u> ; <u>Romieu et al., 2006</u> ; <u>1997</u> ; <u>1996</u>). <u>1997</u>); (<u>1996</u>) <u>Romieu et al. (1997</u>); (<u>1996</u>) found larger increases in
21 22 23 24	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (<u>Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996</u>).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O ₃ at lag 0, than at lag 1 or 2. Recent
21 22 23 24 25	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O ₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of
21 22 23 24 25 26	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O ₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O ₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that
21 22 23 24 25 26 27	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O ₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O ₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O ₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O ₃ concentrations with respiratory symptoms
21 22 23 24 25 26 27 28	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O_3 concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O_3 at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O_3 concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O_3 concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5
21 22 23 24 25 26 27 28 29	Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O_3 concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O_3 at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O_3 concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O_3 concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg).
21 22 23 24 25 26 27 28 29 30	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to
21 22 23 24 25 26 27 28 29 30 31	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to ambient O₃ exposure but did not produce conclusive evidence. Larger O₃-associated
21 22 23 24 25 26 27 28 29 30 31 32	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to ambient O₃ exposure but did not produce conclusive evidence. Larger O₃-associated (8-h avg [10 a.m6 p.m.] or 8-h max) increases in symptoms were found in children
21 22 23 24 25 26 27 28 29 30 31 32 33	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to ambient O₃ exposure but did not produce conclusive evidence. Larger O₃-associated (8-h avg [10 a.m6 p.m.] or 8-h max) increases in symptoms were found in children taking asthma medication, although the specific medications examined differed between
21 22 23 24 25 26 27 28 29 30 31 32 33 34	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to ambient O₃ exposure but did not produce conclusive evidence. Larger O₃-associated (8-h avg [10 a.m6 p.m.] or 8-h max) increases in symptoms were found in children taking asthma medication, although the specific medications examined differed between studies. As with results for PEF, in the NCICAS multicity cohort, O₃-associated increases
21 22 23 24 25 26 27 28 29 30 31 32 33 34 35	 Mexico City, Mexico examined and found increases in respiratory symptoms in association with 1-h max O₃ concentrations (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996).1997); (1996) Romieu et al. (1997); (1996) found larger increases in symptoms in association with increases in 1-h max O₃ at lag 0, than at lag 1 or 2. Recent studies expanded on earlier evidence by indicating associations with multiday averages of O₃ concentrations. Romieu et al. (2006) and Escamilla-Nuñez et al. (2008) found that ORs for associations of ambient 1-h max O₃ concentrations with respiratory symptoms and medication use increased as the number of averaging days increased (up to lag 0-5 avg). Studies of children with asthma examined factors that may modify symptom responses to ambient O₃ exposure but did not produce conclusive evidence. Larger O₃-associated (8-h avg [10 a.m6 p.m.] or 8-h max) increases in symptoms were found in children taking asthma medication, although the specific medications examined differed between studies. As with results for PEF, in the NCICAS multicity cohort, O₃-associated increases in morning symptoms were larger in children taking cromolyn (used to treat asthma with

and Table 6-20) (Mortimer et al., 2000). Among children with asthma in Southern New
England, O3-associated increases in symptoms were limited mostly to children taking
steroids, cromolyn, or leukotriene inhibitors for maintenance (Gent et al., 2003).

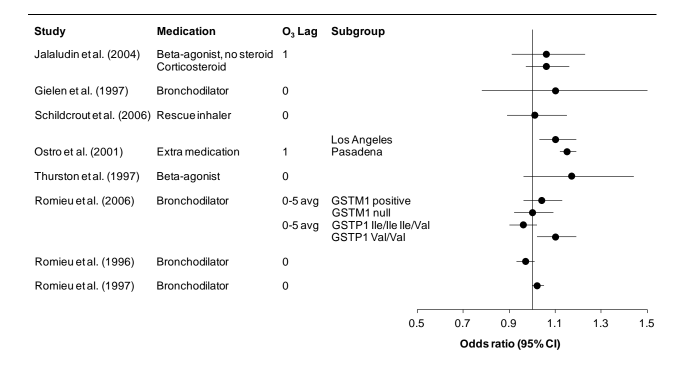
- 4 In most studies of children with asthma, a majority of subjects (52 to 100%) had atopy as 5 determined by sensitization to any examined allergen. While studies found O₃-associated 6 increases in pulmonary inflammation in children with atopy (Section 6.2.3.2) and in 7 animal models of allergy (Section 6.2.3.3), studies did not indicate that the risk of 8 O₃-associated respiratory symptoms differed in children with asthma with and without 9 atopy. In NCICAS, Mortimer et al. (2000) found that an increase in 8-h avg (10 a.m.-6 10 p.m.) O_3 was associated with a similar increased incidence of asthma symptoms among 11 the 79% of subjects with atopy and the 21% of subjects without atopy (Figure 6-11 and 12 Table 6-20). Odds ratios for O_3 did not differ by residential allergen levels. Among 13 children with asthma in Fresno, CA, most associations of single- and multiday lags of 14 8-h max O_3 concentrations (0-14 days) with wheeze were near or below 1.0 among all 15 subjects. Among the various O_3 lags examined, increases in O_3 were not consistently 16 associated with increases in wheeze in subjects with cat or fungi allergy either (Mann et 17 al., 2010).
- 18 Romieu et al. (2006) found differences in O_3 -associated respiratory symptoms by genetic 19 variants in GST enzymes, particularly, GSTP1 and less so for GSTM1. Compared with 20 GSTP1 Ile/Ile or Ile/Val subjects, larger effects were estimated for GSTP1 Val/Val 21 subjects (Figure 6-11 and Table 6-20). The largest OR was found for difficulty breathing 22 in children with asthma who had both GSTM1 null and GSTP1 Val/Val genotypes (OR: 23 1.49 [95% CI: 1.14, 1.93] per 30-ppb increase in lag 0-5 avg of 8-h max O_3). While these 24 results are consistent with those described for antioxidant capacity modifying 25 O_3 -associated changes in lung function (Section 6.2.1.2) and pulmonary inflammation 26 [Section 6.2.3.2 for results in the same cohort (Sienra-Monge et al., 2004)], it is important 27 to note that effect modification by GSTP1 variants has not been consistent. (Romieu et 28 al., 2006) found an O_3 -associated decrease in FEV₁ only in children with GSTP1 Ile/Ile 29 or Ile/Val genotype. Among children in southern California, GSTP1 Ile/Ile was 30 associated with greater risk of asthma onset (Section 7.2.1). Asthma prevalence has not 31 been consistently associated with a particular GSTP1 genotype either (Tamer et al., 2004; 32 Mapp et al., 2002; Hemmingsen et al., 2001).

Asthma Medication Use

Although recent studies contributed mixed evidence, the collective body of evidence
 supports associations between increases in ambient O₃ concentration and increased
 asthma medication use in children (Figure 6-12 and Table 6-21). Most studies examined

1 2 3

1	and found associations with lags 0 or 1 of 1-h max O_3 concentrations; however,
2	associations also were found for multiday average O3 concentrations (lag 0-5 avg in
3	Romieu et al. (2006) and lags 0-2 avg and 0-4 avg in Just et al. (2002). Within several
4	studies, associations were consistent between respiratory symptoms and asthma
5	medication use (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; Schildcrout et al.,
6	2006; Jalaludin et al., 2004; Romieu et al., 1997; Thurston et al., 1997). As an exception,
7	<u>Romieu et al. (1996</u>) found that O_3 was associated with an increase in respiratory
8	symptoms but not bronchodilator use, and Rabinovitch et al. (2004) indicated statistically
9	significant associations with symptoms but not bronchodilator use (OR not reported). A
10	few studies found higher odds of O3-associated increases in asthma medication use than
11	in respiratory symptoms (Just et al., 2002; Ostro et al., 2001).



Note: CS = corticosteroid. Results generally are presented in order of increasing mean ambient O_3 concentration. Odds ratios are from single-pollutant models and are standardized to a 40- ppb for 1-h max O_3 and a 30-ppb increase for 8-h max or 15-h avg O_3 .

Figure 6-12 Associations between ambient ozone concentrations and asthma medication use.

Table 6-21Additional characteristics and quantitative data for studies
presented in Figure 6-12.

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Medication	Subgroup	Standardized OR (95% CI) ^a
<u>Jalaludin et</u> <u>al. (2004</u>)	Sydney, Australia 125 children with asthma,	15-h avg (6 a.m9 p.m.)	1	Beta-agonist, no corticosteroid		1.06 (0.91, 1.23)
	mean age 9.6 yr			Inhaled corticosteroid		1.06 (0.97, 1.16)
Gielen et al.	Amsterdam, Netherlands	8-h max	0	Bronchodilator		1.10 (0.78, 1.55)
<u>(1997</u>)	61 children with asthma, ages 7-13 yr					
Schildcrout et al. (2006)	Albuquerque, NM; Baltimore, MD; Boston, MA; Denver, CO; San Diego, CA; Seattle, WA; St. Louis, MO; Toronto, ON, Canada	1-h max	0	Rescue inhaler		1.01 (0.89, 1.15)
	990 children with asthma, ages 5-12 yr					
Ostro et al.	Los Angeles, CA	1-h max	1	Any extra	Pasadena	1.15 (1.12, 1.19)
<u>(2001</u>)	138 children with moderate/severe asthma, ages 6-13 yr			medication	Los Angeles	1.10 (1.03, 1.19)
<u>Thurston et</u> <u>al. (1997</u>)	CT River Valley, CT 166 children with asthma, ages 7-13 yr	1-h max	0	Beta-agonist		1.17 (0.96, 1.44)
<u>Romieu et</u> <u>al. (2006</u>)	Mexico City, Mexico 151 children with asthma,	1-h max	0-5 avg	Bronchodilator	GSTM1 positive	1.04 (0.96, 1.13)
	mean age 9 yr				GSTM1 null	1.00 (0.92, 1.09)
					GSTP1 lle/lle or lle/Val	0.96 (0.90, 1.02)
					GSTP1 Val/Val	1.10 (1.02, 1.19)
<u>Romieu et</u> <u>al. (1996</u>)	northern Mexico City, Mexico	1-h max	0	Bronchodilator		0.97 (0.93, 1.01)
	71 children with asthma, ages 5-7 yr					
<u>Romieu et</u> <u>al. (1997</u>)	southern Mexico City, Mexico 65 children with asthma, ages 5-13 yr	1-h max	0	Bronchodilator		1.02 (1.00, 1.05)

Study*	Location/Population	O₃ Averaging Time	O₃ Lag	Medication	Subgroup	Standardized OR (95% CI) ^a
<u>Just et al.</u> (2002) ^b	Paris, France 82 Children with asthma, mean (SD) age 10.9 (2.5) yr	24-h avg	0	Beta-agonist, no steroid		3.95 (1.22, 12.9)
Gent et al.	CT, southern MA	1-h max	0	Bronchodilator	O ₃ <43.2 ppb	1.00 (reference)
<u>(2003</u>)⁵	130 children with asthma on maintenance medication				O ₃ 43.2- 51.5 ppb	1.00 (0.96, 1.05)
					O₃ 51.6- 58.8 ppb	1.04 (1.00, 1.09)
					O₃ 58.9- 72.6 ppb	1.02 (0.98, 1.07)
					O ₃ ≥ 72.7 ppb	1.05 (0.97, 1.13)

*Includes studies in Figure 6-12, plus others.

^aEffect estimates are standardized to a 40-ppb increase for 1-h max O_3 , a 30-ppb increase for 8-h max or 15-h avg O_3 , and a 20-ppb increase for 24-h avg O_3 .

^bResults not included in Figure 6-12. Results from Just et al. (2002) were out of range of other estimates, and results from Gent et al. (2003) were presented per quintile of ambient O_3 concentration.

Changes in Activity

2associated diminished activity in children with asthma (O'Connor et al., 2008; Delfino et3al., 2003). These studies examined different O3 averaging times and lags. In the multicity4ICAS cohort, O'Connor et al. (2008) found that a 20-ppb increase in lag 1-19 avg of524-hour O3 was associated with a 10% lower odds (95% CI: -26, 10) of slow play. In a6small (n = 22) panel study conducted in children with asthma in Los Angeles CA,7Delfino et al. (2003) found that a 40-ppb increase in lag 0 of 1-h max O3 was associated8with an increase in symptoms that interfered with daily activity with an OR of 7.419(95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children10with asthma in association with increases in ambient O3 concentration with long lag11periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O316exposure and the potential for residual seasonal confounding with examination of long	1	While investigation has been limited, evidence does not consistently demonstrate O_3 -
4ICAS cohort, O'Connor et al. (2008) found that a 20-ppb increase in lag 1-19 avg of524-hour O3 was associated with a 10% lower odds (95% CI: -26, 10) of slow play. In a6small (n = 22) panel study conducted in children with asthma in Los Angeles CA,7Delfino et al. (2003) found that a 40-ppb increase in lag 0 of 1-h max O3 was associated8with an increase in symptoms that interfered with daily activity with an OR of 7.419(95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children10with asthma in association with increases in ambient O3 concentration with long lag11periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O3	2	associated diminished activity in children with asthma (O'Connor et al., 2008; Delfino et
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7Delfino et al. (2003) found that a 40-ppb increase in lag 0 of 1-h max O3 was associated8with an increase in symptoms that interfered with daily activity with an OR of 7.419(95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children10with asthma in association with increases in ambient O3 concentration with long lag11periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O3	5	24-hour O_3 was associated with a 10% lower odds (95% CI: -26, 10) of slow play. In a
 with an increase in symptoms that interfered with daily activity with an OR of 7.41 (95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children with asthma in association with increases in ambient O₃ concentration with long lag periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008) examined absences for any reason, Gilliland et al. (2001) found associations with absences for respiratory illnesses. Despite this evidence, several limitations are notable, including the lack of a well-characterized mode of action for long lag periods of O₃ 	6	small ($n = 22$) panel study conducted in children with asthma in Los Angeles CA,
 9 (95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children 10 with asthma in association with increases in ambient O₃ concentration with long lag 11 periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland 12 et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008) 13 examined absences for any reason, Gilliland et al. (2001) found associations with 14 absences for respiratory illnesses. Despite this evidence, several limitations are notable, 15 including the lack of a well-characterized mode of action for long lag periods of O₃ 	7	Delfino et al. (2003) found that a 40-ppb increase in lag 0 of 1-h max O ₃ was associated
10with asthma in association with increases in ambient O3 concentration with long lag11periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O3	8	with an increase in symptoms that interfered with daily activity with an OR of 7.41
11periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O3	9	(95% CI: 1.18, 43.2). Several studies reported increases in school absenteeism in children
12et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)13examined absences for any reason, Gilliland et al. (2001) found associations with14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O3	10	with asthma in association with increases in ambient O_3 concentration with long lag
 examined absences for any reason, <u>Gilliland et al. (2001</u>) found associations with absences for respiratory illnesses. Despite this evidence, several limitations are notable, including the lack of a well-characterized mode of action for long lag periods of O₃ 	11	periods (14-day and 30-day distributed lags, 19-day avg) (O'Connor et al., 2008; Gilliland
14absences for respiratory illnesses. Despite this evidence, several limitations are notable,15including the lack of a well-characterized mode of action for long lag periods of O_3	12	et al., 2001; Chen et al., 2000). Whereas Chen et al. (2000) and O'Connor et al. (2008)
15 including the lack of a well-characterized mode of action for long lag periods of O_3	13	examined absences for any reason, Gilliland et al. (2001) found associations with
	14	absences for respiratory illnesses. Despite this evidence, several limitations are notable,
16 exposure and the potential for residual seasonal confounding with examination of long	15	including the lack of a well-characterized mode of action for long lag periods of O_3
	16	exposure and the potential for residual seasonal confounding with examination of long
17 lag periods. In analyses of single-day lags, <u>Gilliland et al. (2001</u>) found associations with	17	lag periods. In analyses of single-day lags, Gilliland et al. (2001) found associations with
18 O ₃ lagged 1 to 5 days, indicating respiratory absences may be affected by O ₃ exposures	18	O_3 lagged 1 to 5 days, indicating respiratory absences may be affected by O_3 exposures
19 with shorter lag periods.	19	with shorter lag periods.

6.2.4.2 Adults with Respiratory Disease

1	Within a small body of studies, several found that increases in ambient O_3 concentration
2	(8-hour or 1-h max) were associated with increases in respiratory symptoms in adults
3	with asthma (Khatri et al., 2009; Feo Brito et al., 2007; Ross et al., 2002). Details from
4	studies of respiratory symptoms in adults with respiratory disease regarding location,
5	time period, and ambient O_3 concentrations are presented in <u>Table 6-22</u> . These studies
6	used different exposure assessment methods: concentrations averaged from sites closest
7	to subjects' location each hour (Khatri et al., 2009) or concentrations measured at one
8	(Ross et al., 2002) or multiple (Feo Brito et al., 2007) city sites. Park et al. (2005a) found
9	inconsistent associations for 24-h avg O_3 measured at 10 city sites among the various
10	symptoms and medication use examined in adults with asthma in Korea during a period
11	of dust storms. In a study of adults with COPD in London, England, increases in lag 1 of
12	8-h max O_3 (at a single city site) were associated with higher odds of dyspnea and sputum
13	changes but lower odds of nasal discharge, wheeze, or upper respiratory symptoms
14	(<u>Peacock et al., 2011</u>).

Table 6-22Mean and upper percentile ozone concentrations in epidemiologic
studies of respiratory symptoms and medication use in adults with
respiratory disease .

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Khatri et al. (2009</u>)	Atlanta, GA	May-September 2003, 2005, 2006	8-h max	61 ^ª	75th: 74 ^ª
<u>Feo Brito et al.</u> (2007)	Ciudad Real and Puertollano, Spain	May-June 2000- 2001	1-h max	65.9 (Ciudad Real) ^b 56.8 (Puertollano) ^b	Max: 101.5 ^b (Ciudad Real); 138.2 ^b (Puertollano)
Eiswerth et al. (2005)	Glendora, CA	October-November 1983	1-h max	NR	NR
Ross et al. (2002)	East Moline, IL	April-October 1994	8-h avg	41.5	Max: 78.3
<u>Peacock et al.</u> (2011)	London, England	All-year 1995-1997	8-h max	15.5	Autumn/Winter Max: 32 Spring/Summer Max: 74
Park et al. (2005a)	Incheon, Korea	March-June 2002	24-h avg	Dust event days: 23.6 Control days: 25.1	NR
Wiwatanadate and Liwsrisakun (2011)	Chiang Mai, Thailand	August 2005-June 2006	24-h avg	17.5	90th: 26.82, Max: 34.65

* Note: Studies presented in order of first appearance in the text of this section.

NR = Not Reported

^aIndividual-level estimates were derived based on time spent in the vicinity of various O_3 monitors.

 b Concentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

1	Some studies that included adults with asthma examined populations with a high
2	prevalence of atopy. In a study of children and adults with asthma (at least 53% with
3	atopy), Ross et al. (2002) found that an increase in lag 1-3 avg of 8-h max O_3 was
4	associated with an increase in symptom score and asthma medication use. Feo Brito et al.
5	(2007) followed 137 adults with asthma in two central Spain cities. All subjects had
6	pollen allergy and were examined during pollen season. In Puertollano, O3 concentrations
7	were obtained from four city monitors, and a 40-ppb increase in lag 3 of 1-h max O_3 was
8	associated with a 14.3% increase (95% CI: 3.6, 26.0) in the number of subjects reporting
9	respiratory symptoms, adjusting only for time trend. The association was much weaker in
10	Ciudad Real (2.3% increase [95% CI: -14, 21%] per 40-ppb increase in lag 4 of 1-h max
11	O ₃), a city characterized by lower ambient air pollution levels and a narrower range of
12	ambient O ₃ concentrations as measured at a single site established by investigators.
13	Cross-sectional studies reported ambient O3-associated decreases in activity in adults
14	with asthma; however, due to various limitations in the collective body of evidence, firm
15	conclusions are not warranted. Although conducted over single seasons, studies did not
16	consider confounding by meteorological factors. In a warm season study in Atlanta, GA
17	(described in Section <u>6.2.1.2</u>), <u>Khatri et al. (2009</u>) found that a 30-ppb increase in lag 2 of

1	8-h max O_3 was associated with a 0.69-point decrease (95% CI: -1.28, -0.11) in the
2	Juniper quality of life score, which incorporates indices for symptoms, mood, and activity
3	limitations (7-point scale). In a fall study conducted in the Los Angeles, CA area in
4	individuals with asthma (age 16 years and older), Eiswerth et al. (2005) found that a
5	40-ppb increase in 1-h max O_3 was associated with a 0.24% (95% CI: 0.08, 0.40%) lower
6	probability of indoor activity but higher probability of outdoor activity. The authors
7	acknowledged that their findings were unexpected and may have been influenced by lack
8	of control for potential confounders but interpreted the decrease in indoor activities as
9	rest replacing chores. In contrast with the aforementioned studies, a panel study of
10	individuals with asthma (ages 13-78 years) in Thailand found that a 20-ppb increase in
11	lag 4 of 24-h avg O_3 was associated with a 26% (95% CI: 4, 43) lower odds of symptoms
12	that interfered with activities (Wiwatanadate and Liwsrisakun, 2011).

6.2.4.3 Populations not Restricted to Individuals with Asthma

13	Locations, time periods, and ambient O3 concentrations for studies of symptoms in
14	populations not restricted to individuals with asthma are presented in Table 6-23. Most
15	studies examined children, and in contrast with lung function results (Section 6.2.1.2),
16	short-term increases in ambient O_3 concentration were not consistently associated with
17	increases in respiratory symptoms in children in the general population (Figure 6-13 and
18	Table 6-24). Because examination of adults was limited, conclusions cannot be drawn.

Table 6-23Mean and upper percentile ozone concentrations in epidemiologic
studies of respiratory symptoms in populations not restricted to
individuals with asthma.

Study*	Location	Study Period	O₃ Averaging Time	Mean/Median Concentration (ppb)	Upper Percentile Concentrations (ppb)
<u>Neas et al.</u> (1995)	Uniontown, PA	June-August 1990	12-h avg (8 a.m8 p.m.)	37.2	Max: 87.5
Linn et al.	Rubidoux, Upland,	September-June	24-h avg personal	5	Max: 16
<u>(1996</u>)	Torrence, CA	1992-1994	24-h avg ambient	23	Max: 53
Hoek and	Deurne and	March-July 1989	1-h max	Deurne: 57	Max: 107
<u>Brunekreef</u> (1995)	Enkhuizen, Netherlands			Enkhuizen: 59	Max: 114
Rodriguez et al.	Perth, Australia	All-year, 1996-	24-h avg	28	Max: 74
<u>(2007</u>)		2003	1-h max	33	Max: 95
<u>Moon et al.</u> (2009)	4 cities, South Korea	April-May 2003	8-h avg (10 a.m6 p.m.)	NR	NR
Ward et al.	Birmingham and	January-March,	24-h avg	Winter median: 13.0	Winter Max: 33
<u>(2002</u>)	Sandwell, England	May-July 1997		Summer median: 22.0	Summer Max: 41
Triche et al.	Southwestern VA	June-August	24-h avg	35.2	75th: 40.6, Max: 56.6
<u>(2006</u>)		1995-1996	8-h max	54.5	75th: 64.1, Max: 87.6
			1-h max	60.8	75th: 70.0, Max: 95.0
<u>Gold et al.</u> (1999)	Mexico City, Mexico	January- November 1991	24-h avg	52.0 ^a	Max: 103ª
<u>Apte et al.</u> (2008)	Multiple U.S. cities (NR)	Winter or summer 1994-	Workday avg (8 a.m 5 p.m.)	34.2 ^b	Max: 86.2 ^b
		1998	24-h avg	25.5 ^b	Max: 67.3 ^b

* Note: Studies presented in order of first appearance in the text of this section.

NR = Not Reported.

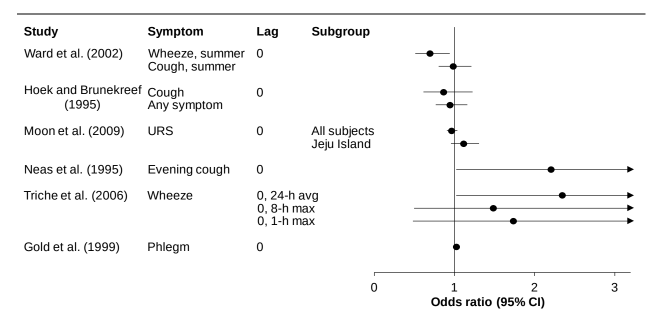
^aMeasured at subject's schools.

^bConcentrations converted from μ g/m³ to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

Children

1	Although evidence of O3-associated increases in respiratory symptoms in children was
2	inconsistent, it did not appear to be attributable to the differences in exposure assessment
3	method among studies [e.g., O ₃ measured at a single site (Linn et al., 1996; Hoek and
4	Brunekreef, 1995), O ₃ averaged across multiple city sites (Rodriguez et al., 2007), O ₃
5	measured at sites near schools (Moon et al., 2009; Ward et al., 2002)]. Some studies that
6	found weak or inconsistent associations between ambient O3 concentrations and
7	respiratory symptoms found O ₃ -associated decrements in lung function (Ward et al.,
8	2002; Linn et al., 1996). In their study of healthy children in Uniontown, PA, Neas et al.
9	(1995) found differences in association with respiratory symptoms between two estimates
10	of O ₃ exposure. Ambient O ₃ concentrations were measured at one central site in town.

1	Subjects spent a mean 5.4 hours outdoors during the 12-hour period (8 a.m8 p.m.) over
2	which O3 concentrations were averaged and symptoms were reported. Evening cough
3	was more strongly associated with O3 concentrations weighted by time spent outdoors
4	(OR: 2.20 [95% CI: 1.02, 4.75] per 30-ppb increase in lag 0 of 12-h avg O ₃) than with
5	unweighted O ₃ concentrations (OR: 1.36 [95% CI: 0.86, 2.13]). Time spent outdoors has
6	been shown to influence O_3 personal-ambient ratios and correlations (Section <u>4.3.3</u>), thus
7	the weighted O ₃ concentrations may have represented personal O ₃ exposures better.



Note: Results generally are presented in increasing order of mean ambient O_3 concentration. URS = Upper respiratory symptoms. Odds ratios are from single-pollutant models and are standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 8-h max (or 12-h avg), and 24-h avg O_3 concentrations, respectively.

Figure 6-13 Associations between ambient ozone concentrations and respiratory symptoms in children in the general population.

Table 6-24	Additional characteristics and quantitative data for studies
	represented inFigure 6-13.

Study*	Location/ Population	O₃ Lag	O ₃ Averaging Time	Symptom	Subgroup	Standardized OR (95% CI) ^a
<u>Ward et al.</u> (2002)	Birmingham and Sandwell, England	0-6 avg	24-h avg	Wheeze, summer Cough, summer		0.69 (0.51, 0.94)
	162 children, age 9 yr					0.98 (0.80, 1.21)
<u>Hoek and</u> <u>Brunekreef</u> (1995)	Enkhuizen, Netherlands 300 children, ages 7 - 11 yr	0	1-h max	Cough Any symptom		0.86 (0.61, 1.22)
						0.94 (0.76, 1.16)
<u>Moon et al.</u> (2009)	4 cities, South Korea 696 children, ages <13 yr	0	8-h avg (10 a.m6	URS	All subjects Jeju Island	0.96 (0.90, 1.03)
	uso children, ages < 15 yr		p.m.)			1.11 (0.95, 1.30)
Neas et al.	Uniontown, PA	0 12-h avg		Evening cough		2.20 (1.02,
<u>(1995</u>)	83 healthy children, 4th and 5th grades		(8 a.m8 p.m.)			4.75) ^b
Triche et al.	Southwestern VA	0	24-h avg	Wheeze		2.34 (1.02,
<u>(2006</u>)	691 infants of mothers with asthma, age <1 yr		8-h max 1-h max			5.37)
						1.48 (0.49, 4.41)
						1.73 (0.48, 6.22)
<u>Gold et al.</u> (1999)	Mexico City, Mexico 40 children, ages 8-11 yr	1	24-h avg	Phlegm		1.02 (1.00, 1.04)
<u>Linn et al.</u> (1996) ^c	Rubidoux, Upland, Torrence, CA 269 children, 4th and 5th grades	0	24-h avg	Evening symptom score		-0.96 (-2.2, 0.26)

*Includes studies in Figure 6-13, plus others.

URS = Upper respiratory symptoms

^aEffect estimates are standardized to a 40-, 30-, and 20-ppb increase for 1-h max, 8-h max (or 8-h avg or 12-h avg), and 24-h avg O_3 , respectively.

 ${}^{\rm b}O_3$ concentrations were weighted by the proportion of time spent outdoors.

^cResults not presented in <u>Figure 6-13</u> because outcome is a continuous variable indicating intensity of symptoms (negative indicates improvement in symptoms).

1	Several other panel studies of children, in which asthma prevalence ranged from 0 to
2	50%, reported null or negative associations between various averaging times and lags of
3	ambient O ₃ concentration and respiratory symptoms (Moon et al., 2009; Rodriguez et al.,
4	2007; Ward et al., 2002; Linn et al., 1996; Hoek and Brunekreef, 1995) (Figure 6-13 and
5	Table 6-24). Among children in Mexico City, Gold et al. (1999) reported an increase in
6	phlegm in association with an increase in lag 1 of 24-h avg O_3 concentration measured at
7	schools; however, investigators acknowledged being unable to distinguish between the
8	effects of O_3 and PM_{10} due to their high correlation (r = 0.75).
9	Unlike other studies that examined ambient O ₃ concentrations from a single monitoring
10	site, Triche et al. (2006) found respiratory symptoms to be associated with O3 measured

1	at a site that for some subjects was located >100 miles away from home (Figure 6-13 and
2	Table 6-24). Subjects included infants in Southwestern VA. Odds ratios were 46-73%
3	larger in the group who had mothers with asthma than among all infants (Triche et al.,
4	<u>2006</u>). Larger ORs were found for 24-h avg than 1-hour or 8-h max O_3 concentrations,
5	particularly for wheeze but less so for difficulty breathing. While these results suggested
6	that children with mothers with asthma may be at increased risk of O_3 -related respiratory
7	morbidity, the authors acknowledged that mothers with asthma may be more likely to
8	report symptoms in their children. Additionally, transient wheeze, which is common in
9	infants, may not predict respiratory morbidity later in life. In another cohort of children
10	with parental history of asthma that was followed to an older age (5 years), increases in
11	ambient O ₃ concentration (increment of effect estimate not reported) were not associated
12	with increases in respiratory symptoms (Rodriguez et al., 2007).

Adults

13	A cross-sectional study of 4,200 adult workers from 100 office buildings across the U.S.
14	found that multiple ambient O_3 metrics, including the 24-h, workday (8 a.m5 p.m.), and
15	late workday (3-6 p.m.) average, were associated with similar magnitudes of increase in
16	building-related symptoms (Apte et al., 2008). It should be noted that office workers
17	likely have a low personal-ambient O_3 correlation and ratio, thus the implications of these
18	findings compared to those of the other respiratory symptom studies are limited.

6.2.4.4 Confounding in Epidemiologic Studies of Respiratory Symptoms and Medication Use

19	Epidemiologic evidence does not indicate that confounding by meteorological factors or
20	copollutant exposures fully accounts for associations observed between short-term
21	increases in ambient O ₃ concentration and respiratory symptoms and medication use.
22	Except where specified in the text, studies found O3-associated increases in respiratory
23	symptoms or medication in statistical models that adjusted for temperature. Thurston et
24	al. (1997) found no independent association between temperature and respiratory
25	symptoms among children with asthma at summer camps. A few studies additionally
26	included humidity in models (Triche et al., 2006; Ross et al., 2002).
27	Several studies that examined populations with a high prevalence of atopy found
28	O ₃ -associated increases in respiratory symptoms and asthma medication use with
29	adjustment for daily pollen counts (Just et al., 2002; Ross et al., 2002; Gielen et al.,
30	1997). Gielen et al. (1997) and Ross et al. (2002) examined populations with a high
31	prevalence of grass pollen allergy (52% and 38%, respectively). In a study conducted

1	over multiple seasons, Ross et al. (2002) found a similar magnitude of association
2	between O ₃ and morning symptoms and medication use with adjustment for pollen
3	counts. Feo Brito et al. (2007) followed adults in central Spain specifically with asthma
4	and pollen allergy. In one city, O ₃ was associated with an increase in the number of
5	subjects reporting symptoms. A smaller increase was estimated for pollen. Conversely, in
6	another city, pollen was associated with an increased reporting of respiratory symptoms,
7	whereas O ₃ was not. The results suggested that O ₃ and pollen may have independent
8	effects that vary by location, depending on the mix of ambient pollutants.
9	Results from copollutant models did not indicate strong confounding by copollutants
10	such as $PM_{2.5}$, PM_{10} , sulfate, SO ₂ , or NO ₂ (<u>Table 6-25</u>). Notably, studies examined
11	different averaging times for O ₃ (1-h max or 8-h avg) and copollutants (3-hour to
12	24-h avg) and reported a range of correlations between O3 and copollutants, which may
13	complicate interpretation of copollutant model results. Information on potential
14	copollutant confounding of asthma medication use results was limited. The association
15	between O_3 and bronchodilator use did not change with adjustment for $PM_{2.5}$ in <u>Gent et</u>
16	al. (2003) but decreased in magnitude with adjustment for 12-h avg sulfate in Thurston et
17	al. (1997). In Thurston et al. (1997) and Gent et al. (2003), 1-h max O ₃ was highly
18	correlated with 12-h avg sulfate (r = 0.74) and 24-h avg $PM_{2.5}$ (r = 0.77), respectively,
19	making it difficult to distinguish the independent effects of O ₃ . Studies conducted
20	concurrently in two areas of Mexico City examined 1-h max O_3 and 24-h avg PM_{10} or
21	PM _{2.5} and found robust ORs for respiratory symptoms for both O ₃ and PM (Romieu et al.,
22	1997; Romieu et al., 1996). Romieu et al. (1997) reported a moderate correlation between
23	1-h max O_3 and 24-h avg PM_{10} (r = 0.47). Associations between O_3 and respiratory
24	symptoms were observed in NCICAS in copollutant models with SO ₂ , NO ₂ , or PM_{10} ,
25	which were examined with different averaging times and lags than was O_3 (Mortimer et
26	<u>al., 2002</u>) (Table 6-25). Also difficult are interpretations of the O_3 -associated increases in
27	respiratory symptoms found with adjustment for two copollutants in the same model
28	(i.e., $PM_{2.5}$ plus NO ₂ or $PM_{10\cdot2.5}$) (Escamilla-Nuñez et al., 2008; Triche et al., 2006).

Table 6-25Associations between ambient ozone concentrations and
respiratory symptoms in single- and co-pollutant models.

Study	Location/Population	O₃ Metrics	Symptom	OR for O ₃ in Single-Pollutant Model (95% CI) ^a	OR for O₃ in Copollutant Model (95% CI) ^ª
<u>Mortimer et</u> al. (2002)	Bronx, East Harlem, NY; Baltimore, MD; Washington, DC; Detroit, MI, Cleveland, OH; Chicago, IL; St. Louis, MO 846 children with asthma, ages 4- 9 yr	8-h avg (10 a.m 6 p.m.) Lag 1-4 avg	Morning symptoms	8 cities with SO ₂ data 1.35 (1.04, 1.74)	With lag 1-2 avg, 3-h avg SO_2 1.23 (0.94, 1.61)
				7 cites with NO ₂ data 1.25 (0.94, 1.67)	With lag 1-6 avg, 24-h avg NO <u>;</u> 1.14 (0.85, 1.55)
				3 cities with PM ₁₀ data 1.21 (0.61, 2.41)	With lag 1-2 avg, 24-h avg PM ₁₀ 1.08 (0.49, 2.39)
<u>Thurston et</u> al. (1997)	CT River Valley 166 children with asthma, ages 7-13 yr	1-h max	Chest symptoms	1.21 (1.12, 1.31) ^b	With lag 0, 12-h avg sulfate 1.19 (1.06, 1.35) ^b
		Lag 0	Beta-agonist use	1.20 (1.09, 1.32) ^b	With lag 0, 12-h avg sulfate 1.07 (0.92, 1.24) ^b
<u>Romieu et al.</u> (1996)	Mexico City, Mexico 71 children with asthma, ages 5-7 yr	1-h max Lag 0	Lower respiratory symptoms	1.07 (1.02, 1.12)	With lag 0, 24-h avg PM _{2.5} 1.06 (1.02, 1.10)
<u>Romieu et al.</u> (1997)	Mexico City, Mexico 65 children with asthma, ages 5- 13 yr	1-h max Lag 0	Lower respiratory symptoms	1.09 (1.04, 1.14)	With lag 0, 24-h avg PM ₁₀ 1.09 (1.01, 1.19)

Results generally are presented in order of increasing mean ambient O_3 concentration.

 ^{a}ORs are standardized to a 40- and 30-ppb increase for 1-h max and 8-h avg O₃, respectively.

^bTemperature not included in models.

6.2.4.5 Summary of Epidemiologic Studies of Respiratory Symptoms and Asthma Medication Use

1	Comprising a majority of available evidence, single-city and -region epidemiologic
2	studies provide consistent evidence for the effects of short-term increases in ambient O ₃
3	exposure on increasing respiratory symptoms and asthma medication use in children with
4	asthma (Figure 6-11 and Figure 6-12 and Table 6-20 and Table 6-21). Evidence from the
5	few available U.S. multicity studies is less consistent (O'Connor et al., 2008; Schildcrout
6	et al., 2006; Mortimer et al., 2002). Findings from a small body of studies indicate
7	O ₃ -associated increases in respiratory symptoms in adults with asthma. Associations
8	between short-term increases in ambient O_3 concentration and reduced activity in
9	children or adults with asthma are not clearly demonstrated. While O3-associated
10	increases in school absenteeism were found in children with asthma, evidence for
11	respiratory-related absences and for O_3 exposure lag periods shorter than 14 days is
12	sparse. Short-term increases in ambient O3 concentration were not consistently associated

with increases in respiratory symptoms in groups comprising children with and without asthma.

- 3 Increases in respiratory symptoms and medication use were associated with increases in 4 ambient O₃ concentration assigned to subjects using various methods. Associations were 5 found with methods likely to represent better ambient exposures, including O_3 measured 6 on site and at the time of children's outdoor activity (Thurston et al., 1997) and 7 concentrations weighted by time spent outdoors (Neas et al., 1995). However, 8 associations also were found with methods that varied in their representation of ambient 9 exposures and spatial variability in ambient concentrations, i.e., concentrations averaged 10 among subjects' locations each hour (Khatri et al., 2009), measured within 5 km of schools or homes (Escamilla-Nuñez et al., 2008; Romieu et al., 2006; 1997; 1996), 11 12 averaged across multiple sites (Feo Brito et al., 2007; Gent et al., 2003; Mortimer et al., 2002), and measured at a single site (Ross et al., 2002; Gielen et al., 1997). 13
- 14 Associations with respiratory symptoms were demonstrated most frequently for 1-h max 15 and 8-h max or avg O₃, and within-study comparisons indicated similar ORs for 1-h max 16 and 8-h max O₃ (Delfino et al., 2003; Gent et al., 2003). Respiratory symptoms also were 17 associated with 12-hour and 24-h avg O₃ (Jalaludin et al., 2004; Gold et al., 1999; Neas et 18 al., 1995). Epidemiologic studies examined respiratory symptoms associated with O₃ 19 concentrations lagged 0 to 5 days and those averaged over 2 to 19 days. While O_3 at lags 20 0 or 1 were consistently associated with respiratory symptoms, several studies found 21 larger ORs for multiday averages (3- to 6-day) of O_3 (Escamilla-Nuñez et al., 2008; 22 Romieu et al., 2006; Just et al., 2002; Mortimer et al., 2002; Ross et al., 2002). 23 Epidemiologic findings for lagged or multiday average O_3 are supported by evidence that 24 O_3 sensitizes bronchial smooth muscle to hyperreactivity and thus acts as a primer for 25 subsequent exposure to antigens such as allergens (Section 5.3.5). Many studies 26 examined populations with asthma with a high prevalence of atopy (52-100%). In these 27 populations, sensitization of airways provides a biologically plausible mode of action by 28 which increases in respiratory symptoms result from increases in O_3 exposure after a lag 29 or accumulated over several days. Further support is provided by findings that airway 30 hyperresponsiveness (Section 6.2.2.1) and some indicators of inflammation 31 (Section 6.2.3.1) remained elevated following repeated O_3 exposures in controlled human 32 exposure studies and by observations from epidemiologic studies that increases in 33 pulmonary inflammation were associated with multiday average O_3 concentrations 34 (Section 6.2.3.2).
- 35There is not strong evidence that O3-associated increases in respiratory symptoms are36confounded by temperature, pollen, or copollutants. In limited analysis, ambient O3 was37associated with respiratory symptoms with adjustment for copollutants, primarily PM.

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1However, identifying the independent effects of O3 in some studies was complicated due2to the high correlations observed between O3 and PM or different lags and averaging3times examined for copollutants. Nonetheless, the consistency of associations among4individuals with asthma with and without adjustment for ambient copollutant5concentrations combined with findings from controlled human exposure studies for the6direct effect of O3 exposure provide substantial evidence for the independent effects of7short-term ambient O3 exposure on increasing respiratory symptoms.

6.2.5 Lung Host Defenses

8 The mammalian respiratory tract has a number of closely integrated defense mechanisms 9 that, when functioning normally, provide protection from the potential health effects 10 attributed to exposure to a wide variety of inhaled particles and microbes. For simplicity, 11 these interrelated defenses can be divided into two major parts: (1) nonspecific (transport, 12 phagocytosis, and bactericidal activity) and (2) specific (immunologic) defense 13 mechanisms. A variety of sensitive and reliable methods have been used to assess the 14 effects of O_3 on these components of the lung's defense system to provide a better 15 understanding of the health effects associated with the inhalation of this pollutant. The 16 previous O₃ AQCD stated that animal toxicological studies provide extensive evidence 17 that acute O_3 exposures as low as 0.08 to 0.5 ppm can cause increases in susceptibility to 18 infectious diseases due to modulation of lung host defenses. Table 6-6 through Table 6-9 19 (U.S. EPA, 1996g, h, i, j) beginning on page 6-41 of the 1996 O₃ AQCD (U.S. EPA, 20 1996a), and Table AX5-7 (U.S. EPA, 2006c), beginning on page AX5-8 of the 2006 O_3 21 AQCD (U.S. EPA, 2006b), present studies on the effects of O_3 on host defense 22 mechanisms. This section discusses the various components of host defenses, such as the 23 mucociliary escalator, the phagocytic, bactericidal, and regulatory role of the alveolar 24 macrophages (AMs), the adaptive immune system, and integrated mechanisms that are 25 studied by investigating the host's response to experimental pulmonary infections.

6.2.5.1 Mucociliary Clearance

The mucociliary system is one of the lung's primary defense mechanisms. It protects the conducting airways by trapping and quickly removing material that has been deposited or is being cleared from the alveolar region by migrating alveolar macrophages. Ciliary movement directs particles trapped on the overlying mucous layer toward the pharynx, where the mucus is swallowed or expectorated.

1 The effectiveness of mucociliary clearance can be determined by measuring such 2 biological activities as the rate of transport of deposited particles; the frequency of ciliary 3 beating; structural integrity of the ciliated cells; and the size, number, and distribution of 4 mucus-secreting cells. Once this defense mechanism has been altered, a buildup of both 5 viable and nonviable inhaled substances can occur on the epithelium and may jeopardize 6 the health of the host, depending on the nature of the uncleared substance. Impaired 7 mucociliary clearance can result in an unwanted accumulation of cellular secretions, 8 increased infections, chronic bronchitis, and complications associated with COPD. A 9 number of previous studies with various animal species have examined the effect of O_3 10 exposure on mucociliary clearance and reported morphological damage to the cells of the 11 tracheobronchial tree from acute and sub-chronic exposure to $O_3 0.2$ ppm and higher. The 12 cilia were either completely absent or had become noticeably shorter or blunt. After 13 placing these animals in a clean-air environment, the structurally damaged cilia 14 regenerated and appeared normal (U.S. EPA, 1986). Based on such morphological 15 observations, related effects such as ciliostasis, increased mucus secretions, and a slowing 16 of mucociliary transport rates might be expected. However, no measurable changes in 17 ciliary beating activity have been reported due to O₃ exposure alone. Essentially no data 18 are available on the effects of prolonged exposure to O_3 on ciliary functional activity or 19 on mucociliary transport rates measured in the intact animal. In general, functional 20 studies of mucociliary transport have observed a delay in particle clearance soon after 21 acute exposure. Decreased clearance is more evident at higher doses (1 ppm), and there is 22 some evidence of attenuation of these effects (U.S. EPA, 1986). However, no recent 23 studies have evaluated the effects of O_3 on mucociliary clearance.

6.2.5.2 Alveolobronchiolar Transport Mechanism

24	In addition to the transport of particles deposited on the mucous surface layer of the
25	conducting airways, particles deposited in the deep lung may be removed either up the
26	respiratory tract or through interstitial pathways to the lymphatic system. The pivotal
27	mechanism of alveolobronchiolar transport involves the movement of AMs with
28	phagocytized particles to the bottom of the mucociliary escalator. Failure of the AMs to
29	phagocytize and sequester the deposited particles from the vulnerable respiratory
30	membrane can lead to particle entry into the interstitial spaces. Once lodged in the
31	interstitium, particle removal is more difficult and, depending on the toxic or infectious
32	nature of the particle, its interstitial location may allow the particle to set up a focus for
33	pathologic processes. Although some studies show reduced early (tracheobronchial)
34	clearance after O ₃ exposure, late (alveolar) clearance of deposited material is accelerated,

presumably due to macrophage influx (which in itself can be damaging due to proteases and oxidative reactions in these cells).

6.2.5.3 Alveolar Macrophages

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3 Within the gaseous exchange region of the lung, the first line of defense against 4 microorganisms and nonviable particles that reach the alveolar surface is the AM. This 5 resident phagocyte is responsible for a variety of activities, including the detoxification 6 and removal of inhaled particles, maintenance of pulmonary sterility via destruction of 7 microorganisms, and interaction with lymphocytes for immunologic protection. Under 8 normal conditions, AMs seek out particles deposited on the alveolar surface and ingest 9 them, thereby sequestering the particles from the vulnerable respiratory membrane. To 10 adequately fulfill their defense function, the AMs must maintain active mobility, a high 11 degree of phagocytic activity, and an optimally functioning biochemical and enzyme 12 system for bactericidal activity and degradation of ingested material. As discussed in 13 previous AOCDs, short periods of O_3 exposure can cause a reduction in the number of 14 free AMs available for pulmonary defense, and these AMs are more fragile, less 15 phagocytic, and have decreased lysosomal enzyme activities required for killing 16 pathogens. For example, in results from earlier work in rabbits, a 2-hour exposure to 17 0.1 ppm O₃ inhibited phagocytosis and a 3-hour exposure to 0.25 ppm decreased 18 lysosomal enzyme activities (Driscoll et al., 1987; Hurst et al., 1970). Similarly, AMs 19 from rats exposed to 0.1 ppm O_3 for 1 or 3 weeks exhibited reduced hydrogen peroxide 20 production (Cohen et al., 2002). A controlled human exposure study reported decrements 21 in the ability of alveolar macrophages to phagocytize yeast following exposure of healthy 22 volunteers to 80 to 100 ppb O_3 for 6.6-hour during moderate exercise (Devlin et al., 23 1991). Although the percentage of phagocytosis-capable macrophages was unchanged by 24 O₃ exposure, the number of yeast engulfed was reduced when phagocytosis was 25 complement-dependent. However, there was no difference in the ability of macrophages 26 to produce superoxide anion after O_3 exposure. These results are consistent with those 27 from another controlled human exposure study in which no changes in the level of 28 lysosomal enzymes or superoxide anion production were observed in macrophages 29 lavaged from healthy human subjects exposed to 400 ppb O₃ for 2 hours with heavy 30 intermittent exercise (Koren et al., 1989). More recently, Lay et al. (2007) observed no 31 difference in phagocytic activity or oxidative burst capacity in macrophages or 32 monocytes from sputum or blood collected from healthy volunteers after a 2-hour 33 exposure to 400 ppb O₃ with moderate intermittent exercise. However, another study 34 found that oxidative burst and phagocytic activity in macrophages increased in GSTM1 35 null subjects compared to GSTM1 positive subjects, who had relatively unchanged

1	macrophage function parameters after an O_3 exposure identical to that of Lay et al.
2	described above (<u>Alexis et al., 2009</u>). Collectively, these studies demonstrate that O_3 can
3	affect multiple steps or aspects required for proper macrophage function, but any C-R
4	relationship appears complex and genotype may be a consideration. A few other recent
5	studies have evaluated the effects of O_3 on macrophage function, but these are of
6	questionable relevance due to the use of in vitro exposure systems and amphibian animal
7	models (Mikerov et al., 2008c; Dohm et al., 2005; Klestadt et al., 2005).

6.2.5.4 Infection and Adaptive Immunity

General Effects on the Immune System

8	The effects of O_3 on the immune system are complex and dependent on the exposure
9	regimen and the observation period. According to toxicological studies it appears that the
10	T-cell-dependent functions of the immune system are more affected than B-cell-
11	dependent functions (U.S. EPA, 2006b). Generally, there is an early immunosuppressive
12	effect that subsides with continued O ₃ exposure, resulting in either a return to normal
13	responses or an enhancement of immune responses. However, this is not always the case
14	as Aranyi et al. Aranyi et al. (1983) showed decreased T-cell mitogen reactions in mice
15	after subchronic (90-day) exposure to 0.1 ppm O ₃ . Earlier studies report changes in cell
16	populations in lymphatic tissues (U.S. EPA, 2006b). A more recent study in mice
17	demonstrated that numbers of certain T-cell subsets in the spleen were reduced after
18	exposure to 0.6 ppm O_3 (10h/day x 15d) (Feng et al., 2006).
19	The inflammatory effects of O_3 involve the innate immune system, and as such, O_3 can
20	affect adaptive (or acquired) immunity via alterations in antigen presentation and
20 21	costimulation by innate immune cells such as macrophages and dendritic cells. Several
22	recent controlled human exposure studies demonstrate increased expression of molecules
23	involved in antigen presentation or costimulation. Lay et al. (2007) collected sputum
24	monocytes from healthy volunteers exposed to 400 ppb O_3 for 2 hours with moderate
25	intermittent exercise and detected increases in HLA-DR, used to present antigen to
26	T-cells, and CD86, a costimulatory marker necessary for T-cell activation. Upregulation
27	of HLA-DR was also observed by <u>Alexis et al. (2009</u>) in sputum dendritic cells and
28	macrophages from GSTM1 null subjects exposed to 400 ppb O_3 for 2 hours with
29	moderate intermittent exercise. On airway monocytes from healthy volunteers 24 hours
30	after exposure to 80 ppb O_3 for 6.6 hours with moderate intermittent exercise, HLA-DR,
31	CD86, and CD14 (a molecule involved in bacterial endotoxin reactivity) were increased,
32	whereas CD80, a costimulatory molecule of more heterogeneous function, was decreased
33	(Alexis et al., 2010). Patterns of expression on macrophages were similar, except that

1	HLA-DR was found to be significantly decreased after O_3 exposure and CD86 was not
2	significantly altered. An increase in IL-12p70, a macrophage and dendritic cell product
3	that activates T-cells, was correlated with increased numbers of dendritic cells. It should
4	be noted that these results are reported as comparisons to baseline as there was no clean
5	air control (<u>Alexis et al., 2010</u> ; <u>Alexis et al., 2009</u>). Another controlled human exposure
6	study reported no increase in IL-12p70 in sputum from healthy, atopic, or atopic
7	asthmatic subjects following a 2-hour exposure to 400 ppb O_3 with intermittent moderate
8	exercise (<u>Hernandez et al., 2010</u>). Levels of HLA-DR, CD14 and CD86 were not
9	increased on macrophages collected from any of these subjects. It is difficult to compare
10	these results to those of Lay et al. (2007) and Alexis et al. (2010) due to differences in O_3
11	concentration, cell type examined, and timing of postexposure analysis.
11	concentration, cen type examined, and timing of postexposure analysis.
12	Although no controlled human exposure studies have examined the effects of O ₃ on the
13	ability to mount antigen-specific responses, upregulation of markers associated with
14	innate immune activation and antigen presentation could potentially enhance adaptive
15	immunity and increase immunologic responses to antigen. While this may bolster
16	defenses against infection, it also may enhance allergic responses (Section $6.2.6$).
17	In animal models, O ₃ has been found to alter responses to antigenic stimulation. For
18	example, antibody responses to a T-cell-dependent antigen were suppressed after a
19	56-day exposure of mice to 0.8 ppm O_3 , and a 14-day exposure to 0.5 ppm O_3 decreased
20	the antiviral antibody response following influenza virus infection (Jakab and Hmieleski,
21	<u>1988</u>); the latter impairment may pave the way for lowered resistance to re-infection. The
22	immune response is highly influenced by the temporal relationship between O ₃ exposure
23	and antigenic stimulation. When O ₃ exposure preceded Listeria infection, there were no
24	effects on delayed-type hypersensitivity or splenic lymphoproliferative responses;
25	however, when O ₃ exposure occurred during or after <i>Listeria</i> infection was initiated,
26	these immune responses were suppressed (Van Loveren et al., 1988). In another study, a
27	reduction in mitogen activated T-cell proliferation was observed after exposure to
28	0.6 ppm O_3 for 15 days that could be ameliorated by antioxidant supplementation.
29	Antigen-specific proliferation decreased by 60%, indicating attenuation of the acquired
30	immunity needed for subsequent memory responses (Feng et al., 2006). O_3 exposure also
31	skewed the ex-vivo cytokine responses elicited by non-specific stimulation toward
32	inflammation, decreasing IL-2 and increasing IFN-y. Modest decreases in immune
33	function assessed in the offspring of O3-exposed dams (mice) were observed by Sharkhuu
34	et al. (2011). The ability to mount delayed-type hypersensitivity responses was
35	significantly suppressed in 42 day-old offspring when dams were exposed to 0.8 or
36	1.2 ppm O_3 , but not 0.4 ppm, from gestational day 9-18. Humoral responses to
37	immunization with sheep red blood cells were unaffected, as were other immune
38	parameters such as splenic populations of CD45+ T-cells, iNKT-cells, and levels of IFN-

1 γ , IL-4, and IL-17 in the BALF. Generally, continuous exposure to O_3 impairs immune2responses for the first several days of exposure, followed by an adaptation to O_3 that3allows a return of normal immune responses. Most species show little effect of O_3 4exposures prior to immunization, but show a suppression of responses to antigen in O_3 5exposures post-immunization.

Microbial Infection

Bacterial infection

6 A relatively large body of evidence shows that O_3 increases susceptibility to bacterial 7 infections. The majority of studies in this area were conducted before the 1996 O₃ AOCD 8 was published and many are included in Table 6-9 (U.S. EPA, 1996j) on page 6-53 of 9 that document. Known contributing factors are impaired mucociliary streaming, altered 10 chemotaxis/motility, defective phagocytosis of bacteria, decreased production of 11 lysosomal enzymes or superoxide radicals by alveolar macrophages, and decreased IFN- γ 12 levels. In animal models of bacterial infection, exposure to 0.08 ppm O_3 increases 13 streptococcus-induced mortality, regardless of whether O_3 exposure precedes or follows 14 infection (Miller et al., 1978; Coffin and Gardner, 1972; Coffin et al., 1967). Increases in 15 mortality are due to the infectious agent, thereby reflecting functional impairment of host 16 defenses. Exercise and copollutants can enhance the effects of O₃ in infectivity models. 17 Although both mice and rats exhibit impaired bactericidal macrophage activity after O_3 18 exposure, mortality due to infection is only observed in mice. Additionally, although 19 mice and humans share many host defense mechanisms, there is little compelling 20 evidence from epidemiologic studies to suggest an association between O_3 exposure and 21 decreased resistance to bacterial infection, and the etiology of respiratory infections is not 22 easily identified via ICD codes (Section 6.2.7.3).

Viral infection

23 Only a few studies, described in previous AQCDs, have examined the effects of O3 24 exposure on the outcome of viral respiratory infection [see Table 6-9 on page 6-53 of the 25 1996 O₃ AQCD (U.S. EPA, 1996)]. Some studies show increased mortality, while others 26 show diminished severity and increased survival time. There is little to no evidence from 27 studies of animals or humans to suggest that O_3 increases the incidence of respiratory 28 viral infection in humans. In human volunteers infected with rhinovirus prior to O₃ 29 exposure (0.3 ppm for 5 consecutive days), no effect on viral titers, IFN- γ production, or 30 blood lymphocyte proliferative responses to viral antigen was observed (Henderson et al., 31 1988). In vitro cell culture studies of human bronchial epithelial cells indicate O_3 -induced 32 exacerbation of human rhinovirus infection (Spannhake et al., 2002), but this is of limited 33 relevance. More recent studies on the interactions of O₃ and viral infections have not been

1	published. Natural killer (NK) cells, which destroy virally infected cells and tumors in the
2	lung, appear to be inhibited by higher concentrations of O_3 and either unaffected or
3	stimulated at lower concentrations. Several studies show decreases in NK cell activity
4	following acute exposures ranging from 0.8 to 1 ppm (Gilmour and Jakab, 1991; Van
5	Loveren et al., 1990; Burleson et al., 1989). However, Van Loveren et al. (1990) showed
6	that a 1-week exposure to 0.2 or 0.4 ppm O_3 increased NK cell activity, and an urban
7	pattern of exposure (base of 0.06 ppm with peaks of 0.25 ppm) had no effect on NK cell
8	activity after 1, 3, 13, 52, or 78 weeks of exposure (Selgrade et al., 1990). A more recent
9	study demonstrated a 35% reduction in NK cell activity after exposure of mice to
10	0.6 ppm O_3 (10h/day x 15d) (Feng et al., 2006). The defective IL-2 production
11	demonstrated in this study may impair NK cell activation. Alternatively, NK cell surface
12	charge may be altered by ROS, decreasing their adherence to target cells (Nakamura and
13	<u>Matsunaga, 1998</u>).

6.2.5.5 Summary of Lung Host Defenses

14	Taken as a whole, the data clearly indicate that an acute O_3 exposure impairs the host
15	defense capability of animals, primarily by depressing AM function and perhaps also by
16	decreasing mucociliary clearance of inhaled particles and microorganisms. Coupled with
17	limited evidence from controlled human exposure studies, this suggests that humans
18	exposed to O ₃ could be predisposed to bacterial infections in the lower respiratory tract.
19	The seriousness of such infections may depend on how quickly bacteria develop
20	virulence factors and how rapidly PMNs are mobilized to compensate for the deficit in
21	AM function. It remains unclear how O ₃ might affect antigen presentation and the
22	costimulation required for T-cell activation, given the mixed results from controlled
23	human exposure studies, but there is toxicological evidence for suppression of T-cell-
24	dependent functions by O ₃ , including reductions in antigen-specific proliferation and
25	antibody production, indicating the potential for impaired acquired immunity and
26	memory responses. To date, a limited number of epidemiologic studies have examined
27	associations between O3 exposure and hospital admissions or ED visits for respiratory
28	infection, pneumonia, or influenza. Results have been mixed, and in some cases
29	conflicting (see Section $6.2.7.2$ and Section $6.2.7.3$). With the exception of influenza, it is
30	difficult to ascertain whether cases of respiratory infection or pneumonia are of viral or
31	bacterial etiology. A study that examined the association between O ₃ exposure and
32	respiratory hospital admissions in response to an increase in influenza intensity did
33	observe an increase in respiratory hospital admissions (Wong et al., 2009), but
34	information from toxicological studies of O_3 and viral infections is ambiguous.

6.2.6 Allergic and Asthma-Related Responses

1	Effects resulting from combined exposures to O3 and allergens have been studied in a
2	variety of animal species, generally as models of experimental asthma. Pulmonary
3	function and airways hyperresponsiveness in animal models of asthma are discussed in
4	Section <u>6.2.1.3</u> and Section <u>6.2.2.2</u> . Previous evidence indicates that O_3 exposure skews
5	immune responses toward an allergic phenotype. For example, Gershwin et al. (1981)
6	reported that O_3 (0.8 and 0.5 ppm for 4 days) exposure caused a 34-fold increase in the
7	number of IgE (allergic antibody)-containing cells in the lungs of mice. In general, the
8	number of IgE-containing cells correlated positively with levels of anaphylactic
9	sensitivity. In humans, allergic rhinoconjunctivitis symptoms are associated with
10	increases in ambient O_3 concentrations (<u>Riediker et al., 2001</u>). Recent controlled human
11	exposure studies have observed O3-induced changes indicating allergic skewing. Airway
12	eosinophils, which participate in allergic disease and inflammation, were observed to
13	increase in atopic, mildly asthmatic volunteers 18 hours following a 7.6-hour exposure to
14	160 ppb O_3 with light intermittent exercise (Peden et al., 1997). No increase in airway
15	eosinophils was observed 4 hours after exposure of healthy, atopic, or atopic asthmatic
16	subjects to 400 ppb O_3 for 2 hours with moderate intermittent exercise (<u>Hernandez et al.</u> ,
17	2010). However, atopic subjects did exhibit increased IL-5, a cytokine involved in
18	eosinophil recruitment and activation, suggesting that perhaps these two studies observed
19	the same effect at different time points. Epidemiologic studies discussed in Section $7.2.5$
20	describe an association between eosinophils and long-term O_3 exposure, consistent with
21	chronic exposure studies in non-human primates. Hernandez et al. (2010) also observed
22	increased expression of high and low affinity IgE receptors on sputum macrophages from
23	atopic asthmatics, which may enhance IgE-dependent inflammation. Sputum levels of
24	IL-4 and IL-13, both pro-allergic cytokines that aid in the production of IgE, were
25	unaltered in all groups. The lack of increase in IL-4 levels in sputum reported by
26	Hernandez et al. (2010), along with increased IL-5, is consistent with results from Bosson
27	et al. (2003), in which IL-5 (but not IL-4 levels) increased in bronchial epithelial biopsy
28	specimens following exposure of mild atopic asthmatics to 200 ppb O_3 for 2 hours with
29	moderate intermittent exercise. IL-5 was not elevated in specimens obtained from healthy
30	(non-asthmatic) O_3 -exposed subjects. Collectively, findings from these studies suggest
31	that O3 can induce or enhance certain components of allergic inflammation in atopic and
32	atopic asthmatic individuals.
33	Ozone enhances inflammatory and allergic responses to allergen challenge in sensitized
34	animals. Short-term exposure (2 days) to 1 ppm O ₃ exacerbated allergic rhinitis and lower
35	airway allergic inflammation in Brown Norway rats, a rat strain that is comparatively less
36	sensitive to O_3 than other rats or humans (<u>Wagner et al., 2009</u> ; <u>2007</u>). OVA-sensitized
37	rats were intranasally challenged with OVA on days 1 and 2, and exposed to 0 or 1 ppm

1	O ₃ (8 h/day) on days 4 and 5. Analysis at day 6 indicated that O ₃ exposure enhanced
2	intraepithelial mucosubstances in the nose and airways, induced cys-LTs, MCP-1, and
3	IL-6 production in BALF, and upregulated expression of the proallergic cytokines IL-5
4	and IL-13. These changes were not evident in non-allergic controls. All of these
5	responses were blunted by gamma-tocopherol (γT ; vitamin E) therapy. γT neutralizes
6	oxidized lipid radicals, and protects lipids and proteins from nitrosative damage from
7	NO-derived metabolites. Farraj et al. (2010) exposed allergen-sensitized adult male
8	BALB/c mice to 0.5 ppm O_3 for 5 hours once per week for 4 weeks. Ozone exposure and
9	O_3 /DEP (2.0 mg/m ³) co-exposure of OVA-sensitized mice elicited significantly greater
10	serum IgE levels than in DEP-exposed OVA-sensitized mice (98% and 89% increases,
11	respectively). Ozone slightly enhanced levels of BAL IL-5, but despite increases in IgE,
12	caused a significant decrease in BAL IL-4 levels. IL-10, IL-13, and IFN- γ levels were
13	unaffected. Lung resistance and elastance were unaffected in allergen sensitized mice
14	exposed solely to 0.5 ppm O_3 once a week for 4 weeks (<u>Farraj et al., 2010</u>). However,
15	co-exposure to O_3 and diesel exhaust particles increased lung resistance.

16 In addition to exacerbating existing allergic responses, O_3 can also act as an adjuvant to 17 produce sensitization in the respiratory tract. In a model of murine asthma, using OVA 18 free of detectable endotoxin, inclusion of 1 ppm O_3 during the initial exposures to OVA 19 (2 h, days 1 and 6) enhanced the inflammatory and allergic responses to subsequent 20 allergen challenge (Hollingsworth et al., 2010). Compared to air exposed animals, 21 O₃-exposed mice exhibited significantly higher levels of total cells, macrophages, 22 eosinophils, and PMNs in BALF, and increased total serum IgE. Pro-allergic cytokines 23 IL-4, and IL-5 were also significantly elevated, along with pleiotropic Th2 cytokine IL-9 24 (associated with bronchial hyperresponsiveness) and pro-inflammatory IL-17, produced 25 by activated T-cells. Based on lower inflammatory, IgE, and cytokine responses in 26 Toll-like receptor 4 deficient mice, the effects of O_3 seem to be dependent on TLR 4 27 signaling, as are a number of other biological responses to O_3 according to studies by 28 Hollingsworth et al. (2004), Kleeberger et al. (2000) and Garantziotis et al. (2010). The 29 involvement of TLR 4, along with its endogenous ligand, hyaluronan, in O_3 -induced 30 responses described in these studies has been corroborated by a controlled human exposure study by Hernandez et al. (2010), who found increased TLR 4 expression and 31 32 elevated levels of hyaluronic acid in atopic and atopic asthmatic volunteers exposed to 33 400 ppb O_3 . This pathway is discussed in more detail in Chapter 5. Examination of 34 dendritic cells (DCs) from the draining thoracic lymph nodes indicated that O₃ did not 35 enhance the migration of DCs from the lungs to the lymph nodes, nor did it alter the 36 expression of functional DC markers such as CD40, MHC class II, or CD83. However, 37 O₃ did increase expression of CD86, which is generally associated with Th2 responses 38 and is detected at higher levels on DCs from allergic asthmatics compared to those from 39 healthy donors Chen et al. (2006b). Increased CD86 has also been observed on airway

1 2 3	cells collected from human subjects following exposure to O_3 in studies by <u>Lay et al.</u> (2007) and <u>Alexis et al. (2009</u>), but not <u>Hernandez et al. (2010</u>) (study details described in Section <u>6.2.5.4</u>).
4	Ozone exposure during gestation has modest effects on allergy and asthma related
5	endpoints in adult offspring. When dams were exposed to 1.2 ppm O_3 (but not 0.8 ppm)
6	from gestational day 9-18, some allergic and inflammatory responses to OVA
7	sensitization and challenge were reduced compared to air exposed controls. This included
8	IgE levels and eosinophils, and was only true of mice that were immunized early in life
9	(PND 3) as opposed to later (PND 42), perhaps due to the proximity of O_3 and antigen
10	exposure. The effects of gestational O ₃ exposure on immune function have not been
11	widely studied, and although reductions in allergic endpoints are not generally observed
12	in association with O ₃ , other parameters of immune function were found to be reduced, so
13	a more global immunosuppression may underlie these effects.
14	In addition to pro-allergic effects, O ₃ could also make airborne allergens more allergenic.
15	When combined with NO ₂ , O ₃ has been shown to enhance nitration of common protein
16	allergens, which may increase their allergenicity Franze et al. (2005).

6.2.7 Hospital Admissions, Emergency Department Visits, and Physicians Visits

6.2.7.1 Summary of Findings from 2006 Ozone AQCD

17	The 2006 O_3 AQCD evaluated numerous respiratory ED visits and hospital admissions
18	studies, which consisted primarily of time-series studies conducted in the U.S., Canada,
19	Europe, South America, Australia and Asia. Upon collectively evaluating the scientific
20	evidence, the 2006 O_3 AQCD concluded that "the overall evidence supports a causal
21	relationship between acute ambient O3 exposures and increased respiratory morbidity
22	resulting in increased ED visits and [hospital admissions] during the warm season" U.S.
23	EPA (2006b). This conclusion was "strongly supported by the human clinical, animal
24	toxicologic[al], and epidemiologic evidence for [O ₃ -induced] lung function decrements,
25	increased respiratory symptoms, airway inflammation, and airway hyperreactivity" U.S.
26	<u>EPA (2006b</u>).
27	Since the completion of the 2006 O_3 AQCD, relatively fewer studies conducted in the
28	U.S., Canada, and Europe have examined the association between short-term exposure to
29	ambient O_3 and respiratory hospital admissions and ED visits with a growing number of
30	studies having been conducted in Asia. This section focuses primarily on multicity

1	studies because they examine the effect of O3 on respiratory-related hospital admissions
2	and ED visits over a large geographic area using a consistent statistical methodology.
3	Single-city studies that encompass a large number of hospital admissions or ED visits, or
4	included a long study-duration were also evaluated because these studies have more
5	power to detect whether an association exists between short-term O ₃ exposure and
6	respiratory hospital admissions and ED visits compared to smaller single-city studies.
7	Additional single-city studies were also evaluated within this section, if they were
8	conducted in locations not represented by the larger single-city and multicity studies, or
9	examined population-specific characteristics not included in the larger studies that may
10	modify the association between short-term O ₃ exposure and respiratory-related hospital
11	admissions or ED visits. The remaining single-city studies identified were not evaluated
12	in this section due to factors such as inadequate study design or insufficient sample size.
12	It should be mostioned that when even ining the approximitien between short terms O
13	It should be mentioned that when examining the association between short-term O_3
14	exposure and respiratory health effects that require medical attention, it is important to
15	distinguish between hospital admissions and ED visits. This is because it is likely that a
16	small percentage of respiratory ED visits will be admitted to the hospital; therefore,
17	respiratory ED visits may represent potentially less serious, but more common outcomes.
18	As a result, in the following sections respiratory hospital admission and ED visit studies
19	are evaluated individually. Additionally, within each section, results are presented as
20	either a collection of respiratory diagnoses or as individual diseases (e.g., asthma, COPD,
21	pneumonia and other respiratory infections) in order to evaluate the potential effect of
22	short-term O ₃ exposure on each respiratory-related outcome. The ICD codes (i.e., ICD-9
23	or ICD-10) that encompass each of these endpoints are presented in Table 6-26 along
24	with the air quality characteristics of the city, or across all cities, included in each study
25	evaluated in this section.

Table 6-26Mean and upper percentile concentrations of respiratory-related
hospital admission and emergency department (ED) visit studies
evaluated

Study	Location	Type of Visit (ICD9/10)	Averaging Time	Mean Concentration (ppb) ^a	Upper Percentile Concentrations (ppb) ^a
Katsouyanni et al. (2009) ^{bc}	90 U.S. cities (NMMAPS) ^d 32 European cities (APHEA) ^d 12 Canadian cities	Hospital Admissions: NMMAPS: All respiratory (460-519) APHEA: All respiratory (460-519) 12 Canadian cities: All respiratory (460-519) ^e	1-h max	NMMAPS: 50th: 34.9-60.0 APHEA: 50th: 11.0-38.1 12 Canadian cities: 50th: 6.7-8.3	NMMAPS: 75th: 46.8-68.8 APHEA: 75th: 15.3-49.4 12 Canadian cities: 75th: 8.4-12.4
<u>Cakmak et al.</u> (2006b)	10 Canadian cities	Hospital Admissions: All respiratory (466, 480-486, 490, 491, 492, 493, 494, 496)	24-h avg	17.4	Max: 38.0-79.0
<u>Biggeri et al.</u> (2005) ^c	4 Italian cities ^f	Hospital Admissions: All respiratory (460-519)	8-h max	Warm season (May-September): 5.7-60.0	95th: 86.1-90.0 Max: 107.5-115.1
<u>Dales et al.</u> (2006)	11 Canadian cities	Hospital Admissions: Respiratory disorders (486, 768.9, 769, 770.8, 786, 799.0, 799.1)	24-h avg	17.0	95th: 24.9-46.0
<u>Lin et al.</u> (2008a)	11 New York regions	Hospital Admissions: Respiratory diseases (466, 490- 493, 496)	8-h max ^g	44.1	75th: 54.0 Max: 217.0
<u>Wong et al.</u> (2009) ^c	Hong Kong	Hospital Admissions: All respiratory (460-519) COPD (490-496)	8-h max ⁹	18.8	75th: 25.9 Max: 100.3
<u>Medina-Ramon</u> et al. (2006) ^h	36 U.S. cities	Hospital Admissions: COPD (490-496, excluding 493) Pneumonia (480-487)	8-h max	Warm (May-September): 45.8 Cool (October-April): 27.6	NR
Yang et al. (2005b)	Vancouver, Canada	Hospital Admissions: COPD (490-492, 494, 496)	24-h avg	All year: 14.1 Winter (January-March): 13.2 Spring (April-June): 19.4 Summer (July-September): 13.8 Fall (October-December): 10.0	Max: 38.6
Zanobetti and Schwartz (2006) ^b	Boston, MA	Hospital Admissions: Pneumonia (480-487)	24-h avg	22.4	75th: 31.0 95th: 47.6
Silverman and Ito (2010) ^b	New York, NY	Hospital Admissions: Asthma (493)	8-h max	Warm (April-August): 41.0	75th: 53 90th: 68

Study	Location	Type of Visit (ICD9/10)	Averaging Time	Mean Concentration (ppb) ^a	Upper Percentile Concentrations (ppb) ^a
<u>Stieb et al.</u> (2009)	7 Canadian cities	ED Visits: Asthma (493) COPD (490-492, 494-496) Respiratory infection (464, 466, 480-487)	24-h avg	18.4	75th: 19.3-28.6
<u>Tolbert et al.</u> (2007)	Atlanta, GA	ED Visits: All respiratory (460-465, 460.0, 466.1, 466.11, 466.19, 477, 480- 486, 491, 492, 493, 496, 786.07, 786.09)	8-h max	Warm: 53.0	75th: 67.0 90th: 82.1 Max: 147.5
<u>Darrow et al.</u> (<u>2011a</u>)	Atlanta, GA	ED Visits: All respiratory (460-466, 477, 480-486, 491, 492, 493, 496,	8-h max	Warm (March-October): 8-h max: 53	8-h max :75th: 67 8-h max :Max: 148
		786.09)	1-h max	Warm (March-October): 1-h max: 62	1-h max :75th: 76 1-h max :Max: 180
			24-h avg	Warm (March-October): 24-h avg: 30	24-h avg :75th: 37 24-h avg :Max: 81
			Commute	Warm (March-October): Commute: 35 ⁱ	Commute :75th: 45 Commute :Max: 106
			Day-time	Warm (March-October): Day-time: 45 ⁱ	Day-time :75th: 58 Day-time :Max: 123
			Night-time	Warm (March-October): Night-time: 14 ⁱ	Night-time :75th: 22 Night-time :Max: 64
Villeneuve et al. (2007) ⁶	Alberta, CAN	ED Visits: Asthma (493)	8-h max	Summer (April-September): 38.0 Winter (October-March): 24.3	Summer: 75th: 46.0 Winter: 75th: 31.5
<u>ito et al. (2007b</u>)	New York, NY	ED Visits: Asthma (493)	8-h max	All year: 30.4 Warm (April-September): 42.7 Cold (October-March): 18.0	All year: 95th: 68.0 Warm months: 95th: 77.0 Cold months: 95th: 33.0
<u>Strickland et al.</u> (2010)	Atlanta, GA	ED Visits: Asthma (493) Wheeze (786.07 after 10/1/98, 786.09 before 10/1/98)	8-h max	All year: 45.4 ⁱ Warm (May-October): 55.2 ⁱ Cold (November-April): 34.5 ⁱ	NR
<u>Mar and Koenig</u> (2009)	Seattle, WA	ED Visits: Asthma (493-493.9)	1-h max 8-h max	Warm (May-October): 1-h max: 38.6 8-h max: 32.2	75th: 1-h max: 45.5 8-h max: 39.2
A <u>rbex et al.</u> (2009)	Sao Paulo, Brazil	ED Visits: COPD (J40-44)	1-h max	48.8	75th: 61.0 Max: 143.8

Study	Location	Type of Visit (ICD9/10)	Averaging Time	Mean Concentration (ppb) ^a	Upper Percentile Concentrations (ppb) ^a
<u>Orazzo et al.</u> (2009) ^c	6 Italian cities	ED Visits: Wheezing	8-h max ^k	Summer (April-September): 21.1-44.3	NR
				Winter (October-March): 11.5-27.9	
Burra et al.	Toronto,	Physician Visits:	1-h max	33.3	95th: 66
(2009) Car	Canada	ED Asthma (493)			Max: 121
Villeneuve et al. (2006b)	Toronto, Canada	Physician Visits: Allergic rhinitis (177)	8-h max	30.0	Max: 98.7
<u>Sinclair et al.</u> (2010) ^I	Atlanta, GA	Physician Visits: Asthma	8-h max	Total Study Period: All-year: 44.0	NR
		Upper respiratory infection Lower respiratory infection		25 mo Period: All-year: 47.9 Warm: 61.2 Cold: 27.8	
				28 mo Period: All-year: 40.7 Warm: 51.8 Cold: 26.0	

^aSome studies did not present an overall value for the mean, middle and/or upper percentiles of the O₃ distribution; as a result, the range of the mean, middle, and/or upper percentiles across all of the cities included in the study are presented.

^bStudy only presented median concentrations.

^cStudy presented concentrations as μg/m³ Concentration was converted to ppb using the conversion factor of 0.51 assuming standard temperature (25°C) and pressure (1 atm).

^dA subset of the European and U.S. cities included in the mortality analyses were used in the hospital admissions analyses: 8 of the 32 European cities and 14 of 90 U.S. cities.

^eHospital admission data was coded using three classifications (ICD-10-CA, ICD-9, and ICD-9-CM). Attempts were made by the original investigators to convert diagnosis from ICD-10-CA back to ICD-9.

^fOnly 4 of the 8 cities included in the study collected O₃ data.

 ${}^{g}O_{3}$ measured from 10:00 a.m. to 6:00 p.m.

 h Only 35 of the 36 cities included in the analysis had O₃ data.

ⁱCommute (7:00 a.m. to 10:00 a.m., 4:00 p.m. to 7:00 p.m.); day-time (8:00 a.m. to 7:00 p.m.); Night-time (12:00 a.m. to 6:00 a.m.).

^jMeans represent population-weighted O₃ concentrations.

 kO_3 measured from 8:00 a.m. to 4:00 p.m.

¹This study did not report the ICD codes used for the conditions examined. The 25-month period represents August 1998-August 2000, and the 28-month period represents September 2000-December 2002. This study defined the warm months as April – October and the cold months as November-March.

6.2.7.2 Hospital Admission Studies

Respiratory Diseases

1	The association between exposure to an air pollutant, such as O ₃ , and daily respiratory-
2	related hospital admissions has primarily been examined using all respiratory-related
3	hospital admissions within the range of ICD-9 codes 460-519. Recent studies published
4	since the 2006 AQCD (U.S. EPA, 2006b) attempt to further examine the effect of O_3
5	exposure on respiratory-related hospital admissions through a multicity design that
6	examines O3 effects across countries using a standardized methodology; multicity studies

that examine effects within one country; and multi- and single-city studies that attempt to examine potential modifiers of the O_3 -respiratory-related hospital admission relationship.

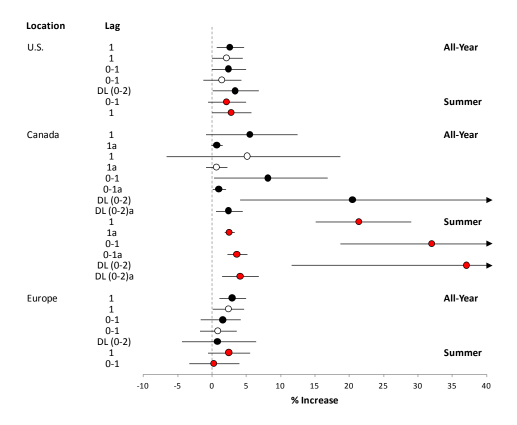
- 3 The Air Pollution and Health: A European and North American Approach (APHENA) 4 study combined data from existing multicity study databases from Canada, Europe 5 (APHEA2) (Katsouyanni et al., 2001), and the U.S. (NMMAPS) (Samet et al., 2000) in 6 order to "develop more reliable estimates of the potential acute effects of air pollution on 7 human health [and] provide a common basis for [the] comparison of risks across 8 geographic areas" (Katsouyanni et al., 2009). In an attempt to address both of these 9 issues, the investigators conducted extensive sensitivity analyses to evaluate the 10 robustness of the results to different model specifications (e.g., penalized splines [PS] 11 versus natural splines [NS]) and the extent of smoothing to control for seasonal and 12 temporal trends. The trend analyses consisted of subjecting the models to varying extent 13 of smoothing selected either a priori (i.e., 3 df/year, 8 df/year, and 12 df/year), which was 14 selected through exploratory analyses using between 2 and 20 df, or by using the absolute 15 sum of the residuals of the partial autocorrelation function (PACF). Although the 16 investigators did not identify the model they deemed to be the most appropriate for 17 comparing the results across study locations, they did specify that "overall effect 18 estimates (i.e., estimates pooled over several cities) tended to stabilize at high degrees of 19 freedom" (Katsouyanni et al., 2009). Therefore, in discussion of the results across the 20 three study locations below, the 8 df/year results are presented for both the PS and NS 21 models because: (1) 8 df/year is most consistent with the extent of temporal adjustment 22 used in previous and recent large multicity studies in the U.S. (e.g., NMMAPS); (2) the 23 risk estimates for 8 df/year and 12 df/year are comparable for all three locations; (3) the 24 models that used the PACF method did not report the actual degrees of freedom chosen; 25 and (4) the 3 df/year and the PACF method resulted in negative O_3 risk estimates, which 26 is inconsistent with the results obtained using more aggressive seasonal adjustments and 27 suggests inadequate control for seasonality. Additionally, in comparisons of results across 28 studies in figures, only the results from one of the spline models (i.e., NS) are presented 29 because it has been previously demonstrated that alternative spline models result in 30 relatively similar effect estimates (HEI, 2003). This observation is consistent with the 31 results of the APHENA analysis that was conducted with a higher number of degrees of 32 freedom (e.g., ≥ 8 df/year) to account for temporal trends.
- 33Katsouyanni et al. (2009) examined respiratory hospital admissions for people aged3465 years and older using 1-h max O3 data. The extent of hospital admission and O3 data35varied across the 3 datasets: Canadian dataset included 12 cities with data for 3 years36(1993-1996) per city; European dataset included 8 cities with each city having data for37between 2 and 8 years from 1988-1997; and the U.S. dataset included 14 cities with each38city having data for 4 to 10 years from 1985-1994 and 7 cities having only summer O3

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1	data. The investigators used a three-stage hierarchical model to account for within-city,
2	within region, and between region variability. Results were presented individually for
3	each region (Figure 6-14; Table 6-27). Ozone and PM_{10} concentrations were weakly
4	correlated in all locations in the summer ($r = 0.27-0.40$), but not in the winter.
5	In the Canadian cities, using all-year data, a 40 ppb increase in 1-h max O_3
6	concentrations at lag 0-1 was associated with an increase in respiratory hospital
7	admissions of 8.9% (95% CI: 0.79, 16.8%) in a PS model and 8.1% (95% CI: 0.24,
8	16.8%) in a NS model (Katsouyanni et al., 2009). The results were somewhat sensitive to
9	the lag day selected, reduced when using a single-day lag (e.g., lag 1) (PS: 6.0%; NS:
10	5.5%) and increased when using a distributed lag model (PS: 18.6%; NS: 20.4%). When
11	adjusting for PM_{10} , the magnitude of the effect estimate was attenuated, but remained
12	positive with it being slightly larger in the NS model (5.1% [95% CI: -6.6, 18.6%])
13	compared to the PS model (3.1% [95% CI: -8.3, 15.9%]). However, in the Canadian
14	dataset the copollutant analysis was only conducted using a 1-day lag. The large
15	confidence intervals for both models could be attributed to the reduction in days included
16	in the copollutant analyses as a result of the every-6th-day PM sampling schedule. When
17	the analysis was restricted to the summer months, stronger associations were observed
18	between O3 and respiratory hospital admissions across the lags examined, ranging from
19	\sim 22 to 37% (the study does not specify whether these effect estimates are from a NS or
20	PS model). Because O ₃ concentrations across the cities included in the Canadian dataset
21	are low (median concentrations ranging from 6.7-8.3 ppb [Table 6-26]), the standardized
22	increment of 40 ppb for a 1-h max increase in O ₃ concentrations represents an unrealistic
23	increase in O_3 concentrations in Canada and increases the magnitude, not direction, of the
24	observed risk estimate. As a result, calculating the O ₃ risk estimate using the standardized
25	increment does not accurately reflect the observed risk of O3-related respiratory hospital
26	admissions. Although this increment adequately characterizes the distribution of 1-h max
27	O_3 concentrations across the U.S. and European datasets, it misrepresents the observed O_3
28	concentrations in the Canadian dataset. As a result in summary figures, for comparability,
29	effect estimates from the Canadian dataset are presented for both a 5.1 ppb increase in
30	1-h max O ₃ concentrations (i.e., an approximate interquartile range [IQR] increase in O ₃
31	concentrations across the Canadian cities) as well as the standardized increment used
32	throughout the ISA.
33	In Europe, weaker but positive associations were also observed in year round analyses;
34	2.9% (95% CI: 0.63, 5.0%) in the PS model and 1.6% (95% CI: -1.7, 4.2%) in the NS
35	model at lag 0-1 for a 40 ppb increase in 1-h max O ₃ concentrations (Katsouyanni et al.,
36	<u>2009</u>). Additionally, at lag 1, associations between O_3 and respiratory hospital admissions
37	were also reduced, but in contrast to the lag 0-1 analysis, greater effects were observed in
38	the NS model (2.9% [95% CI: 1.0, 4.9%]) compared to the PS model (1.5% [95% CI:

1	-2.2, 5.4]). Unlike the Canadian analysis, a distributed lag model provided limited
2	evidence of an association between O3 and respiratory hospital admissions. To compare
3	with the Canadian results, with adjustment for PM_{10} at lag 1, O_3 effect estimates were
4	increased in the PS model (2.5% [95% CI: 0.39-4.8%]) and remained robust in the NS
5	model (2.4% [95% CI: 0.08, 4.6%]). However, the European analysis also examined the
6	effect of adjusting for PM_{10} at lag 0-1 and found results were attenuated, but remained
7	positive in both models (PS: 0.8% [95% CI: -2.3, 4.0%]; NS: 0.8% [95% CI: -1.8,
8	3.6%]). Unlike the Canadian and U.S. datasets, the European dataset consisted of daily
9	PM data. The investigators did not observe stronger associations in the summer-only
10	analyses for the European cities at lag 0-1 (PS: 0.4% [95% CI: -3.2, 4.0%]; NS: 0.2%
11	[95% CI: -3.3, 3.9%]), but did observe some evidence for larger effects during the
12	summer, an ~2.5% increase, at lag 1 in both models (the study does not present the extent
13	of temporal smoothing used for these models).



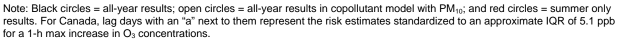


Figure 6-14 Percent increase in respiratory hospital admissions from natural spline models with 8 df/yr for a 40 ppb increase in 1-h max ozone concentrations for each location of the APHENA study.

Location*	Season	Lag ^a	Copollutant	% Increase (95% CI) ^b
U.S.	All-year	1		2.62 (0.63, 4.64)
		1	PM ₁₀	2.14 (-0.08, 4.40)
		0-1		2.38 (0.00, 4.89)
		0-1	PM ₁₀	1.42 (-1.33, 4.23)
		DL(0-2)		3.34 (0.02-6.78)
	Summer	0-1		2.14 (-0.63, 4.97)
		1		2.78 (-0.02, 5.71)
Canada	All-year	1		5.54 (-0.94, 12.4)
		1a		0.69 (-0.12, 1.50) ^a
		1	PM ₁₀	5.13 (-6.62, 18.6)
		1a	PM ₁₀	0.64 (-0.87, 2.20) ^a
		0-1		8.12 (0.24, 16.8)
		0-1a		1.00 (0.03, 2.00) ^a
		DL(0-2)		20.4 (4.07, 40.2)
		DL(0-2) ^a		2.4 (0.51, 4.40) ^a
	Summer	1		21.4 (15.0, 29.0)
		1a		2.50 (1.80, 3.30) ^a
		0-1		32.0 (18.6, 47.7)
		0-1 ^ª		3.60 (2.20, 5.10) ^a
		DL(0-2)		37.1 (11.5, 67.5)
		DL(0-2) ^a		4.1 (1.40, 6.80) ^a
Europe	All-year	1		2.94 (1.02, 4.89)
		1	PM ₁₀	2.38 (0.08, 4.64)
		0-1		1.58 (-1.71, 4.15)
		0-1	PM ₁₀	0.87 (-1.79, 3.58)
		DL(0-2)		0.79 (-4.46, 6.37)
	Summer	1		2.46 (-0.63, 5.54)
		0-1		0.24 (-3.32, 3.91)

Table 6-27 Corresponding effect estimates for Figure 6-14.

*For effect estimates in Figure 6-14.

^aFor Canada, lag days with an "a" next to them represent the risk estimates standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations.

^bUnless noted, risk estimates standardized to 40 ppb for a 1-h max increase in O_3 concentrations.

1	For the U.S. in year round analyses, the investigators reported a 1.4% (95% CI: -0.9,
2	3.9%) increase in the PS model and 2.4% (95% CI: 0.0, 4.9%) increase in the NS model
3	in respiratory hospital admissions at lag 0-1 for a 40 ppb increase in 1-h max O_3
4	concentrations with similar results for both models at lag 1 (Katsouyanni et al., 2009).
5	The distributed lag model provided results similar to those observed in the European

- 1 dataset with the PS model (1.1% [95% CI: -3.0, 5.3%]), but larger effects in the NS 2 model (3.3% [95% CI: 0.02, 6.8%]), which is consistent with the Canadian results. With 3 adjustment for PM₁₀ using the U.S. data (i.e., every-6th-day PM data), results were 4 attenuated, but remained positive at lag 0-1 (PS: 0.6% [95% CI: -2.0, 3.3%]; NS: 1.4% 5 [95% CI: -1.3, 4.2%]) which is consistent with the results presented for the European 6 dataset. However, at lag 1, U.S. risk estimates remained robust to the inclusion of PM_{10} in 7 copollutant models as was observed in the Canadian and European datasets. Compared to 8 the all-year analyses, the investigators did not observe stronger associations in the 9 summer-only analysis at either lag 0-1 (\sim 2.2%) or lag 1 (\sim 2.8%) in both the PS and NS 10 models (the study does not present the extent of temporal smoothing used for these 11 models).
- 12 Several additional multicity studies examined respiratory disease hospital admissions in 13 Canada and Europe. Cakmak et al. (2006b) evaluated the association between ambient O_3 14 concentrations and respiratory hospital admissions for all ages in 10 Canadian cities from 15 April 1993 to March 2000. The primary objective of this study was to examine the 16 potential modification of the effect of ambient air pollution on daily respiratory hospital 17 admissions by education and income using a time-series analysis conducted at the city-18 level. The authors calculated a pooled estimate across cities for each pollutant using a 19 random effects model by first selecting the lag day with the strongest association from the 20 city-specific models. For O_3 , the mean lag day across cities that provided the strongest 21 association and for which the pooled effect estimate was calculated was 1.2 days. In this 22 study, all-year O_3 concentrations were used in the analysis, and additional seasonal 23 analyses were not conducted. Cakmak et al. (2006b) reported a 4.4% increase (95% CI: 24 2.2, 6.5%) in respiratory hospital admissions for a 20 ppb increase in 24-hour average O_3 25 concentrations. The investigators only examined the potential effect of confounding by 26 other pollutants through the use of a multipollutant model (i.e., two or more additional 27 pollutants included in the model), which is difficult to interpret due to the potential 28 multicollinearity between pollutants. Cakmak et al. (2006b) also conducted an extensive 29 analysis of potential modifiers, specifically sex, educational attainment, and family 30 income, on the association between air pollution and respiratory hospital admissions. 31 When stratifying by sex, the increase in respiratory hospital admissions due to short-term 32 O₃ exposure were similar in males (5.2% [95% CI: 3.0, 7.3%]) and females (4.2% 33 [95% CI: 1.8, 6.6%]). In addition, the examination of effect modification by income 34 found no consistent trend across the quartiles of family income. However, there was 35 evidence that individuals with an education level less than the 9th grade were 36 disproportionately affected by O_3 exposure (4.6% [95% CI: 1.8, 7.5%]) compared to 37 individuals that completed grades 9-13 (1.7% [95% CI: -1.9, 5.3%]), some university or 38 trade school (1.4% [95% CI: -2.0, 5.1%]), or have a university diploma (0.66% [95% CI: 39 -3.3, 4.7%]). The association between O₃ and respiratory hospital admissions in

individuals with an education level less than the 9th grade was the strongest association across all of the pollutants examined.

- 3 A multicity study conducted in Europe by Biggeri et al. (2005) examined the association 4 between short-term O₃ exposure and respiratory hospital admissions for all ages in four 5 Italian cities from 1990 to 1999. In this study, O_3 was only measured during the warm 6 season (May-September). The authors examined associations between daily respiratory 7 hospital admissions and short-term O₃ exposure at the city-level using a time-series 8 analysis. Pooled estimates were calculated by combining city-specific estimates using 9 fixed and random effects models. The investigators found no evidence of an association 10 between O_3 exposure and respiratory hospital admissions in the warm season in both the 11 random (0.1% [95% CI: -5.2, 5.7%]; distributed lag 0-3) and fixed effects (0.1% 12 [95% CI: -5.2, 5.7%]; distributed lag 0-3) models for a 30 ppb increase in 8-h max O₃ 13 concentrations.
- 14 Additional studies examined associations between short-term O₃ exposure and respiratory 15 hospital admissions specifically in children. In a multicity study conducted in Canada, 16 Dales et al. (2006) examined the association between all-year ambient O₃ concentrations 17 and neonatal (ages 0-27 days) respiratory hospital admissions in 11 Canadian cities from 18 1986 to 2000. The investigators used a statistical analysis approach similar to Cakmak et 19 al. (2006b) (i.e., time-series analysis to examine city-specific associations, and then a 20 random effects model to pool estimates across cities). The authors reported that for O_3 21 the mean lag day across cities that provided the strongest association was 2 days. The 22 authors reported a 5.4% (95% CI: 2.9, 8.0%) increase in neonatal respiratory hospital 23 admissions for a 20 ppb increase in 24-h avg O₃ concentrations at lag-2 days. The results 24 from Dales et al. (2006) provide support for the associations observed in a smaller scale 25 study that examined O_3 exposure and pediatric respiratory hospital admissions in 26 New York state (Lin et al., 2008a). Lin et al. (2008a), when examining single-day lags of 27 0 to 3 days, observed a positive association between O_3 and pediatric (i.e., <18 years) 28 respiratory admissions at lag 2 (results not presented quantitatively) in a two-stage 29 Bayesian hierarchical model analysis of 11 geographic regions of New York state from 30 1991 to 2001. Additionally, in copollutant models with PM_{10} , collected every-6th day, the 31 authors found region-specific O₃ associations with respiratory hospital admissions 32 remained relatively robust.
- 33Overall, the evidence from epidemiologic studies continues to support an association34between short-term O3 exposure and respiratory-related hospital admissions, but it35remains unclear whether certain factors (individual- or population-level) modify this36association. Wong et al. (2009) examined the potential modification of the relationship37between ambient O3 (along with NO2, SO2, and PM10) and respiratory hospital

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2

 air pollution concentrations were estimated by centering non-missing daily air pollution data on the annual mean for each monitor and then an overall daily concentration was calculated by taking the average of the daily centered mean across all monitors. Influenza intensity was defined as a continuous variable using the proportion of weekly specimens positive for influenza A or B instead of defining influenza epidemics. This approach was
 4 calculated by taking the average of the daily centered mean across all monitors. Influenza 5 intensity was defined as a continuous variable using the proportion of weekly specimens 6 positive for influenza A or B instead of defining influenza epidemics. This approach was
 5 intensity was defined as a continuous variable using the proportion of weekly specimens 6 positive for influenza A or B instead of defining influenza epidemics. This approach was
6 positive for influenza A or B instead of defining influenza epidemics. This approach was
7 used to avoid any potential bias associated with the unpredictable seasonality of influenza
8 in Hong Kong where there are traditionally two seasonal peaks, which is in contrast to the
9 single peaking influenza season in the U.S. (Wong et al., 2009). In models that examined
10 the baseline effect (i.e., without taking into consideration influenza intensity) of short-
11 term O_3 exposure, the authors found a 3.6% (95% CI: 1.9, 5.3%) and 3.2% (95% CI: 1.0,
12 5.4%) increase in respiratory hospital admissions at lag 0-1 for a 30 ppb increase in
13 8-h max O_3 concentrations for the all age and ≥ 65 age groups, respectively. When
14 examining influenza intensity, <u>Wong et al. (2009</u>) reported that the association between
15 short-term exposure to O ₃ and respiratory hospital admissions was stronger with higher
16 levels of influenza intensity: additional increase in respiratory hospital admissions above
17 baseline of 1.4% (95% CI: 0.24, 2.6%) for all age groups and 2.4% (95% CI: 0.94, 3.8%)
18 for those 65 and older when influenza activity increased from 0% to 10%. No difference
19 in effects was observed when stratifying by sex.

Cause-Specific Respiratory Outcomes

20	In the 2006 O ₃ AQCD a limited number of studies were identified that examined the
21	effect of short-term O ₃ exposure on cause-specific respiratory hospital admissions. The
22	limited evidence "reported positive O_3 associations with asthma and COPD,
23	especially during the summer or warm season" (U.S. EPA, 2006b). Of the studies
24	evaluated since the completion of the 2006 O ₃ AQCD, more have focused on identifying
25	whether O3 exposure is associated with specific respiratory-related hospital admissions,
26	including COPD, pneumonia, and asthma, but the overall body of evidence remains
27	small.

Chronic Obstructive Pulmonary Disease

28Medina-Ramon et al. (2006) examined the association between short-term exposure to29ambient O_3 and PM_{10} concentrations and Medicare hospital admissions among30individuals \geq 65 years of age for COPD in 35 cities in the U.S. for the years 1986-1999.31The cities included in this analysis were selected because they monitored PM_{10} on a daily32basis. In this study, city-specific results were obtained using a monthly time-stratified33case-crossover analysis. A meta-analysis was then conducted using random effects34models to combine the city-specific results. All cities measured O_3 from May through

1	September, while only 16 of the cities had year-round measurements. The authors
2	reported a 1.6% increase (95% CI: 0.48, 2.9%) in COPD admissions for lag 0-1 in the
3	warm season for a 30 ppb increase in 8-h max O_3 concentrations. When examining
4	single-day lags, stronger associations were observed for lag 1 (2.9% [95% CI: 1.8, 4.0%])
5	compared to lag 0 (-1.5% [95% CI: -2.7, -0.24%]). The authors found no evidence of
6	associations in cool season (-1.9% [95% CI: -3.6, -0.06%]; lag 0-1) or year round (0.24%
7	[95% CI: -0.78, 1.2%]; lag 0-1) analyses. In a copollutant model restricted to days in
8	which PM_{10} was available, the association between O_3 and COPD hospital admissions
9	remained robust. Of note, the frequency of PM_{10} measurements varied across cities with
10	measurements collected either every 2, 3, or 6 days. The authors conducted additional
11	analyses to examine potential modification of the warm season estimates for O_3 and
12	COPD admissions by several city-level characteristics: percentage living in poverty,
13	emphysema mortality rate (as an indication of smoking), daily summer apparent
14	temperature, and percentage of households using central air conditioning. Of the city-
15	level characteristics examined, stronger associations were only reported for cities with a
16	smaller variability in daily apparent summer temperature.
17	In a single-city study conducted in Vancouver from 1994-1998, a location with low
18	ambient O ₃ concentrations (Table 6-26), Yang et al. (2005b) examined the association
19	between O_3 and COPD. Ozone was moderately inversely correlated with CO (r = -0.56),
20	NO ₂ (r = -0.32), and SO ₂ (r = -0.34), and weakly inversely correlated with PM_{10}
21	(r = -0.09), suggesting that the observed O_3 effect is likely not only due to a positive
22	correlation with other pollutants. Yang et al. (2005b) examined 1- to 7-day (e.g., (0-
23	6 days) lagged moving averages and observed an 8.8% (95% CI: -12.5, 32.6%) increase
24	in COPD admissions for lag 0-3 per 20 ppb increase in 24-h avg O_3 concentrations. In
25	two-pollutant models with every-day data for NO ₂ , SO ₂ , and PM ₁₀ at lag 0-3, O_3 risk
26	estimates remained robust, but were increased slightly when CO was added to the model
27	(<u>Figure 6-19;</u> <u>Table 6-29</u>).
28	In the study discussed above, Wong et al. (2009) also examined the potential
29	modification of the relationship between ambient O_3 and COPD hospital admissions by
30	influenza intensity. The authors also found evidence of an additional increase in COPD
31	admissions above baseline when influenza activity increased from 0% to 10% of 1.0%
32	(95% CI: -0.82, 2.9%) for all age groups and 2.4% (95% CI: 0.41, 4.4%) for those 65 and
33	older. The baseline increase in COPD hospital admissions at lag 0-1 for a 30 ppb increase
34	in 8-h max O_3 concentrations was 8.5% (95% CI: 5.6, 11.4%) for the all age and 4.2%
35	$(95\% \text{ CI: } 1.1, 7.3\%) \ge 65 \text{ age groups.}$

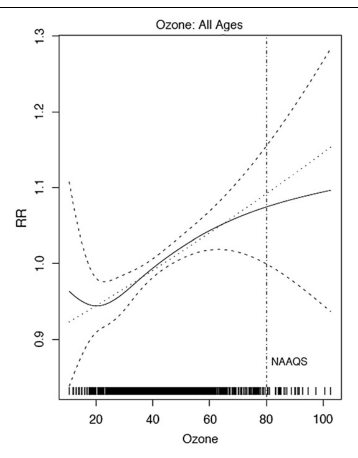
Pneumonia

1	In addition to COPD, Medina-Ramon et al. (2006) examined the association between
2	short-term exposure to ambient O_3 and PM_{10} concentrations and Medicare hospital
3	admissions among individuals \geq 65 years of age for pneumonia (ICD-9: 480-487). The
4	authors reported an increase in pneumonia-hospital admissions in the warm season (2.5%
5	[95% CI: 1.6, 3.5%] for a 30 ppb increase in 8-h max O ₃ concentrations; lag 0-1). Similar
6	to the results observed for COPD hospital admissions, pneumonia-hospital admissions
7	associations were stronger at lag 1 (2.6% [95% CI: 1.8, 3.4%]) compared to lag 0 (0.06%
8	[95% CI: -0.72, 0.78%]), and no evidence of an association was observed in the cool
9	season or year round. In two-pollutant models restricted to days for which PM_{10} data was
10	available, as discussed above, the association between O ₃ exposure and pneumonia-
11	hospital admissions remained robust (results not presented quantitatively). The authors
12	also examined potential effect modification of the warm season estimates for O3-related
13	pneumonia-hospital admissions, as was done for COPD, by several city-level
14	characteristics. Stronger associations were reported in cities with a lower percentage of
15	central air conditioning use. Across the cities examined, the percentage of households
16	having central air conditioning ranged from 6 to 93%. The authors found no evidence of
17	effect modification of the O3-pneumonia-hospital admission relationship when examining
18	the other city-level characteristics.
19	Results from a single-city study conducted in Boston did not support the results presented
20	by Medina-Ramon et al. (2006). Zanobetti and Schwartz (2006) examined the association
21	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was
21 22	•
	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was
22	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with
22 23	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis,
22 23 24	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia
22 23 24 25	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O ₃ concentrations at lag 0 and a
22 23 24 25 26	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O ₃ concentrations at lag 0 and a 6.0% (95% CI: -11.1, -1.4%) decrease for the average of lags 0 and 1. It should be noted
22 23 24 25 26 27	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O ₃ concentrations at lag 0 and a 6.0% (95% CI: -11.1, -1.4%) decrease for the average of lags 0 and 1. It should be noted that the mean daily counts of pneumonia admissions was low for this study, ~14
22 23 24 25 26 27 28	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O ₃ concentrations at lag 0 and a 6.0% (95% CI: -11.1, -1.4%) decrease for the average of lags 0 and 1. It should be noted that the mean daily counts of pneumonia admissions was low for this study, ~14 admissions per day compared to ~271 admissions per day for <u>Medina-Ramon et al.</u>
22 23 24 25 26 27 28 29	of O_3 and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with $PM_{2.5}$ (r = 0.20) and weakly inversely correlated with black carbon, NO ₂ , and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O ₃ concentrations at lag 0 and a 6.0% (95% CI: -11.1, -1.4%) decrease for the average of lags 0 and 1. It should be noted that the mean daily counts of pneumonia admissions was low for this study, ~14 admissions per day compared to ~271 admissions per day for <u>Medina-Ramon et al.</u> (2006). However, in analyses with other pollutants <u>Zanobetti and Schwartz (2006</u>) did
22 23 24 25 26 27 28 29 30	 of O₃ and pneumonia Medicare hospital admissions for the period 1995-1999. Ozone was weakly positively correlated with PM_{2.5} (r = 0.20) and weakly inversely correlated with black carbon, NO₂, and CO (-0.25, -0.14, and -0.30, respectively). In an all-year analysis, the investigators reported a 3.8% (95% CI: -7.9, -0.1%) decrease in pneumonia admissions for a 20 ppb increase in 24-hour average O₃ concentrations at lag 0 and a 6.0% (95% CI: -11.1, -1.4%) decrease for the average of lags 0 and 1. It should be noted that the mean daily counts of pneumonia admissions was low for this study, ~14 admissions per day compared to ~271 admissions per day for Medina-Ramon et al. (2006). However, in analyses with other pollutants Zanobetti and Schwartz (2006) did observe positive associations with pneumonia-hospital admissions, indicating that the low

Asthma

33There are relatively fewer studies that examined the association between short-term34exposure to O_3 and asthma hospital admissions, presumably due to the limited power35given the relative rarity of asthma hospital admissions compared to ED or physician

1	visits. A study from New York City examined the association of 8-h max O ₃
2	concentrations with severe acute asthma admissions (i.e., those admitted to the Intensive
3	Care Unit [ICU]) during the warm season in the years 1999 through 2006 (Silverman and
4	<u>Ito, 2010</u>). In this study, O_3 was moderately correlated with PM_{10} (r = 0.59). When
5	stratifying by age, the investigators reported positive associations with ICU asthma
6	admissions for the 6- to 18-year age group (26.8% [95% CI: 1.4, 58.2%] for a 30 ppb
7	increase in maximum 8-h avg O_3 concentrations at lag 0-1), but little evidence of
8	associations for the other age groups examined (<6 years, 19-49, 50+, and all ages).
9	However, positive associations were observed for each age-stratified group and all ages
10	for non-ICU asthma admissions, but again the strongest association was reported for the
11	6- to 18-years age group (28.2% [95% CI: 15.3, 41.5%]; lag 0-1). In two-pollutant
12	models, O ₃ effect estimates for both non-ICU and ICU hospital admissions remained
13	robust to adjustment for PM _{2.5} . In an additional analysis, using a smooth function, the
14	authors examined whether the shape of the C-R curve for O_3 and asthma hospital
15	admissions (i.e., both general and ICU for all ages) is linear. To account for the potential
16	confounding effects of PM _{2.5} , Silverman and Ito (2010) also included a smooth function
17	of $PM_{2.5}$ lag 0-1. When comparing the curve to a linear fit line the authors found that the
18	linear fit is a reasonable approximation of the C-R relationship between O_3 and asthma
19	hospital admissions around and below the level of the 1997 O_3 NAAQS (Figure 6-15).



Note: The average of 0-day and 1-day lagged 8-hour O_3 was used in a two-pollutant model with $PM_{2.5}$ lag 0-1, adjusting for temporal trends, day of the week, and immediate and delayed weather effects. The solid lines are smoothed fit data, with long broken lines indicating 95% confidence bands. The density of lines at the bottom of the figure indicates sample size. Source: Reprinted with permission of the American Academy of Allergy, Asthma & Immunology (Silverman and Ito, 2010).

Figure 6-15 Estimated relative risks (RRs) of asthma hospital admissions for 8-h max ozone concentrations at lag 0-1 allowing for possible nonlinear relationships using natural splines.

Averting Behavior

1	The studies discussed above have found consistent positive associations between short-
2	term O ₃ exposure and respiratory-related hospital admissions, however, the strength of
3	these associations may be underestimated due to the studies not accounting for averting
4	behavior. As discussed in Section 4.6.5, a recent study (Neidell and Kinney, 2010;
5	Neidell, 2009) conducted in Southern California demonstrate that controlling for
6	avoidance behavior increases O_3 effect estimates for respiratory hospital admissions,
7	specifically for children and older adults. These studies show that on days where no
8	public alert was issued warning of high O3 concentrations there was an increase in asthma
9	hospital admissions. Although only one study has examined averting behavior and this

1study is limited to the outcome of asthma hospital admissions in one location (i.e., Los2Angeles, CA) for the years 1989-1997, it does provide preliminary evidence indicating3that epidemiologic studies may underestimate associations between O3 exposure and4health effects by not accounting for behavioral modification when public health alerts are5issued.

6.2.7.3 Emergency Department Visit Studies

Overall, relatively fewer studies have examined the association between short-term O_3 exposure and respiratory-related ED visits, compared to hospital admissions. In the 2006 O_3 AQCD, positive, but inconsistent, associations were observed between O_3 and respiratory-related ED visits with effects generally occurring during the warm season. Since the completion of the previous AQCD, larger studies have been conducted, in terms of sample size, study duration, and in some cases multiple cities, to examine the association between O_3 and ED visits for all respiratory diseases, COPD, and asthma.

Respiratory Disease

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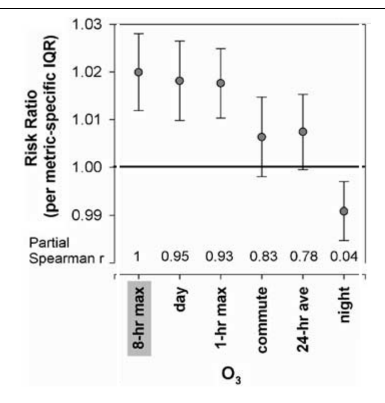
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- 13 A large single-city study conducted in Atlanta, by Tolbert et al. (2007), and subsequently 14 re analyzed by <u>Darrow et al. (2011a)</u> using different air quality data, provides evidence 15 for an association between short-term exposures to ambient O₃ concentrations and 16 respiratory ED visits. Tolbert et al. (2007) examined the association between air 17 pollution, both gaseous pollutants and PM and its components, and respiratory disease 18 ED visits in all ages from 1993 to 2004. The correlations between O₃ and the other 19 pollutants examined ranged from 0.2 for CO and SO₂ to 0.5-0.6 for the PM measures. 20 Using an a priori average of lags 0-2 for each air pollutant examined, the authors reported 21 a 3.9% (95% CI: 2.7, 5.2%) increase in respiratory ED visits for a 30 ppb increase in 22 8-h max O_3 concentrations during the warm season [defined as March-October in Darrow 23 et al. (2011a)]. In copollutant models, limited to days in which data for all pollutants were 24 available, O₃ respiratory ED visits associations with CO, NO₂, and PM₁₀, were attenuated, 25 but remained positive (results not presented quantitatively).
- 26 Darrow et al. (2011a) examined the same health data as Tolbert et al. (2007), but used air 27 quality data from one centrally located monitor instead of the average of multiple 28 monitors. This study primarily focused on exploring whether differences exist in the 29 association between O_3 exposure and respiratory-related ED visits depending on the 30 exposure metric used (i.e., 8-h max, 1-h max, 24-hour average, commuting period [7:00 31 a.m. to 10:00 a.m.; 4:00 p.m. to 7:00 p.m.], day-time [8:00 a.m. to 7:00 p.m.] and night-32 time [12:00 a.m. to 6:00 a.m.]). An ancillary analysis of the spatial variability of each 33 exposure metric conducted by Darrow et al. (2011a) found a rather homogenous spatial

1	distribution of O_3 concentrations (r~>0.8) as the distance from the central monitor
2	increased from 10 km to 60 km for all exposure durations, except the night-time metric.
3	The relatively high spatial correlation gives confidence in the use of a single monitor and
4	the resulting risk estimates. To examine the association between the various O_3 exposure
5	metrics and respiratory ED visits, the authors conceptually used a time-stratified case-
6	crossover framework where control days were selected as those days within the same
7	calendar month and maximum temperature as the case day. However, instead of
8	conducting a traditional case-crossover analysis, the authors used a Poisson model with
9	indicator variables for each of the strata (i.e., parameters of the control days). Darrow et
10	al. (2011a) found using an a prior lag of 1 day, the results were somewhat variable across
11	exposure metrics. The strongest associations with respiratory ED visits were found when
12	using the 8-h max, 1-h max, and day-time exposure metrics with weaker associations
13	using the 24-h avg and commuting period exposure metrics; a negative association was
14	observed when using the night-time exposure metric (Figure 6-16). These results indicate
15	that using the 24-h avg exposure metric may lead to smaller O ₃ -respiratory ED visits risk
16	estimates due to: (1) the dilution of relevant O ₃ concentrations by averaging over hours
17	(i.e., nighttime hours) during which O_3 concentrations are known to be low and (2)
18	potential negative confounding by other pollutants (e.g., CO, NO ₂) during the nighttime
19	hours (<u>Darrow et al., 2011a</u>).



Source: Reprinted with permission of Nature Publishing Group (Darrow et al., 2011a).

Figure 6-16 Risk ratio for respiratory ED visits and different ozone exposure metrics in Atlanta from 1993-2004.

1	In an additional study conducted in 6 Italian cities, <u>Orazzo et al. (2009</u>) examined respiratory ED visits for ages 0-2 years in 6 Italian cities from 1996 to 2000. However,
3	instead of identifying respiratory ED visits using the traditional approach of selecting
4	ICD codes as was done by Tolbert et al. (2007) and Darrow et al. (2011a), Orazzo et al.
5	(2009) used data on wheeze extracted from medical records as an indicator of lower
6	respiratory disease. This study examined daily counts of wheeze in relation to air
7	pollution using a time-stratified case-crossover approach in which control days were
8	matched on day of week in the same month and year as the case day. The authors found
9	no evidence of an association between 8-h max O ₃ concentrations and respiratory ED
10	visits in children aged 0-2 years in models that examined both single-day lags and
11	moving averages of lags from 0-6 days in year-round and seasonal analyses (i.e., warm
12	and cool seasons). In all-year analyses, the percent increase in total wheeze ranged from -
13	1.4% to -3.3% for a 0-1 to 0-6 day lag, respectively.

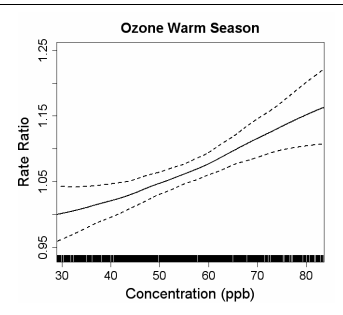
COPD

- 1 Stieb et al. (2009) also examined the association between short-term O_3 exposure and 2 COPD ED visits in 7 Canadian cities. Across cities, in an all-year analysis, O_3 was found 3 to be positively associated with COPD ED visits (2.4% [95% CI: -1.9, 6.9%] at lag 1 and 4 4.0% [95% CI: -0.54, 8.6%] at lag 2 for a 20 ppb increase in 24-h avg O_3 concentrations). 5 In seasonal analyses, larger effects were observed between O₃ and COPD ED visits 6 during the warm season (i.e., April-September) 6.8% [95% CI: 0.11, 13.9%] (lag day not 7 specified); with no associations observed in the winter season. Stieb et al. (2009) also 8 examined associations between respiratory-related ED visits, including COPD, and air 9 pollution at sub-daily time scales (i.e., 3-h avg of ED visits versus 3-h avg pollutant 10 concentrations) and found no evidence of consistent associations between any pollutant 11 and any respiratory outcome.
- 12 In a single-city study, Arbex et al. (2009) examined the association between COPD and 13 several ambient air pollutants, including O₃, in Sao Paulo, Brazil for the years 2001-2003 14 for individuals over the age of 40. Associations between O₃ exposure and COPD ED 15 visits were examined in both single-day lag (0-6 days) and polynomial distributed lag 16 models (0-6 days). In all-year analyses, O₃ was not found to be associated with an 17 increase in COPD ED visits (results not presented quantitatively). The authors also 18 conducted stratified analyses to examine the potential modification of the air pollutant-19 COPD ED visits relationship by age (e.g., 40-64, >64) and sex. In these analyses O₃ was 20 found to have an increase in COPD ED visits for women, but not for men or either of the 21 age groups examined.

Asthma

22 In a study of 7 Canadian cities, Stieb et al. (2009) also examined the association between 23 exposure to air pollution (i.e., CO, NO₂, O₃, SO₂, PM₁₀, PM_{2.5}, and O₃) and asthma ED 24 visits. Associations between short-term O_3 exposure and asthma ED visits were examined 25 at the city level and then pooled using either fixed or random effects models depending 26 on whether heterogeneity among effect estimates was found to be statistically significant. 27 Across cities, in an all-year analysis, the authors found that short-term O_3 exposure was 28 associated with an increase (4.7% [95% CI: -1.4, 11.1%] at lag 1 and 3.5% [95% CI: 29 0.33, 6.8%] at lag 2 for a 20 ppb increase in 24-h avg O₃ concentrations) in asthma ED 30 visits. The authors did not present the results from seasonal analyses for asthma, but 31 stated that no associations were observed between any pollutant and respiratory ED visits 32 in the winter season. As stated previously, in analyses of 3-h avg O_3 concentrations, the 33 authors observed no evidence of consistent associations between any pollutant and any 34 respiratory outcome, including asthma. A single-city study conducted in Alberta, Canada 35 Villeneuve et al. (2007) from 1992-2002 among individuals two years of age and older

- 1 provides additional support for the findings from Stieb et al. (2009), but also attempts to 2 identify those lifestages (i.e., 2-4, 5-14, 15-44, 45-64, 65-74, or 75+) at greatest risk to 3 O₃-induced asthma ED visits. In a time-referent case-crossover analysis, Villeneuve et al. 4 found an increase in asthma ED visits in an all-year analysis across all ages (12.0% 5 [95% CI: 6.8, 17.2] for a 30 ppb increase in max 8-h avg O₃ concentrations at lag 0-2) 6 with associations being stronger during the warmer months (19.0% [95% CI: 11.9, 28.1]). 7 When stratified by age, the strongest associations were observed in the warm season for 8 individuals 5-14 (28.1% [95% CI: 11.9, 45.1]; lag 0-2) and 15-44 (19.0% [95% CI: 8.5, 9 31.8]; lag 0-2). These associations were not found to be confounded by the inclusion of 10 aeroallergens in age-specific models. 11 Several additional single-city studies have also provided evidence of an association
- 12 between asthma ED visits and ambient O_3 concentrations. Ito et al. (2007b) examined the 13 association between short-term exposure to air pollution and asthma ED visits for all ages 14 in New York City from 1999 to 2002. Similar to Darrow et al. (2011a), when examining 15 the spatial distribution of O_3 concentrations, Ito et al. (2007b) found a rather homogenous 16 distribution (r $\sim \geq 0.80$) when examining monitor-to-monitor correlations at distances up to 17 20 miles. Ito et al. (2007b) used three different weather models with varying extent of 18 smoothing to account for temporal relationships and multicollinearity among pollutants 19 and meteorological variables (i.e., temperature and dew point) to examine the effect of 20 model selection on the air pollutant-asthma ED visit relationship. When examining O_3 , 21 the authors reported a positive association with asthma ED visits, during the warm season 22 across the models (ranging from 8.6 to 16.9%) and an inverse association in the cool 23 season (ranging from -23.4 to -25.1%), at lag 0-1 for a 30 ppb increase in 8-h max O_3 24 concentrations. Ito et al. (2007b) conducted copollutant models using a simplified version 25 of the weather model used in NMMAPS analyses (i.e., terms for same-day temperature 26 and 1-3 day average temperature). The authors found that O_3 risk estimates were not 27 substantially changed in copollutant models that used every-day data for PM_{2.5}, NO₂, 28 SO₂, and CO during the warm season (Figure 6-19; Table 6-29).



Note: The reference for the rate ratio is the estimated rate at the 5th percentile of the pollutant concentration. Estimates are presented for the 5th percentile through the 95th percentile of pollutant concentrations due to instability in the C-R estimates at the distribution tails.

Source: Reprinted with permission of American Thoracic Society (Strickland et al., 2010).

Figure 6-17 Loess C-R estimates and twice-standard error estimates from generalized additive models for associations between 8-h max 3-day average ozone concentrations and ED visits for pediatric asthma.

1	Strickland et al. (2010) examined the association between O_3 exposure and pediatric
2	asthma ED visits (ages 5-17 years) in Atlanta between 1993 and 2004 using air quality
3	data over the same years as Darrow et al. (2011a) and Tolbert et al. (2007). However,
4	unlike Darrow et al. (2011a) and Tolbert et al. (2007), which used single centrally located
5	monitors or an average of monitors, respectively, Strickland et al. (2010) used
6	population-weighting to combine daily pollutant concentrations across monitors. In this
7	study, the authors developed a statistical model using hospital-specific time-series data
8	that is essentially equivalent to a time-stratified case-crossover analysis (i.e., using
9	interaction terms between year, month, and day-of-week to mimic the approach of
10	selecting referent days within the same month and year as the case day). The authors
11	observed a 6.4% (95% CI: 3.2, 9.6%) increase in ED visits for a 30 ppb increase in
12	8-h max O ₃ concentrations at lag 0-2 in an all-year analysis. In seasonal analyses,
13	stronger associations were observed during the warm season (i.e., May-October) (8.4%
14	[95% CI: 4.4, 12.7%]; lag 0-2) than the cold season (4.5% [95% CI: -0.82, 10.0%]; lag 0-
15	2). Strickland et al. (2011) confirmed these findings in an additional analysis using the
16	same dataset, and found that the exposure assignment approach used (i.e., centrally

located monitor, unweighted average across monitors, and population-weighted average
 across monitors) did not influence pediatric asthma ED visit risk estimates for spatially
 homogeneous pollutants such as O₃.

4 In copollutant analyses conducted over the entire dataset for the gaseous pollutants 5 (i.e., (CO, NO₂), and limited to a subset of years (i.e., 1998-2004) for which daily PM 6 data (i.e., PM_{2.5} elemental carbon, PM_{2.5} sulfate) were available, Strickland et al. (2010) 7 found that O₃ risk estimates were not substantially changed when controlling for other 8 pollutants (results not presented quantitatively). The authors also examined the C-R 9 relationship between O3 exposure and pediatric asthma ED visits and found that both 10 quintile and loess C-R analyses (Figure 6-17) suggest that there are elevated associations 11 with O₃ at 8-h max concentrations as low as 30 ppb. These C-R analyses do not provide 12 evidence of a threshold level.

13 In a single-city study conducted on the West coast, Mar and Koenig (2009) examined the 14 association between O₃ exposure and asthma ED visits (ICD-9 codes: 493-493.9) for 15 children (<18) and adults (\geq 18) in Seattle, WA from 1998 to 2002. Of the total number 16 of visits over the study duration, 64% of visits in the age group <18 comprised boys, and 17 70% of visits in the > 18 age group comprised females. Mar and Koenig (2009) 18 conducted a time-series analysis using both 1-h max and max 8-h avg O_3 concentrations. 19 A similar magnitude and pattern of associations was observed at each lag examined using 20 both metrics. Mar and Koenig (2009) presented results for single day lags of 0 to 5 days, 21 but found consistent positive associations across individual lag days which supports the 22 findings from the studies discussed above that examined multi-day exposures. For 23 children, consistent positive associations were observed across all lags, ranging from a 24 19.1-36.8% increase in asthma ED visits for a 30 ppb increase in 8-h max O₃ 25 concentrations with the strongest associations observed at lag 0 (33.1% [95% CI: 3.0, 26 68.5]) and lag 3 (36.8% [95% CI: 6.1, 77.2]). O₃ was also found to be positively 27 associated with asthma ED visits for adults at all lags, ranging from 9.3-26.0%, except at 28 lag 0. The slightly different lag times for children and adults suggest that children may be 29 more immediately responsive to O_3 exposures than adults Mar and Koenig (2009).

Respiratory Infection

30Although an increasing number of studies have examined the association between O331exposure and cause-specific respiratory ED visits this trend has not included an extensive32examination of the association between O3 exposure and respiratory infection ED visits.33Stieb et al. (2009) also examined the association between short-term O3 exposure and34respiratory infection ED visits in 7 Canadian cities. In an all-year analysis, there was no35evidence of an association between O3 exposure and respiratory infection ED visits at any36lag examined (i.e., 0, 1, and 2). Across cities, respiratory infections comprised the single

largest diagnostic category, approximately 32%, of all the ED visits examined, which also included myocardial infarction, heart failure, dysrhythmia, asthma, and COPD.

6.2.7.4 Outpatient and Physician Visit Studies

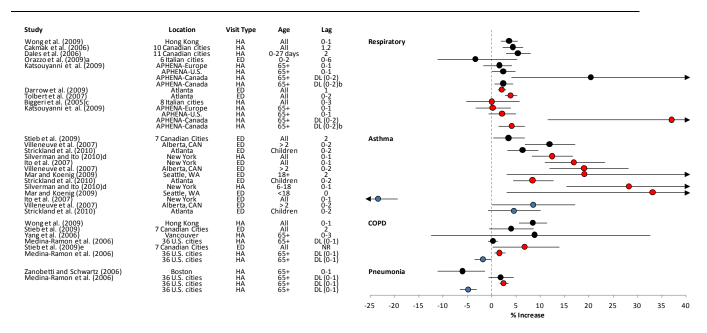
3	Several studies have examined the association between ambient O ₃ concentrations and
4	physician or outpatient (non-hospital, non-eD) visits for acute conditions in various
5	geographic locations. Burra et al. (2009) examined asthma physician visits among
6	patients aged 1-17 and 18-64 years in Toronto, Canada from 1992 to 2001. The authors
7	found little or no evidence of an association between asthma physician visits and O ₃ ;
8	however, seasonal analyses were not conducted. It should be noted that in this study,
9	most of the relative risks for O ₃ were less than one and statistically significant, perhaps
10	indicating an inverse correlation with another pollutant or an artifact of the strong
11	seasonality of asthma visits. Villeneuve et al. (2006b) also focused on physician visits to
12	examine the effect of short-term O3 exposure on allergic rhinitis among individuals aged
13	65 or older in Toronto from 1995 to 2000. The authors did not observe any evidence of
14	an association between allergic rhinitis physician visits and ambient O ₃ concentrations in
15	single-day lag models in an all-year analysis (results not presented quantitatively).
16	In a study conducted in Atlanta, Sinclair et al. (2010) examined the association of acute
17	asthma and respiratory infection (e.g., upper respiratory infections and lower respiratory
18	infections) outpatient visits from a managed care organization with ambient O_3
19	concentrations as well as multiple PM size fractions and species from August 1998
20	through December 2002. The authors separated the analysis into two time periods (the
21	first 25 months of the study period and the second 28 months of the study period), in
22	order to compare the air pollutant concentrations and relationships between air pollutants
23	and acute respiratory visits for the 25-month time-period examined in Sinclair and
24	Tolsma (2004) to an additional 28-month time-period of available data from the Atlanta
25	Aerosol Research Inhalation Epidemiology Study (ARIES). The authors found little
26	evidence of an association between O_3 and asthma visits, for either children or adults, or
27	respiratory infection visits in all-year analyses and seasonal analyses. For example, a
28	slightly elevated relative risk (RR) for childhood asthma visits was observed during the
29	25-month period in the cold season (RR: 1.12 [95% CI: 0.86, 1.41]; lag 0-2 for a 30 ppb
30	increase in 8-h max O ₃), but not in the warm season (RR: 0.97 [95% CI: 0.86, 1.10]; lag
31	0-2). During the 28-month period at lag 0-2, a slightly larger positive effect was observed
32	during the warm season (RR: 1.06 [95% CI: 0.97, 1.17]), compared to the cold season
33	(RR: 1.03 [95% CI: 0.87, 1.21]). Overall, these results contradict those from Strickland et
34	<u>al. (2010</u>) discussed above. Although the mean number of asthma visits and O_3
35	concentrations in Sinclair et al. (2010) and Strickland et al. (2010) are similar the

1 2 1difference in results between the two studies could potentially be attributed to the severity2of O3-induced asthma exacerbations (i.e., more severe symptoms requiring a visit to a3hospital) and behavior, such as delaying a visit to the doctor for less severe symptoms.

6.2.7.5 Summary

4	The results of the recent studies evaluated largely support the conclusion of the 2006 O_3
5	AQCD. While fewer studies were published overall since the previous review, several
6	multicity studies (e.g., Cakmak et al., 2006b; Dales et al., 2006) and a multi-continent
7	study (Katsouyanni et al., 2009) provide supporting evidence for an association between
8	short-term O3 exposure and an increase in respiratory-related hospital admissions and ED
9	visits. Across studies, different ICD-9 codes were used to define total respiratory causes,
10	which may contribute to some heterogeneity in the magnitude of association. These
11	findings are supported by single-city studies that used different exposure assignment
12	approaches (i.e., average of multiple monitors, single monitor, population-weighted
13	average) and averaging times (i.e., 1-h max and 8-h max).
14	Collectively, in both single-city and multicity studies there is continued evidence for
15	increases in both hospital admissions and ED visits when examining all respiratory
16	outcomes combined. Additionally, recent studies published since the 2006 O ₃ AQCD
17	support an association between short-term O_3 exposure and asthma (Strickland et al.,
18	2010; Stieb et al., 2009) and COPD (Stieb et al., 2009; Medina-Ramon et al., 2006)
19	hospital admissions and ED visits, with more limited evidence for pneumonia-hospital
20	admissions and ED visits (Medina-Ramon et al., 2006; Zanobetti and Schwartz, 2006).
21	As with total respiratory causes, studies used slightly different ICD-9 codes to define
22	specific conditions. In seasonal analyses, stronger associations were observed in the
23	warm season or summer months compared to the cold season, particularly for asthma
24	(Strickland et al., 2010; Ito et al., 2007b) and COPD (Medina-Ramon et al., 2006)
25	(Figure 6-18; Table 6-28), which is consistent with the conclusions of the 2006 O_3
26	AQCD. There is also continued evidence that children are particularly at greatest risk to
27	O3-induced respiratory effects (Silverman and Ito, 2010; Strickland et al., 2010; Mar and
28	Koenig, 2009; Villeneuve et al., 2007; Dales et al., 2006). Of note, the consistent
29	associations observed across studies for short-term O3 exposure and respiratory-related
30	hospital admissions and ED visits was not supported by studies that focused on
31	respiratory-related outpatient or physician visits. These differences could potentially be
32	attributed to the severity of O3-induced respiratory effects requiring more immediate
33	treatment or behavioral factors that result in delayed visits to a physician. Although the
34	collective evidence across studies indicates a consistent positive association between O_3
35	exposure and respiratory-related hospital admissions and ED visits, the magnitude of

1	these associations may be underestimated due to behavioral modification in response to
2	forecasted air quality (Neidell and Kinney, 2010; Neidell, 2009) (Section 4.6.5).
3	The studies that examined the potential confounding effects of copollutants found that O ₃
4	effect estimates remained relatively robust upon the inclusion of PM (measured using
5	different sampling strategies ranging from every-day to every-6th day) and gaseous
6	pollutants in two-pollutant models (Figure 6-19; Table 6-29). Additional studies that
7	conducted copollutant analyses, but did not present quantitative results, also support these
8	conclusions (Strickland et al., 2010; Tolbert et al., 2007; Medina-Ramon et al., 2006).
9	Overall, recent studies provide copollutant results that are consistent with the studies
10	evaluated in the 2006 O ₃ AQCD [(<u>U.S. EPA, 2006b</u>), Figure 7-12, page 7-80 of the 2006
11	O ₃ AQCD], which found that O ₃ respiratory hospital admissions risk estimates remained
12	robust to the inclusion of PM in copollutant models.



Note: Effect estimates are for a 20 ppb increase in 24-h; 30 ppb increase in 8-h max; and 40 ppb increase in 1-h max O_3 concentrations. HA=hospital admission; ED=emergency department. Black=All-year analysis; Red=Summer only analysis; Blue=Winter only analysis.

^aWheeze used as indicator of lower respiratory disease.

^b APHENA-Canada results standardized to approximate IQR of 5.1 ppb for 1-h max O₃ concentrations.

 $^{\circ}$ Study included 8 cities; but of those 8, only 4 had O₃ data.

^d non-ICU effect estimates.

^e The study did not specify the lag day of the summer season estimate.

Figure 6-18 Percent increase in respiratory-related hospital admission and ED visits in studies that presented all-year and/or seasonal results.

Study*	ED Visit or Hospital Admission	Location	Age	Lag	Avg Time	% Increase (95% CI)
Respiratory						
All-year						
Wong et al. (2009)	Hospital Admission	Hong Kong	All	0-1	8-h max	3.58 (1.90, 5.29)
Cakmak et al. (2006b)	Hospital Admission	10 Canadian cities	All	1.2	24-h avg	4.38 (2.19, 6.46)
Dales et al. (2006)	Hospital Admission	11 Canadian cities	0-27 days	2	24-h avg	5.41 (2.88, 7.96)
<u>Orazzo et al. (2009</u>) ^a	ED Visit	6 Italian cities	0-2	0-6	8-h max	-3.34 (-11.2, 5.28)
Katsouyanni et al. (2009)	Hospital	APHENA-europe	65+	0-1	1-h max	1.58 (-1.71, 4.15)
	Admission	APHENA-U.S.	65+	0-1	1-h max	2.38 (0.00, 4.89)
		APHENA-Canada	65+	DL(0-2)	1-h max	20.4 (4.07, 40.2)
		APHENA-Canada	65+	DL(0-2) ^b	1-h max	2.4 (0.51, 4.40)
Warm						
Darrow et al. (2011a)	ED Visit	Atlanta	All	1	8-h max	2.08 (1.25, 2.91)
<u>Tolbert et al. (2007)</u>	ED Visit	Atlanta	All	0-2	8-h max	3.90 (2.70, 5.20)
Biggeri et al. (2005) ^c Hospital Admission		8 Italian cities	All	0-3	8-h max	0.06 (-5.24, 5.66)
<u>Katsouyanni et al. (2009)</u>	Hospital	APHENA-europe	65+	0-1	1-h max	0.24 (-3.32, 3.91)
	Admission	APHENA-U.S.	65+	0-1	1-h max	2.14 (-0.63, 4.97)
		APHENA-Canada	65+	DL(0-2)	1-h max	37.1 (11.5, 67.5)
		APHENA-Canada	65+	DL(0-2) ^b	1-h max	4.1 (1.40, 6.80)
Asthma						
All-year						
<u>Stieb et al. (2009</u>)	ED Visit	7 Canadian cities	All	2	24-h avg	3.48 (0.33, 6.76)
Villeneuve et al. (2007)	ED Visit	Alberta, CAN	>2	0-2	8-h max	11.9 (6.8, 17.2)
Strickland et al. (2010)	ED Visit	Atlanta	Children	0-2	8-h max	6.38 (3.19, 9.57)
Warm						
Silverman and Ito (2010) ^d	Hospital Admission	New York	All	0-1	8-h max	12.5 (8.27, 16.7)
<u>lto et al. (2007b</u>)	ED Visit	New York	All	0-1	8-h max	16.9 (10.9, 23.4)
<u>Villeneuve et al. (2007)</u>	ED Visit	Alberta, CAN	>2	0-2	8-h max	19.0 (11.9, 28.1)
Mar and Koenig (2009)	ED Visit	Seattle, WA	18+	2	8-h max	19.1 (3.00, 40.5)
Strickland et al. (2010)	ED Visit	Atlanta	Children	0-2	8-h max	8.43 (4.42, 12.7)
Silverman and Ito (2010) ^d	Hospital Admission	New York	6-18	0-1	8-h max	28.2 (15.3, 41.5)
Mar and Koenig (2009)	ED Visit	Seattle, WA	<18	0	8-h max	33.1 (3.00, 68.5)

Table 6-28Corresponding Effect Estimates for Figure 6-18.

Study*	ED Visit or Hospital Admission	Location	Age	Lag	Avg Time	% Increase (95% CI)
Cold						
<u>Ito et al. (2007b</u>)	ED Visit	New York	All	0-1	8-h max	-23.4 (-27.3, -19.3)
Villeneuve et al. (2007)	ED Visit	Alberta, CAN	>2	0-2	8-h max	8.50 (0.00, 17.2)
Strickland et al. (2010)	ED Visit	Atlanta	Children	0-2	8-h max	4.52 (-0.82, 10.1)
COPD						
All-year						
<u>Stieb et al. (2009)</u>	ED Visit	7 Canadian cities	All	2	24-h avg	4.03 (-0.54, 8.62)
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	0.24 (-0.78, 1.21)
<u>Yang et al. (2005b)</u>	Hospital Admission	Vancouver	65+	0-3	24-h avg	8.80 (-12.5, 32.6)
Warm						
Stieb et al. (2009) ^e	ED Visit	7 Canadian cities	All	NR	24-h avg	6.76 (0.11, 13.9)
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	1.63 (0.48, 2.85)
Cold						
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	-1.85 (-3.60, -0.06)
Pneumonia						
All-year						
Zanobetti and Schwartz (2006)	Hospital Admission	Boston	65+	0-1	24-h avg	-5.96 (-11.1, -1.36)
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	1.81 (-0.72, 4.52)
Warm						
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	2.49 (1.57, 3.47)
Cold						
Medina-Ramon et al. (2006)	Hospital Admission	36 U.S. cities	65+	0-1	8-h max	-4.88 (-6.59, -3.14)

*Includes studies in Figure 6-18.

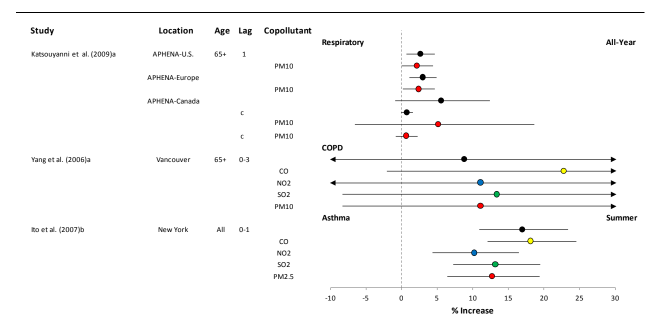
^aWheeze used as indicator of lower respiratory disease.

^bAPHENA-Canada results standardized to approximate IQR of 5.1 ppb for 1-h max O₃ concentrations.

 $^{\circ}Study$ included 8 cities, but of those 8 only 4 had O_{3} data.

^dNon-ICU effect estimates.

^eThe study did not specify the lag day of the summer season estimate.



Notes: Effect estimates are for a 20 ppb increase in 24 hours; 30 ppb increase in 8-h max; and 40 ppb increase in 1-h max O_3 concentrations.

^aStudies that examined hospital admissions,

^bA study that examined ED visits,

^cRisk estimates from APHENA -Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations. Black = results from single-pollutant models; Red = results from copollutant models with PM₁₀ or PM_{2.5}; Yellow = results from copollutant models with CO; Blue = results from copollutant models with NO₂; Green = results from copollutant models with SO₂.

Figure 6-19 Percent increase in respiratory-related hospital admissions and ED visits for studies that presented single and copollutant model results.

Study ^{*,a}	Location	Visit Type	Age	Lag	Copollutant	% Increase (95% CI)
All-year: Res	spiratory					
	APHENA-U.S.	Hospital	65+	1		2.62 (0.63, 4.64)
<u>et al. (2009</u>)		Admission			PM ₁₀	2.14 (-0.08, 4.40)
	APHENA-europe					2.94 (1.02, 4.89)
					PM ₁₀	2.38 (0.08, 4.64)
	APHENA-					5.54 (-0.94, 12.4)
	Canada					0.69 (-0.12, 1.50)b
					PM ₁₀	5.13 (-6.62, 18.6)
					PM ₁₀	0.64 (-0.87, 2.20)b
	COPD					
Yang et al.	Vancouver	Hospital Admission	65+	0-3		8.80 (-12.5, 32.6)
<u>(2005b</u>)					СО	22.8 (-2.14, 50.7)
					NO ₂	11.1 (-10.4, 37.6)
					SO ₂	13.4 (-8.40, 40.2)
					PM ₁₀	11.1 (-8.40, 37.6)
Summer: As	thma					
Ito et al.	New York	ED	All	0-1		16.9 (10.9, 23.4)
<u>(2007b</u>)				_	CO	18.1 (12.1, 24.5)
					NO ₂	10.2 (4.29, 16.4)
					SO ₂	13.1 (7.16, 19.5)
					PM _{2.5}	12.7 (6.37, 19.3)

Table 6-29 Corresponding effect estimates for Figure 6-19.

*Studies include in Figure 6-19.

^aAveraging times: <u>Katsouyanni et al. (2009</u>) = 1-h max; <u>Yang et al. (2005b</u>) = 24-h avg; and <u>Ito et al. (2007b</u>) = 8-h max. ^bRisk estimates standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations.

1 To date only a few studies have examined the C-R relationship between short-term O_3 2 exposure and respiratory-related hospital admissions and ED visits. A preliminary 3 examination of the C-R relationship found no evidence of a deviation from linearity when 4 examining the association between short-term O₃ exposure and asthma hospital 5 admissions (Silverman and Ito, 2010). Additionally, an examination of the C-R 6 relationship for O₃ exposure and pediatric asthma ED visits found no evidence of a 7 threshold with elevated associations with O_3 at concentrations as low as 30 ppb 8 (Silverman and Ito, 2010; Strickland et al., 2010). However, in both studies there is 9 uncertainty in the shape of the C-R curve at the lower end of the distribution of O₃ 10 concentrations due to the low density of data in this range.

1	In totality, building upon the conclusions of the 2006 AQCD, the evidence from recent
2	studies continues to support an association between short-term O ₃ exposure and
3	respiratory-related hospital admissions and ED visits. Additional evidence also supports
4	stronger associations during the warm season for specific respiratory outcomes such as
5	asthma and COPD.

6.2.8 Respiratory Mortality

- 6 The epidemiologic, controlled human exposure, and toxicological studies discussed 7 within this section (Section 6.2) provides evidence for multiple respiratory effects in 8 response to short-term O_3 exposure. Additionally, the evidence from experimental studies 9 indicates multiple potential pathways of O₃-induced respiratory effects, which support the 10 continuum of respiratory effects that could potentially result in respiratory-related 11 mortality. The 2006 O₃ AQCD found inconsistent evidence for an association between 12 short-term O₃ exposure and respiratory mortality (U.S. EPA, 2006b). Although some 13 studies reported a strong positive association between O₃ exposure and respiratory 14 mortality, additional studies reported a small association or no association. The majority 15 of recent multicity studies found consistent positive associations between short-term O₃ 16 exposure and respiratory mortality, specifically during the summer months.
- 17 The APHENA study, described earlier in Section 6.2.7.2, (Katsouyanni et al., 2009) also 18 examined associations between short-term O₃ exposure and mortality and found 19 consistent positive associations for respiratory mortality in all-year analyses, except in the 20 Canadian data set for ages \geq 75, with an increase in the magnitude of associations in 21 analyses restricted to the summer season across data sets and age ranges. Additional 22 multicity studies from the U.S. (Zanobetti and Schwartz, 2008b), Europe (Samoli et al., 23 2009), Italy (Stafoggia et al., 2010), and Asia (Wong et al., 2010) that conducted summer 24 season and/or all-year analyses provide additional support for an association between 25 short-term O_3 exposure and respiratory mortality (Figure 6-36).
- 26 Of the studies evaluated, only the APHENA study (Katsouyanni et al., 2009) and the Italian multicity study (Stafoggia et al., 2010) conducted an analysis of the potential for 27 28 copollutant confounding of the O₃-respiratory mortality relationship. In the APHENA 29 study, specifically the European dataset, focused on the natural spline model with 30 8 df/year (as discussed in Section 6.2.7.2) and lag 1 results (as discussed in 31 Section 6.6.2.1), respiratory mortality risk estimates were robust to the inclusion of PM_{10} 32 in copollutant models in all-year analyses with O₃ respiratory mortality risk estimates 33 increasing in the Canadian and U.S. datasets compared to single-pollutant model results. 34 In summer season analyses, respiratory O_3 mortality risk estimates were robust in the

1 U.S. dataset and attenuated in the European dataset. Similarly, in the Italian multicity 2 study (Stafoggia et al., 2010), which was limited to the summer season, respiratory 3 mortality risk estimates were attenuated in copollutant models with PM₁₀. Based on the 4 APHENA and Italian multicity results, O₃ respiratory mortality risk estimates appear to 5 be moderately to substantially sensitive (e.g., increased or attenuated) to inclusion of 6 PM₁₀. However, in the APHENA study, the mostly every-6th-day sampling schedule for 7 PM₁₀ in the Canadian and U.S. datasets greatly reduced their sample size and limits the 8 interpretation of these results.

6.2.9 Summary and Causal Determination

9	The 2006 O_3 AQCD concluded that there was clear, consistent evidence of a causal
10	relationship between short-term O_3 exposure and respiratory effects (U.S. EPA, 2006b).
11	This conclusion was substantiated by evidence from controlled human exposure and
12	toxicological studies indicating a range of respiratory effects in response to short-term O ₃
13	exposure, including pulmonary function decrements and increases in respiratory
14	symptoms, lung inflammation, lung permeability, and airway hyperresponsiveness.
15	Toxicological studies provided additional evidence for O ₃ -induced impairment of host
16	defenses. Combined, these findings from experimental studies provided support for
17	epidemiologic evidence, in which short-term increases in O ₃ concentration were
18	consistently associated with decreases in lung function in populations with increased
19	outdoor exposures, children with asthma, and healthy children; increases in respiratory
20	symptoms and asthma medication use in children with asthma; and increases in
21	respiratory-related hospital admissions and asthma-related ED visits. Short-term
22	increases in ambient O3 concentration also were consistently associated with increases in
23	all-cause and cardiopulmonary mortality; however, the contribution of respiratory causes
24	to these findings was uncertain.
25	Building on the large body of evidence presented in the 2006 O ₃ AQCD, recent studies
26	support associations between short-term O3 exposure and respiratory effects. Controlled
27	human exposure studies continue to provide the strongest evidence for lung function
28	decrements in young healthy adults over a range of O ₃ concentrations. Studies previously
29	reported mean O_3 -induced FEV ₁ decrements of 6-8% at 80 ppb O_3 (Adams, 2006a,
30	2003a; McDonnell et al., 1991; Horstman et al., 1990), and recent evidence additionally
31	indicates mean FEV ₁ decrements of 6% at 70 ppb O_3 (Schelegle et al., 2009) and 2-3% at
32	60 ppb O ₃ (Kim et al., 2011; Brown et al., 2008; Adams, 2006a) (Section 6.2.1.1). In
33	healthy young adults, O_3 -induced decrements in FEV ₁ do not appear to depend on sex
34	(Hazucha et al., 2003), body surface area or height (McDonnell et al., 1997), lung size or

baseline FVC (Messineo and Adams, 1990). There is limited evidence that blacks may

35

1	experience greater O_3 -induced decrements in FEV ₁ than do age-matched whites (Que et
2	al., 2011; Seal et al., 1993). Healthy children experience similar spirometric responses
3	but lesser symptoms from O_3 exposure relative to young adults (McDonnell et al.,
4	<u>1985a</u>). On average, spirometric and symptom responses to O_3 exposure appear to decline
5	with increasing age beyond about 18 years of age (McDonnell et al., 1999b; Seal et al.,
6	<u>1996</u>). There is also a tendency for slightly increased spirometric responses in mild
7	asthmatics and allergic rhinitics relative to healthy young adults (Jorres et al., 1996).
8	Spirometric responses in asthmatics appear to be affected by baseline lung function,
9	i.e., responses increase with disease severity (Horstman et al., 1995).
10	Available information from controlled human exposure studies on recovery from O ₃
11	exposure indicates that an initial phase of recovery in healthy individuals proceeds
12	relatively rapidly, with acute spirometric and symptom responses resolving within about
13	2 to 4 hours (Folinsbee and Hazucha, 1989). Small residual lung function effects are
14	almost completely resolved within 24 h. Effects of O3 on the small airways persisting
15	a day following exposure, assessed by persistent decrement in $\text{FEF}_{25-75\%}$ and altered
16	ventilation distribution, may be due in part to inflammation (Frank et al., 2001; Foster et
17	al., 1997). In more responsive individuals, this recovery in lung function takes longer (as
18	much as 48 hours) to return to baseline. Some cellular responses may not return to
19	baseline levels in humans for more than 10-20 days following O_3 exposure (Devlin et al.,
20	<u>1997</u>). Airway hyperresponsiveness and increased epithelial permeability are also
21	observed as late as 24 hours postexposure (Que et al., 2011).
22	With repeated O ₃ exposures over several days, spirometric and symptom responses
23	become attenuated in both healthy individuals and asthmatics, but this attenuation is lost
24	after about a week without exposure (Gong et al., 1997a; Folinsbee et al., 1994; Kulle et
25	al., 1982). Airway responsiveness also appears to be somewhat attenuated with repeated
26	O ₃ exposures in healthy individuals, but becomes increased in individuals with
27	preexisting allergic airway disease (Gong et al., 1997a; Folinsbee et al., 1994). Some
28	indicators of pulmonary inflammation are attenuated with repeated O ₃ exposures.
29	However, other markers such as epithelial integrity and damage do not show attenuation,
30	suggesting continued tissue damage during repeated O_3 exposure (<u>Devlin et al., 1997</u>).
31	Consistent with controlled human exposure study findings, epidemiologic evidence
32	indicates that lung function decrements are related to short-term increases in ambient O_3
33	concentration (Section $6.2.1.2$). As described in the 1996 and 2006 O ₃ AQCDs, the most
34	consistent observations were those in populations engaged in outdoor recreation,
35	exercise, or work. Epidemiologic evidence also demonstrates that increases in ambient O_3
36	concentration are associated with decreases in lung function in children with asthma
37	(Figure 6-6 and Figure 6-7 and Table 6-8 and Table 6-9) and children without asthma

1	(Figure 6-8 and Table 6-12). Evidence in adults with respiratory disease and healthy
2	adults is inconsistent. In children with asthma, lung function mostly was found to
3	decrease by $<1-2\%$ per unit increase in O ₃ concentration ¹ . However, in children with
4	asthma, O_3 -associated lung function decrements were found in conjunction with O_3 -
5	assocated increases in respiratory symptoms (Just et al., 2002; Mortimer et al., 2002;
6	Ross et al., 2002; Gielen et al., 1997; Romieu et al., 1997; Thurston et al., 1997; Romieu
7	et al., 1996). Biological plausibility for O_3 -associated decrements in lung function in
8	controlled human exposure, epidemiologic, and animal studies is provided by the well-
9	documented effects of O_3 on activation of bronchial C-fibers (Section <u>5.3.2</u>).
9	documented effects of O_3 on activation of biolicinal C-fibers (Section $5.5.2$).
10	Across disciplines, studies have examined factors that may potentially increase the risk of
11	O ₃ -induced decrements in lung function. In the controlled human exposure studies, there
12	is a large degree of intersubject variability in lung function decrements, symptomatic
13	responses, pulmonary inflammation, airway hyperresponsiveness, and altered epithelial
14	permeability in healthy adults exposed to O ₃ (Que et al., 2011; Holz et al., 2005;
15	McDonnell, 1996). The magnitude of pulmonary inflammation, airway
16	hyperresponsiveness, and increases in epithelial permeability do not appear to be
17	correlated, nor are these responses to O_3 correlated with changes in lung function,
18	suggesting that different mechanisms may be responsible for these processes (Que et al.,
19	2011; Balmes et al., 1997; Balmes et al., 1996; Aris et al., 1995). However, these
20	responses tend to be reproducible within a given individual over a period of several
21	months indicating differences in the intrinsic responsiveness of individuals (Holz et al.,
22	2005; Hazucha et al., 2003; Holz et al., 1999; McDonnell et al., 1985b). Numerous
23	reasons for differences in the risk of individuals to O ₃ exposure have been reported in the
24	literature. These include dosimetric and mechanistic differences (Section 5.4). Further,
25	evidence in all three disciplines suggests a role for antioxidant defenses (i.e., vitamin
26	supplementation, genetic variants in oxidative metabolizing enzymes) in modulating
27	respiratory responses to O ₃ . The biological plausibility of these findings is provided by
28	the well-characterized evidence for O ₃ exposure leading to the formation of secondary
29	oxidation products that subsequently activate neural reflexes that mediate lung function
30	decrements (Section $5.3.2$) and that initiate pulmonary inflammation (Section $5.3.3$).
21	
31	Recent controlled human exposure studies (Section <u>6.2.3.1</u>) and toxicological studies
32	(Section $\underline{6.2.3.3}$) also continue to demonstrate lung injury and inflammatory responses
33	upon O_3 exposure. Evidence from more than a hundred toxicological studies clearly
34	indicates that O_3 induces damage and inflammation in the lung, and studies continue to
35	elucidate the mechanistic pathways involved (Section 5.3). Though inflammation may
36	resolve, continued cellular damage may alter the structure and function of pulmonary

¹ Effect estimates were standardized to a 40-ppb increase for 1-h max O_3 , a 30-ppb increase for 8-h max O_3 , and a 20-ppb increase for 24-h avg O_3 .

1	tissues. Recent controlled human studies support previous findings for pulmonary
2	inflammation but demonstrate effects at 60 ppb O ₃ , the lowest concentration evaluated.
3	Building on the extensive experimental evidence, recent epidemiologic studies, most of
4	which were conducted in Mexico City, indicate ambient O3-associated increases in
5	pulmonary inflammation in children with asthma. Multiple studies examined and found
6	increases in eNO (Berhane et al., 2011; Khatri et al., 2009; Barraza-Villarreal et al.,
7	<u>2008</u>). In subjects with asthma, these O_3 -associated increases in pulmonary inflammation
8	were found concomitantly with O3-associated increases in respiratory symptoms (Khatri
9	et al., 2009; Barraza-Villarreal et al., 2008). Although more limited in number,
10	epidemiologic studies also found associations with cytokines such as IL-6 or IL-8
11	(Barraza-Villarreal et al., 2008; Sienra-Monge et al., 2004), eosinophils (Khatri et al.,
12	2009), antioxidants (Sienra-Monge et al., 2004), and indicators of oxidative stress
13	(<u>Romieu et al., 2008</u>) (Section $6.2.3.2$). This epidemiologic evidence is coherent with
14	results from controlled human exposure and toxicological studies that demonstrated an
15	induction or reduction of these same endpoints after O ₃ exposure.
16	The evidence for O ₃ -induced pulmonary inflammation and airway hyperresponsiveness,
17	largely demonstrated in controlled human exposure and toxicological studies, provides
18	mechanistic support for O3-associated increases in respiratory symptoms observed in both
19	controlled human exposure and epidemiologic studies. Controlled human exposure
20	studies of healthy, young adults demonstrate increases in respiratory symptoms induced
21	by O_3 exposures <80 ppb (<u>Schelegle et al., 2009</u> ; <u>Adams, 2006a</u>) (Section <u>6.2.1.1</u>).
22	Adding to this evidence, epidemiologic studies find effects in children with asthma.
23	Although the epidemiologic evidence was less consistent in the few available U.S.
24	multicity studies (O'Connor et al., 2008; Schildcrout et al., 2006; Mortimer et al., 2002),
25	the weight of evidence, provided by a larger body of single-city and -region studies,
26	indicates that short-term increases in ambient O ₃ concentration are associated with
27	increases in respiratory symptoms and asthma medication use in children with asthma
28	(Section <u>6.2.4.1</u>). Several epidemiologic studies found associations between ambient O_3
29	concentrations and respiratory symptoms in populations with asthma that also had a high
30	prevalence of allergy (52-100%) (Escamilla-Nuñez et al., 2008; Feo Brito et al., 2007;
31	Romieu et al., 2006; Just et al., 2002; Mortimer et al., 2002; Ross et al., 2002; Gielen et
32	al., 1997). The strong evidence in populations with asthma and allergy is supported by
33	observations of O_3 -induced inflammation in animal models of allergy (Section <u>6.2.3.3</u>),
34	and may be explained mechanistically by the action of O ₃ to sensitize bronchial smooth
35	muscle to hyperreactivity and thus, potentially act as a primer for subsequent exposure to
36	antigens such as allergens (Section $5.3.5$).
37	Modification of innate and adaptive immunity is emerging as a mechanistic pathway
38	contributing to the effects of O_3 on asthma and allergic airways disease (Section <u>5.3.6</u>).

1	While the majority of evidence comes from animal studies, controlled human exposure
2	studies have found differences between asthmatics and healthy controls in O ₃ -mediated
3	innate and adaptive immune responses (Section $5.4.2.2$), suggesting that these pathways
4	may be relevant to humans and may lead to the induction and exacerbation of asthma
5	(<u>Alexis et al., 2010; Hernandez et al., 2010; Alexis et al., 2009; Bosson et al., 2003</u>).
6	The subclinical and overt respiratory effects observed across disciplines, as described
7	above, collectively provide support for epidemiologic studies that demonstrate
8	consistently positive associations between short-term O ₃ exposure and respiratory-related
9	hospital admissions and ED visits (Section $6.2.7$). Consistent with evidence presented in
10	the 2006 O ₃ AQCD, recent multicity studies and a multicontinent study (i.e., APHENA)
11	(Katsouyanni et al., 2009) found risk estimates ranging from an approximate 1.6 to 5.4%
12	increase in all respiratory-related hospital admissions and ED visits in all-year analyses
13	for a unit increase in ambient O_3 concentration (as described in Section 2.1). Positive
14	associations persisted in analyses restricted to the summer season, but the magnitude
15	varied depending on the study location (Figure 6-18). Compared with studies reviewed in
16	the 2006 O ₃ AQCD, a larger number of recent studies examined hospital admissions and
17	ED visits for specific respiratory outcomes. Although limited in number, both single- and
18	multi-city studies found consistent, positive associations between short-term O ₃
19	exposures and asthma and COPD hospital admissions and ED visits, with more limited
20	evidence for pneumonia. Consistent with the conclusions of the 2006 O_3 AQCD, in
21	studies that conducted seasonal analyses, risk estimates were elevated in the warm season
22	compared to cold season or all-season analyses, specifically for asthma and COPD.
23	Although recent studies did not include detailed age-stratified results, the increased risk
24	of asthma hospital admissions (Silverman and Ito, 2010; Strickland et al., 2010; Dales et
25	<u>al., 2006</u>) observed for children strengthens the conclusion from the 2006 O_3 AQCD that
26	children are potentially at increased risk of O_3 -induced respiratory effects (U.S. EPA,
27	2006b). Although the C-R relationship has not been extensively examined, preliminary
28	examinations found no evidence of a threshold between short-term O_3 exposure and
29	asthma hospital admissions and pediatric asthma ED visits, with uncertainty in the shape
30	of the C-R curve at the lower limit of ambient concentrations in the U.S. (<u>Silverman and</u>
31	<u>Ito, 2010; Strickland et al., 2010</u>).
32	Recent evidence extends the potential range of well-established O ₃ -associated respiratory
33	effects by demonstrating associations between short-term ambient O_3 -associated respiratory
34	respiratory-related mortality. In all-year analyses, a multicontinent (APHENA) and
35	multicity (PAPA) study found consistent, positive associations with respiratory mortality
36	
	with evidence of an increase in the magnitude of associations in analyses restricted to the
37	summer months. Further, additional multicity studies conducted in the U.S. and Europe

provide evidence supporting stronger O_3 -respiratory mortality associations during the summer season (Section <u>6.2.8</u>).

- 3 Several studies of respiratory morbidity and mortality evaluated the potential confounding effects of copollutants, in particular, PM₁₀, PM_{2.5}, or NO₂. In most cases, 4 5 effect estimates remained robust to the inclusion of copollutants. In some studies of lung 6 function and respiratory symptoms, larger effects were estimated for O₃ when 7 copollutants were added to models. Ozone effect estimates for respiratory-related hospital 8 admissions and ED visits remained relatively robust upon the inclusion of PM and 9 gaseous pollutants in two-pollutant models (Strickland et al., 2010; Tolbert et al., 2007; 10 Medina-Ramon et al., 2006). Although copollutant confounding was not extensively 11 examined in studies of cause-specific mortality, O₃-respiratory mortality risk estimates 12 remained positive but were moderately to substantially sensitive (e.g., increased or 13 attenuated) to the inclusion of PM_{10} in copollutant models (Stafoggia et al., 2010; 14 Katsouyanni et al., 2009). However, interpretation of these results requires caution due to 15 the limited PM datasets used in these studies as a result of the every 3rd- or 6th-day PM 16 sampling schedule employed in most cities. Together, these copollutant-adjusted findings 17 across respiratory endpoints provide support for the independent effects of short-term 18 exposures to ambient O_3 .
- 19 Across the respiratory endpoints examined in epidemiologic studies, associations were 20 found using several different exposure assessment methods that likely vary in how well 21 ambient O_3 concentrations represent ambient exposures and between-subject variability 22 in exposures. Evidence clearly demonstrated O_3 -associated lung function decrements in 23 populations with increased outdoor exposures for whom ambient O_3 concentrations 24 measured on site of outdoor activity and/or at the time of outdoor activity have been more 25 highly correlated and similar in magnitude to personal O_3 exposures (Section 4.3.3). 26 However, associations with respiratory effects also were found with ambient O₃ 27 concentrations expected to have weaker personal-ambient relationships, including those 28 measured at home or school, measured at the closest site, averaged from multiple 29 community sites, and measured at a single site. Overall, there was no clear indication that 30 a particular method of exposure assessment produced stronger findings.
- 31An additional consideration in the evaluation of the epidemiologic evidence is the impact32of behavioral modifications on observed associations. A study demonstrated that the33magnitude of O_3 -associated asthma hospitalizations in Los Angeles, CA was34underestimated due to behavioral modification in response to forecasted air quality35(Section 4.6.5). It is important to note that the study was limited to one metropolitan area36and used air quality data for the years 1989-1997, when the O_3 concentration that37determines the designation of an O_3 action day, was much higher than it is currently.

1

2

1	Both panel and time-series epidemiologic studies found increases in respiratory effects in
2	association with increases in O3 concentrations using various exposure metrics
3	(i.e., 24-h avg, 1-h max, and 8-h max O ₃ concentrations). However, for respiratory
4	symptoms and pulmonary inflammation, a majority of studies examined and found
5	associations with 1-h max or 8-h max and 8-h max or daytime avg O ₃ , respectively.
6	Within study comparisons of associations among various exposure metrics with lung
7	function and respiratory symptoms yielded mixed evidence. Within some studies, larger
8	effects were estimated for shorter O3 averaging times whereas in other studies, larger
9	effects were estimated for longer averaging times or no difference was found among
10	averaging times. Comparisons in a limited number of time-series studies indicate rather
11	comparable risk estimates across exposure metrics with some evidence indicating that
12	24-h avg O_3 was associated with a smaller increase in risk of respiratory ED visits
13	(Section $6.2.7.3$). Overall, there was no indication that the consistency or magnitude of
14	the observed association was stronger for a particular O ₃ exposure metric. In examination
15	of the lag structure of associations, the weight of epidemiologic evidence for the range of
16	respiratory endpoints supports associations with ambient O3 concentrations lagged 0 to
17	1 day, which is consistent with the O ₃ -induced respiratory effects observed in controlled
18	human exposure studies. Several studies also found increased respiratory morbidity in
19	association with O_3 concentrations averaged over multiple days (2 to 5 days). Across
20	respiratory endpoints examined in epidemiologic studies, there was not strong evidence
21	that the magnitude of association was larger for any particular lag.

22 In summary, recent studies evaluated since the completion of the 2006 O₃ AQCD support 23 and expand upon the strong body of evidence that indicated a causal relationship between 24 short-term O_3 exposure and respiratory health effects. Controlled human exposure studies 25 continue to demonstrate O_3 -induced decreases in FEV₁ and pulmonary inflammation at 26 concentrations as low as 60 ppb. Epidemiologic studies provide evidence that increases in 27 ambient O_3 exposure can result in lung function decrements, increases in respiratory 28 symptoms, and pulmonary inflammation in children with asthma; increases in 29 respiratory-related hospital admissions and ED visits; and increases in respiratory 30 mortality. Recent toxicological studies demonstrating O₃-induced inflammation, airway 31 hyperresponsiveness, and impaired lung host defense have continued to support the 32 biological plausibility for the O₃-induced respiratory effects observed in the controlled 33 human exposure and epidemiologic studies. Additionally, recent epidemiologic studies 34 further confirm that respiratory morbidity and mortality associations are stronger during 35 the warm/summer months and remain relatively robust after adjustment for copollutants. 36 The recent evidence integrated across toxicological, controlled human exposure, and 37 epidemiologic studies, along with the total body of evidence evaluated in previous 38 AQCDs, is sufficient to conclude that there is a causal relationship between short-39 term O₃ exposure and respiratory health effects.

6.3 Cardiovascular Effects

1	Overall, there have been a relatively small number of studies that have examined the
2	potential effect of short-term O_3 exposure on the cardiovascular system. This was
3	reflected in the 1996 O ₃ AQCD by the limited discussion on possible O ₃ -related
4	cardiovascular effects. The 2006 O ₃ AQCD (<u>U.S. EPA, 2006b</u>) built upon the limited
5	evidence described in the 1996 O_3 AQCD and further explored the potential relationship
6	between short-term O_3 exposure and cardiovascular outcomes. The 2006 O_3 AQCD
7	concluded that "O3 directly and/or indirectly contributes to cardiovascular-related
8	morbidity" but added that the body of evidence was limited. This conclusion was based
9	on a controlled human exposure study that included hypertensive adult males, a few
10	epidemiologic studies of physiologic effects, heart rate variability, arrhythmias,
11	myocardial infarctions, and hospital admissions, and toxicological studies of heart rate,
12	heart rhythm, and blood pressure.

6.3.1 Controlled Human Exposure

13	O3 reacts rapidly on contact with respiratory system tissue and is not absorbed or
14	transported to extrapulmonary sites to any significant degree as such. Controlled human
15	exposure studies discussed in the previous AQCDs failed to demonstrate any consistent
16	extrapulmonary effects. Some controlled human exposure studies have attempted to
17	identify specific markers of exposure to O_3 in blood. Buckley et al. (1975) reported a
18	28% increase in serum α -tocopherol and a 26% increase in erythrocyte fragility in healthy
19	males immediately following exposure to 500 ppb O_3 for 2.75 hours with exercise
20	(unspecified activity level). However, in healthy adult males exposed during exercise
21	(\dot{V}_E =44 L/min) to 323 ppb O ₃ (on average) for 130 min on 3 consecutive days, <u>Foster et</u>
22	al. (1996) found a 12% reduction in serum α -tocopherol 20 hours after the third day of O_3
23	exposure. Liu et al. (1999); (1997) used a salicylate metabolite, 2,3, dehydroxybenzoic
24	acid (DHBA), to indicate increased levels of hydroxyl radical which hydroxylates
25	salicylate to DHBA. Increased DHBA levels after exposure to 120 and 400 ppb suggest
26	that O_3 increases production of hydroxyl radical. The levels of DHBA were correlated
27	with changes in spirometry. Interestingly, simultaneous exposure of healthy adults to O_3
28	(120 ppb for 2 hours at rest) and concentrated ambient particles (CAPs) resulted in a
29	diminished systemic IL-6 response compared with exposure to CAPs alone (Urch et al.,
30	<u>2010</u>).
31	Gong et al. (1998) exposed hypertensive ($n = 10$) and healthy ($n = 6$) adult males, 41 to
32	78 years of age, to FA and on the subsequent day to 300 ppb O_3 for 3 hours with
33	intermittent exercise ($\dot{V}_E = 30$ L/min). The overall results did not indicate any major acute

1 2 3 4 5 6 7 8 9 10 11	cardiovascular effects of O_3 in either the hypertensive individuals or healthy controls. Statistically significant O_3 effects for both groups combined were increases in heart rate, rate-pressure product, and the alveolar-to-arterial PO ₂ gradient, suggesting that impaired gas exchange was being compensated for by increased myocardial work. The mechanism for the decrease in arterial oxygen tension in the <u>Gong et al. (1998</u>) study could be due to an O ₃ -induced ventilation-perfusion mismatch. <u>Gong et al. (1998</u>) suggested that by impairing alveolar-arterial oxygen transfer, the O ₃ exposure could potentially lead to adverse cardiac events by decreasing oxygen supply to the myocardium. The subjects in the <u>Gong et al. (1998</u>) study had sufficient functional reserve so as to not experience significant ECG changes or myocardial ischemia and/or injury. In studies evaluating the exercise performance of healthy adults, no significant effect of O ₃ on arterial O ₂
12	saturation has been observed (Schelegle and Adams, 1986).
13 14	<u>Fakhri et al. (2009</u>) evaluated changes in HRV among adult volunteers (n = 50; 27 ± 7 years) during 2-hour exposures to PM _{2.5} CAPs ($127 \pm 62 \ \mu g/m^3$) and O ₃
15 16	$(114 \pm 7 \text{ ppb})$, alone and in combination. High frequency HRV was increased following
10	CAPs-only ($p = 0.046$) and O ₃ -only ($p = 0.051$) exposures, but not in combination. The standard deviation of NN intervals and the square root of the mean squared differences of
17	successive NN intervals also showed marginally significant ($0.05) increase due$
19	to O_3 but not CAPS. Ten of the subjects in this study were characterized as "mildly"
20	asthmatic, however, asthmatic status was not found to modify these effects. <u>Power et al.</u>
20	(2008) also investigated HRV in a small group of mild-to-moderate allergic asthmatics
22	(n = 5; mean age = 37 years) exposed for 4 hours during moderate intermittent exercise to
23	FA, carbon and ammonium nitrate particles $(313 \pm 20 \ \mu\text{g/m}^3)$, and carbon and ammonium
24	nitrate particles $(255 \pm 37 \ \mu g/m^3) + O_3$ (200 ppb). Changes in frequency-domain variables
25	for the particle and particle + O_3 exposures were not statistically significant compared
26	with FA. Seemingly in contrast to Fakhri et al. (2009), the standard deviation of NN
27	intervals and the square root of the mean squared differences of successive NN intervals
28	also showed a significant ($p = 0.01$) decrease for both the particle and particle + O_3
29	exposures relative to FA responses. Using a similar protocol, Sivagangabalan et al.
30	(2011) concluded that spatial dispersion of cardiac repolarization was most affected by
31	the combined pollutant exposure of $CAP + O_3$ compared to FA in healthy adults.
32 33 34 35 36 37 38	Diastolic blood pressure increased by 2 mmHg following the combined $O_3 + CAPs$ exposure, but was not altered by either O_3 or CAPs alone in the Fakhri et al. (2009) study. For a subset of the subjects without asthma in the Fakhri et al. (2009) study, Urch et al. (2005) previously reported a 6 mmHg increase in diastolic blood pressure following a 2-hour resting exposure to O_3 (120 ppb) + PM _{2.5} CAPs (150 µg/m ³) in healthy adults (n = 23; 32 ± 10 years), which was statistically different from the 1 mmHg increase seen following FA exposure. Brook et al. (2002) found O_3 (120 ppb) + PM _{2.5} CAPs (150

1	μ g/m ³) in healthy adults (n = 25; 35 ± 10 years) caused brachial artery vasoconstriction.
2	However, minimal change in diastolic blood pressure (0.9 mmHg increase) relative to FA
3	(0.4 mmHg decrease) was observed. More recently, Sivagangabalan et al. (2011)
4	observed reported a 4.2 mmHg increase in diastolic blood pressure following a 2-hour
5	resting exposure to O ₃ (110 ppb) + PM _{2.5} CAPs (150 μ g/m ³) in healthy adults (n = 25;
6	27 ± 8 years), which was statistically different from the 1.7 mmHg increase seen
7	following the FA exposure. The CAP exposure alone also caused a 3 mmHg increase in
8	diastolic blood pressure which was significantly more than following FA. However,
9	similar to FA, the O ₃ exposure alone caused a 1.8 mmHg increase in diastolic blood
10	pressure. Overall, these studies indicate an effect of CAPs and CAP + O_3 , but not O_3
11	alone, on diastolic blood pressure.

6.3.2 Epidemiology

12	The 2006 O ₃ AQCD concluded that the "generally limited body of evidence is highly
13	suggestive that O3 directly and/or indirectly contributes to cardiovascular-related
14	morbidity," including physiologic effects (e.g., release of platelet activating factor
15	[PAF]), HRV, arrhythmias, and myocardial infarctions, although the available body of
16	evidence reviewed during the 2006 O3 AQCD does not "fully substantiate links between
17	ambient O_3 exposure and adverse cardiovascular outcomes" (U.S. EPA, 2006b). Since
18	the completion of the 2006 O ₃ AQCD an increasing number of studies have examined the
19	relationship between short-term O_3 exposure and cardiovascular morbidity and mortality.
20	These recent studies, as well as evidence from the previous AQCDs, are presented within
21	this section.

6.3.2.1 Arrhythmia

22	In the 2006 O_3 AQCD, conflicting results were observed when examining the effect of O_3
23	on arrhythmias (Dockery et al., 2005; Rich et al., 2005). A study by Dockery et al. (2005)
24	reported no association between O3 concentration and ventricular arrhythmias among
25	patients with implantable cardioverter defibrillators (ICD) living in Boston, MA,
26	although when O_3 concentration was categorized into quintiles, there was weak evidence
27	of an association with increasing O ₃ concentration (median O ₃ concentration: 22.9 ppb).
28	Rich et al. (2005) performed a re-analysis of this cohort using a case-crossover design
29	and detected a positive association between O ₃ concentration and ventricular arrhythmias.
30	Recent studies were conducted in various locations and each used a different cardiac
31	episode to define an arrhythmic event and a different time period of exposure, which may

Table 6-30	Characterization of ozone concentrations (in ppb) from studies of
	arrhythmias.

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
Metzger et al. (2007)	Atlanta, GA	8-h max Summer only	53.9 (23)	Max: 148
<u>Rich et al. (2006b</u>)	Boston, MA	1-h	22.2*	75th: 33 Max: 119.5
		24-h	22.6*	75th: 30.9 Max: 77.5
Rich et al. (2006a)	St. Louis, MO	24-h	21*	75th: 31
Anderson et al. (2010)	London, England	8-h max	8.08	75th: 11.5
(<u>Sarnat et al., 2006b</u>)	Steubenville, OH	24-h Summer and Fall only	21.8 (12.6)	75th: 28.5 Max: 74.8
		5 days	22.2 (9.1)	75th: 29.1 Max: 44

Note: Median presented (information on mean not given); studies presented in order of first appearance in the text of this section.

3	Multiple studies examined O_3 -related effects on individuals with ICDs. A study of 518
4	ICD patients who had at least 1 tachyarrythmia within a 10-year period (totaling 6,287
5	tachyarrhythmic event-days; 1993-2002) was conducted in Atlanta, Georgia (Metzger et
6	al., 2007). Tachyarrhythmic events were defined as any ventricular tachyarrhythmic
7	event, any ventricular tachyarrhythmic event that resulted in electrical therapy, and any
8	ventricular tachyarrhythmic event that resulted in defibrillation. In the primary analysis,
9	no evidence of an association was observed for a 30 ppb increase in 8-h max O_3
10	concentrations and tachyarrhythmic events (OR: 1.00 [95% CI: 0.92, 1.08]; lag 0).
11	Season-specific as well as several sensitivity analyses (including the use of an
12	unconstrained distributed lag model [lags 0-6]) were conducted resulting in similar null
13	associations.
1.4	
14	In a case-crossover analysis, a population of ICD patients in Boston, previously examined
15	by (Rich et al., 2005) was used to assess the association between air pollution and
16	paroxysmal atrial fibrillation (PAF) episodes (Rich et al., 2006b). In addition to
17	ventricular arrhythmias, ICD devices may also detect supraventricular arrhythmias, of
18	which atrial fibrillation is the most common. Although atrial fibrillation is generally not
19	considered lethal, it has been associated with increased premature mortality as well as
20	hospitalization and stroke. Ninety-one electrophysiologist-confirmed episodes of PAF

- 1 were ascertained among 29 patients. An association (OR: 3.86 [95% CI: 1.44, 10.28] per 2 40 ppb increase in 1-h max O_3 concentrations) was observed between increases in O_3 3 concentration during the concurrent hour (lag 0-h) and PAF episodes. The estimated OR 4 for the 24-hour moving average concentration was elevated (OR: 1.81 [95% CI: 0.86, 5 3.83] per 20 ppb), but weaker than the estimate for the shorter exposure window. The 6 association between PAF and O₃ concentration in the concurrent hour during the cold 7 months was comparable to that during the warm months. In addition, no evidence of a 8 deviation from linearity between O₃ concentration and the log odds of PAF was observed. 9 Authors report that the difference between O₃ concentration and observed effect between 10 this study (PAF and 1-hour O_3) and their previous study (ventricular arrhythmias and 24-hour moving average O₃) (Rich et al., 2005) suggest a more rapid response to air 11 12 pollution for PAF (Rich et al., 2006b).
- 13 In an additional study, Rich et al. (2006a) employed a case-crossover design to examine 14 the association between air pollution and 139 confirmed ventricular arrhythmias among 15 56 ICD patients in St Louis, Missouri. The authors observed a positive association with 16 O₃ concentration (OR: 1.17 [95% CI: 0.58, 2.38] per 20 ppb increase in 24-hour moving 17 avg O_3 concentrations [lags 0-23 hours]). Although the authors concluded these results 18 were similar to their results from Boston (Rich et al., 2005), they postulated that the 19 pollutants responsible for the increased risk in ventricular arrhythmias are different (O_3 20 and PM_{2.5} in Boston and sulfur dioxide in St Louis).
- 21 Anderson et al. (2010) used a case-crossover framework to assess air pollution and 22 activation of ICDs among patients from all 9 ICD clinics in the London National Health 23 Service hospitals. "Activation" was defined as tachycardias for which the defibrillator 24 delivered treatment. Investigators modeled associations using unconstrained distributed 25 lags from 0 to 5 days. The sample consisted of 705 patients with 5,462 activation days 26 (O₃ concentration information was for 543 patients and 4,092 activation days). Estimates 27 for the association with O_3 concentration were consistently positive, although weak (OR: 28 1.09 [95% CI: 0.76, 1.55] per 30 ppb increase in 8-h max O₃ concentrations at 0-1 day 29 lag; OR: 1.04 [95% CI: 0.60, 1.81] per 30 ppb increase in 8-h max O₃ concentrations at 30 0-5 day lag) (Anderson et al., 2010).
- 31In contrast to arrhythmia studies conducted among ICD patients, Sarnat et al. (2006b)32recruited non-smoking adults (age range: 54-90 years) to participate in a study of air33pollution and arrhythmias conducted over two 12-week periods during summer and fall34of 2000 in a region characterized by industrial pollution (Steubenville, Ohio). Continuous35ECG data acquired on a weekly basis over a 30-minute sampling period were used to36assess ectopy, defined as extra cardiac depolarizations within the atria (supraventricular37ectopy, SVE) or the ventricles (ventricular ectopy, VE). Increases in the 5-day moving

1	average (days 1-5) of O_3 concentration were associated with an increased odds of SVE
2	(OR: 2.17 [95% CI: 0.93, 5.07] per 20 ppb increase in 24-h avg O ₃ concentrations). A
3	weaker association was observed for VE (OR: 1.62 [95% CI: 0.54, 4.90] per 20 ppb
4	increase in 24-h avg O_3 concentrations). The results of the effect of 5-day O_3
5	concentration on SVE were robust to the inclusion of SO_4^{2-} in the model [OR: 1.62
6	(95% CI: 0.54, 4.90)]. The authors indicate that the strong associations observed at the
7	5-day moving averages, as compared to shorter time periods, suggests a relatively long-
8	acting mechanistic pathways, such as inflammation, may have promoted the ectopic beats
9	in this population (Sarnat et al., 2006b).

10Although many studies report positive associations, collectively, studies of arrhythmias11report inconsistent results. This may be due to variation in study populations, length and12season of averaging time, and outcome under study.

6.3.2.2 Heart Rate/Heart Rate Variability

13	In the 2006 O ₃ AQCD, two large population-based studies of air pollution and HRV were
14	summarized (Park et al., 2005b; Liao et al., 2004a). In addition, the biological
15	mechanisms and potential importance of HRV were discussed. Briefly, the study of acute
16	effects of air pollution on cardiac autonomic control is based on the hypothesis that
17	increased air pollution levels may stimulate the autonomic nervous system and lead to an
18	imbalance of cardiac autonomic control characterized by sympathetic activation
19	unopposed by parasympathetic control (U.S. EPA, 2006b). Examples of HRV indices
20	include the standard deviation of normal-to-normal intervals (SDNN), the square root of
21	the mean of the sum of the squares of differences between adjacent NN intervals (r-
22	MSSD), high-frequency power (HF), low-frequency power (LF), and the LF/HF ratio.
23	Liao et al. (2004a) examined the association between air pollution and cardiac autonomic
24	control in the fourth cohort examination (1996-1998) of the U.Sbased Atherosclerosis
25	Risk in Communities Study. A decrease in log-transformed HF was associated with an
26	increase in O_3 concentration among white study participants. Park et al. (2005b)
27	examined the effects of air pollution on indices of HRV in a population-based study
28	among men from the Normative Aging Study in Boston, Massachusetts. Several
29	associations were observed with O_3 concentration and HRV outcomes. A reduction in LF
30	was associated with increased O_3 concentration, which was robust to inclusion of $PM_{2.5}$.
31	The associations with all HRV indices and O_3 concentration were stronger among those
32	with ischemic heart disease and hypertension. In addition to the population-based studies
33	included in the 2006 O_3 AQCD was a study by Schwartz et al. (2005), who conducted a
34	panel study to assess the relationship between exposure to summertime air pollution and
35	HRV. A weak association of O_3 concentration during the hour immediately preceding the

1	health measures was observed with r-MSSD among a study population that consisted of
2	mostly older female participants. In summary, these studies suggest that short-term
3	exposures to ambient O_3 concentrations are predictors of decreased HRV and that the
4	relationship may be stronger among certain subgroups. More recent studies that examined
5	the association between O_3 concentration and HRV are described below. Study-specific
6	characteristics and O_3 concentrations for these studies are presented in <u>Table 6-31</u> .

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
Park et al. (2007)	Boston, MA	24-h	Range of 17.0-29.1	
<u>Park et al. (2008</u>)	Boston, MA	24-h	23.4 (13)	
<u>Baja et al. (2010</u>)	Boston, MA	0 lag 10-h lag	23 (16) 21 (15)	
Wheeler et al. (2006)	Atlanta, GA	4-h 24-h	18.5 29.4	75th: 22.5
<u>Zanobetti et al. (2010)</u>	Boston, MA	0.5-h 2-h 3-D 5-D	20.7* 20.5* 21.9* 22.8*	75th: 30.33 75th: 30.08 75th: 28.33 75th: 29.28
<u>Chan et al. (2005a</u>)	Taipei, Taiwan	1-h	21.9 (15.4)	Max: 114.9
<u>Wu et al. (2010)</u>	Taipei, Taiwan	Working period	24.9 (14.0)	Max: 59.2
Ruidavets et al. (2005a)	Toulouse, France	8-h max	38.3 (14.8)	75th: 46.9 Max: 80.3
<u>Chuang et al. (2007a)</u>	Taipei, Taiwan	24-h 48-h 72-h	28.4 (12.1) 33.3 (8.9) 33.8 (7.1)	Max: 49.3 Max: 47.8 Max: 48.3
Chuang et al. (2007b)	Taipei, Taiwan	1-h	35.1	Max: 192.0

Table 6-31Characterization of ozone concentrations (in ppb) from studies of
heart rate variability.

*Note: Median presented (information on mean not given); studies presented in order of first appearance in the text of this section.

7 Several follow-up examinations of HRV were conducted among the participants of the 8 Normative Aging Study in Boston. A trajectory cluster analysis was used to assess 9 whether pollution originating from different locations had varying relationships with 10 HRV (Park et al., 2007). Subjects who were examined on days when air parcels 11 originated in the west had the strongest associations with O_3 ; however, the O_3 12 concentration in this cluster was low (24-h avg, 17.0 ppb) compared to the other clusters 13 (24-h avg of 21.3-29.1 ppb). LF and SDNN decreased with increases in the 4-hour 14 moving average of O_3 concentration from the west (LF decreased by 51.2% [95% CI: 1.6, 15 75.9%] and SDNN decreased by 28.2% [95% CI: -0.5, 48.7%] per 30 ppb increase in 16 4-h avg O₃ concentrations) (Park et al., 2007). The Boston air mass originating in the

1	west traveled over Illinois, Indiana, and Ohio; states typically characterized by coal-
2	burning power plants. Due to the low O ₃ concentrations observed in the west cluster, the
3	authors hypothesize that O_3 concentration on those days could be capturing the effects of
4	other, secondary and/or transported pollutants from the coal belt or that the relationship
5	between ambient O_3 concentration and personal exposure to O_3 is stronger during that
6	period (supported by a comparatively low apparent temperature which could indicate a
7	likelihood to keep windows open and reduced air conditioning use) (Park et al., 2007).
8	An additional follow-up evaluation using the Normative Aging Study examined the
9	potential for effect modification by chronic lead (Pb) exposure on the relationship
10	between air pollution and HRV (Park et al., 2008). Authors observed graded reductions in
11	HF and LF of HRV in relation to O ₃ (and sulfate) concentrations across increasing
12	quartiles of tibia and patella lead (HF: percent change 32.3% [95% CI: -32.5, 159.3] for
13	the first quartile of tibia Pb and -59.1 [95% CI: -77.3, -26.1] for the fourth quartile of
14	tibia Pb per 30 ppb increase in 4-h avg O ₃ concentrations; LF: percent change 8.0%
15	[95% CI: -36.9, 84.9] for the first quartile of tibia Pb and -59.3 [95% CI: -74.6, -34.8] for
16	the fourth quartile of tibia Pb per 30 ppb increase in 4-h avg O ₃ concentrations). In
17	addition, associations were similar when education and cumulative traffic-adjusted bone
18	Pb levels were used in analyses. Authors indicate the possibility that O ₃ (which has low
19	indoor concentrations) was acting as a proxy for sulfate (correlation coefficient for O ₃
20	and sulfate $= 0.57$). Investigators of a more recent follow-up to the Normative Aging
21	Study hypothesized that the relationships between short-term air pollution exposures and
22	ventricular repolarization, as measured by changes in the heart-rate corrected QT interval
23	(QTc), would be modified by participant characteristics (e.g., obesity, diabetes, smoking
24	history) and genetic susceptibility to oxidative stress (Baja et al., 2010). No evidence of
25	an association between O ₃ concentration (using a quadratic constrained distributed lag
26	model and hourly exposure lag models over a 10-hour time window preceding the visit)
27	and QTc was reported (change in mean QTc -0.74 [95% CI: -3.73, 2.25]); therefore,
28	potential effect modification of personal and genetic characteristics with O ₃ concentration
29	was not assessed (Baja et al., 2010). Collectively, the results from studies that examined
30	the Normative Aging Study cohort found an association between increases in short-term
31	O ₃ concentration and decreases in HRV (Park et al., 2008; Park et al., 2007; Park et al.,
32	2005b) although not consistently in all of the studies (Baja et al., 2010). Further, observed
33	relationships appear to be stronger among those with ischemic heart disease,
34	hypertension, and elevated bone lead levels, as well as when air masses arrive from the
35	west (the coal belt). However, it is not clear if O_3 concentration is acting as a proxy for
36	other, secondary particle pollutants (such as sulfate) (Park et al., 2008). In addition, since
37	the Normative Aging Study participants were older, predominately white men, results
38	may not be generalizable to the a large proportion of the U.S. population.

1	Additional studies of populations not limited to the Normative Aging Study have also
2	examined associations between O ₃ exposure and HRV. A panel study among 18
3	individuals with COPD and 12 individuals with recent myocardial infarction (MI) was
4	conducted in Atlanta, Georgia (Wheeler et al., 2006). HRV was assessed for each
5	participant on 7 days in fall 1999 and/or spring 2000. Ozone concentrations were not
6	associated with HRV (SDNN) among all subjects (percent change of 2.36% [95% CI:
7	-10.8%, 17.5%] per 30 ppb 4-hour O_3 increase) or when stratified by disease type
8	(COPD, recent MI, and baseline FEV_1) (Wheeler et al., 2006).
9	HRV and air pollution was assessed in a panel study among 46 predominately white male
10	patients (study population: 80.4% male, 93.5% white) aged 43-75 years in Boston,
11	Massachusetts, with coronary artery disease (Zanobetti et al., 2010). Up to four home
12	visits were made to assess HRV over the year following the index event. Pollution lags
13	used in analyses ranged between 30 minutes to a few hours and up to 5 days prior to the
14	HRV assessments, calculated from hourly O_3 measurements averaged over three
15	monitoring sites in Boston. Decreases in r-MSSD were reported for all averaging times of
16	O ₃ concentration (percent change of -5.18% [95% CI: -7.89, -2.30] per 20 ppb of 5-day
17	moving average of O_3 concentration), but no evidence of an association between O_3
18	concentration and HF was observed (quantitative results not provided). In two-pollutant
19	models with O_3 and either $PM_{2.5}$ or BC, O_3 associations remained robust.
20	A few recent studies were conducted outside of the U.S. that examined the relationship
20 21	A few recent studies were conducted outside of the U.S. that examined the relationship between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010</u> ; <u>Chuang et</u>
	*
21	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010;</u> <u>Chuang et</u>
21 22	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a</u>). No
21 22 23	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a</u>). No associations were reported between O_3 concentration and HRV among CHD patients and
21 22 23 24	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O_3 concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O_3
21 22 23 24 25	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010</u> ; <u>Chuang et al., 2007b</u> ; <u>Chuang et al., 2007a</u> ; <u>Chan et al., 2005a</u> ; <u>Ruidavets et al., 2005a</u>). No associations were reported between O_3 concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (<u>Chan et al.,</u> 2007).
21 22 23 24 25 26	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010</u> ; <u>Chuang et al., 2007b</u> ; <u>Chuang et al., 2007a</u> ; <u>Chan et al., 2005a</u> ; <u>Ruidavets et al., 2005a</u>). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (<u>Chan et al., 2005a</u>). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between
21 22 23 24 25 26 27	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI:
21 22 23 24 25 26 27 28	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (Wu et al., 2010). In addition,
21 22 23 24 25 26 27 28 29	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (Wu et al., 2010). In addition, no consistent relationships were identified between O ₃ concentration and resting heart
21 22 23 24 25 26 27 28 29 30	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (Wu et al., 2010). In addition, no consistent relationships were identified between O ₃ concentration and resting heart rate among middle-aged (35-64 years) participants residing in Toulouse, France
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21 22 23 24 25 26 27 28 29 30 31 32	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a</u>). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (<u>Chan et al., 2005a</u>). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (<u>Wu et al., 2010</u>). In addition, no consistent relationships were identified between O ₃ concentration and resting heart rate among middle-aged (35-64 years) participants residing in Toulouse, France (<u>Ruidavets et al., 2005a</u>). A negative trend was reported for the 3-day cumulative (lag days 1-3) concentration of 8-h max O ₃ with heart rate (p for trend = 0.02); however,
21 22 23 24 25 26 27 28 29 30 31 32 33	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (Wu et al., 2010). In addition, no consistent relationships were identified between O ₃ concentration and resting heart rate among middle-aged (35-64 years) participants residing in Toulouse, France (Ruidavets et al., 2005a). A negative trend was reported for the 3-day cumulative (lag days 1-3) concentration of 8-h max O ₃ with heart rate (p for trend = 0.02); however, the individual odds ratios comparing quintiles of exposure showed no association (OR for
21 22 23 24 25 26 27 28 29 30 31 32 33 34	between air pollution concentrations and heart rate and HRV (<u>Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a</u>). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (<u>Chan et al., 2005a</u>). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (<u>Wu et al., 2010</u>). In addition, no consistent relationships were identified between O ₃ concentration and resting heart rate among middle-aged (35-64 years) participants residing in Toulouse, France (<u>Ruidavets et al., 2005a</u>). A negative trend was reported for the 3-day cumulative (lag days 1-3) concentration of 8-h max O ₃ with heart rate (p for trend = 0.02); however,
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21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36	between air pollution concentrations and heart rate and HRV (Wu et al., 2010; Chuang et al., 2007b; Chuang et al., 2007a; Chan et al., 2005a; Ruidavets et al., 2005a). No associations were reported between O ₃ concentration and HRV among CHD patients and patients with one or more major CHD risk factors residing in Taipei, Taiwan (Chan et al., 2005a). Another study in Taipei, Taiwan examined mail carriers and reported O ₃ concentration measured using personal monitors. No association was observed between O ₃ concentration and the measures of HRV (percent change for SDNN: 0.57 [95% CI: -21.27, 28.46], r-MSSD: -7.10 [95% CI: -24.24, 13.92], HF: -1.92 [95% CI: -23.68, 26.02], LF: -4.82 [95% CI: -25.34, 21.35] per 40 ppb O ₃) (Wu et al., 2010). In addition, no consistent relationships were identified between O ₃ concentration and resting heart rate among middle-aged (35-64 years) participants residing in Toulouse, France (Ruidavets et al., 2005a). A negative trend was reported for the 3-day cumulative (lag days 1-3) concentration of 8-h max O ₃ with heart rate (p for trend = 0.02); however, the individual odds ratios comparing quintiles of exposure showed no association (OR for O ₃ concentraction of 0.93 [95% CI: 0.86, 1.01] for the highest quintile of resting heart rate compared to the lowest). When stratified by current smoking status, non-smokers had a

1	Taiwan to assess the relationship between air pollutants and inflammation, oxidative
2	stress, blood coagulation, and autonomic dysfunction (Chuang et al., 2007b; Chuang et
3	al., 2007a). Participants were apparently healthy college students (aged 18-25 year) who
4	were living in a university dormitory in metropolitan Taipei. Health endpoints were
5	measured three times from April to June in 2004 or 2005. Ozone concentration was
6	assessed in statistical models using the average of the 24, 48, and 72 hours before the
7	hour of each blood sampling. Decreases in HRV (measured as SDNN, r-MSSD, LF, and
8	HF) were associated with increases in O ₃ concentrations in single-pollutant models
9	(percent change for SDNN: -13.45 [95% CI: -16.26, -10.60], r-MSSD -13.76 [95% CI:
10	-21.62, -5.44], LF -9.16 [95% CI: -13.29, -4.95], HF -10.76 [95% CI: -18.88, -2.32] per
11	20 ppb cumulative 3-day avg O_3 concentrations) and remained associated with 3-day O_3
12	concentrations in two-pollutant models with sulfate. Another study in Taiwan recruited
13	individuals with CHD or at risk for cardiovascular disease from outpatient clinics during
14	the study period (two weeks in February) (Chuang et al., 2007b). No association was
15	observed between O ₃ concentration and HRV measures (SDNN, r-MSSD, LF, HF)
16	(numerical results not provided in publication).

17Overall, studies of O3 concentration and HRV report inconsistent results. Multiple studies18conducted in Boston observed positive associations but the authors of many of these19studies postulated that O3 concentration was possibly acting as a proxy for other20pollutants. The majority of other studies, both in the U.S. and internationally, report null21findings. The inconsistencies observed are further complicated by the different HRV22measures and averaging times used by the studies.

6.3.2.3 Stroke

23	The 2006 O ₃ AQCD did not identify any studies that examined the association between
24	short-term O3 exposure and stroke. However, recent studies have attempted to examine
25	this relationship. Lisabeth et al. (2008) used a time-series approach to assess the
26	relationship between daily counts of ischemic stroke and transient ischemic attack (TIA)
27	with O_3 concentrations in a southeast Texas community among residents 45 years and
28	older (2001-2005; median age of cases, 72 years). The median O_3 concentration (hourly
29	average per 24-hour time-period) was 25.6 ppb (IQR 18.1-33.8). The associations
30	between same-day O ₃ concentrations and stroke/TIA risk were positive (RR: 1.03
31	[95% CI: 0.96, 1.10] per 20 ppb increase in 24-h avg O ₃ concentrations) and previous-day
32	(RR: 1.05 [95% CI: 0.99, 1.12] per 20 ppb increase in 24-h avg O ₃ concentrations).
33	Associations were robust to adjustment for PM _{2.5} .

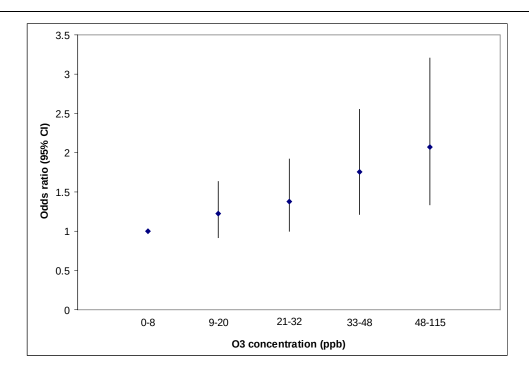
1	A case-crossover design was used in a study conducted in Dijon, France between March
2	1994 and December 2004, among those 40 years of age and older who presented with
3	first-ever stroke (<u>Henrotin et al., 2007</u>). The mean O_3 concentration, calculated over
4	8-hour daytime periods, was 14.95 ppb (IQR: 6-22 ppb). No association was observed
5	between O_3 concentration at any of the single-day lags examined (i.e., 0-3 days) and
6	hemorrhagic stroke. However, an association between ischemic stroke occurrence and O_3
7	concentrations with a 1-day lag was observed (OR: 1.54 [95% CI: 1.14, 2.09] per 30 ppb
8	increase in 8-h max O_3 concentrations). The observed association between short-term O_3
9	exposure and ischemic stroke persisted in two-pollutant models with PM ₁₀ , SO ₂ , NO ₂ , or
10	CO. This association was stronger among men (OR: 2.12 [95% CI: 1.36, 3.30] per 30 ppb
11	increase in 8-h max O ₃ concentrations) than among women (OR: 1.17 [95% CI: 0.77,
12	1.78] per 30 ppb increase in 8-h max O_3 concentrations) in single pollutant models. When
13	stroke was examined by subtype among men, an association was observed for ischemic
14	strokes of large arteries and for transient ischemic attacks, but not for cardioembolic or
15	lacunar ischemic strokes. The subtype analysis was not performed for women.
16	Additionally, for men a linear exposure-response was observed when O ₃ concentration
17	was assessed based on quintiles (p for trend = 0.01) (Figure 6-20). A potential limitation
18	of this study is that 67.4% of the participating men were smokers compared to 9.3% of
19	the women.
20	Another case-crossover study performed in Dijon, France examined the association
21	between O3 concentration and incidence of fatal and non-fatal ischemic cerebrovascular
22	events (ICVE) (Henrotin et al., 2010). Mean 8-hour O ₃ concentration was 19.1 ppb (SD
23	12.2 ppb). A positive association was observed between recurrent ICVE and 8-h O_3
24	concentration with a 3-day lag (OR: 1.92 [95% CI 1.17, 3.12]), but not for other lags (0,
25	1, 2, 4) or cumulative days (0-1, 0-2, 1-2, 1-3). Although some ORs for incident ICVEs
26	were elevated, none were statistically significant. Results for associations using the
27	maximum daily 1-hour O_3 concentrations were similar to the 8-hour results but slightly
28	attenuated. ORs were similar in two pollutant models with SO ₂ , NO ₂ , CO, and PM_{10} (data
29	not given). In stratified analyses, the association between 1-day lagged O_3 concentration

30

31

and incident and recurrent ICVE was greater among individuals with diabetes or

individuals with multiple preexisting vascular conditions.



Source: Henrotin et al. (2007).

Figure 6-20 Odds ratio (95% confidence interval) for ischemic stroke by quintiles of ozone exposure

6.3.2.4 Biomarkers

1	An increasing number of studies have examined the relationship between air pollution
2	and biomarkers in an attempt to elucidate the biological mechanisms linking air pollution
3	and cardiovascular disease. A wide range of markers assessed as well as different types
4	of study designs and locations chosen make comparisons across studies difficult.
5	<u>Table 6-32</u> provides an overview of the O_3 concentrations reported in each of the studies
6	evaluated.

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
<u>Liao et al. (2005</u>)	3 U.S. counties	8-h	40 (20)	
<u>Thompson et al.</u> (2010)	Toronto, Ontario	1-h / 1 yr	21.94 (15.78)	
Rudez et al. (2009)	Rotterdam, the Netherlands	24-h	22*	75th: 31.5
				Max: 90
Chuang et al.	Taipei, Taiwan	24-h	28.4 (12.1)	Max: 49.3
<u>(2007a</u>)		48-h	33.3 (8.9)	Max: 47.8
		72-h	33.8 (7.1)	Max: 48.3
Steinvil et al. (2008)	Tel-Aviv, Israel	0.5-h	29.2 (9.7)	75th: 36
Chen et al. (2007a)	Los Angeles and	8-h / 2 weeks	30.8*	Max: 47.9
	San Francisco, CA	8-h / 1 mo	28.3*	Max: 43.1
<u>Wellenius et al.</u> (2007)	Boston, MA	1-h / 24-h	25.1 (12.9)	
Goldberg et al. (2008)	Montreal, Quebec	24-h	NS	
Baccarelli et al.	Lombardia, Italy	1-h	18.3*	75th: 35.1
<u>(2007</u>)				Max: 202.3
<u>Chuang et al.</u> (2010)	Taiwan		26.83 (9.7)	Max: 62.1

Table 6-32Characterization of ozone concentrations (in ppb) from studies of
biomarkers.

*Note: Median presented (information on mean not given); studies presented in order of first appearance in the text of this section.

Hemostasis and coagulation markers

1	Multiple studies used various markers to examine if associations were present between
2	short-term O_3 exposure and hemostasis and coagulation. Some of the markers included in
3	these studies were as follows: fibrinogen, von Willebrand factor (vWF), plasminogen
4	activator fibrinogen inhibitor-1 (PAI-1), tissue-type plasminogen activator (tPA), platelet
5	aggregation, and thrombin generation.
6	A population-based study in the United States was conducted to assess the relationship
7	between short-term exposure to air pollution and markers of blood coagulation using the
8	Atherosclerosis Risk in Communities (ARIC) study cohort (Liao et al., 2005). Significant
9	curvilinear associations were observed for O_3 (1 day prior to blood draw) and fibrinogen
10	and vWF (quantitative results not provided for regression models although adjusted
11	means [SE] of vWF were given as 118% [0.79%] for O_3 concentrations <40 ppb, 117%
12	[0.86%] for O ₃ concentrations 40-70 ppb, and 124% [1.97%] for O ₃ concentrations of
13	70 ppb). The association between short-term O_3 exposure and fibrinogen was more

- 1 pronounced among those with a history of cardiovascular disease (CVD) and was 2 statistically significant among only this subgroup of the population. The curvilinear 3 relationship between concentration and outcome suggested stronger relationships at 4 higher concentrations of O_3 . The authors note that the most pronounced associations 5 occurred when the pollutant concentrations were 2-3 standard deviations above the mean. 6 The results from this relatively large-scale cross-sectional study suggest weak 7 associations with between short-term O_3 exposure and increases in fibrinogen (among 8 those with a history of CVD) and vWF. A retrospective repeated measures analysis was 9 performed in Toronto, Canada among adults aged 18-40 years (n = 45) between the years 10 of 1999 and 2006 (Thompson et al., 2010). Single pollutant models were used with moving averages up to 7 days. No evidence of an association was observed between 11 12 short-term O₃ exposure and increases in fibrinogen.
- 13 A repeated measures study was conducted among 40 healthy individuals living or 14 working in the city center of Rotterdam, the Netherlands to assess the relationship 15 between air pollution and markers of hemostatis and coagulation (platelet aggregation, 16 thrombin generation, and fibrinogen) (Rudez et al., 2009). Each participant provided between 11 and 13 blood samples throughout a 1-year period (498 samples on 197 days). 17 18 Examined lags ranged from 6 hours to 3 days prior to blood sampling. No consistent 19 evidence of an association was observed between O3 concentration and any of the 20 biomarkers (percent change of max platelet aggregation: -6.87 [95% CI: -21.46, 7.70] per 21 20 ppb increase in 24-h avg O_3 concentration at 4-day average; percent change of 22 endogenous thrombin potential: 0.95 [95% CI: -3.05, 4.95] per 20 ppb increase in 23 24-h avg O₃ concentration at 4-day avg; percent change of fibrinogen: -0.57 [95% CI: 24 -3.05, 2.00] per 20 ppb increase in 24-h avg O₃ concentration at lag 1-day). Some 25 associations with O_3 were in the opposite direction to that hypothesized which may be 26 explained by the negative correlation between O_3 and other pollutants (correlation 27 coefficients ranged from -0.4 to -0.6). The statistically significant inverse effects 28 observed in single-pollutant models with O_3 were no longer apparent when PM_{10} was 29 included in the model (Rudez et al., 2009).
- 30 A panel study in Taiwan measured health endpoints using blood samples from healthy 31 individuals (n = 76) at three times from April to June in 2004 or 2005 (Chuang et al., 32 2007a). Increases in fibrinogen and PAI-1 were associated with increases in O_3 33 concentrations in single-pollutant models (percent change in fibrinogen: 11.76 [95% CI: 34 4.03, 19.71] per 20 ppb 3-day cumulative avg O₃ concentration; percent change in PAI-1: 35 6.08 [95% CI: 38.91, 84.27] per 20 ppb 3-day cumulative avg O₃ concentration). These 36 associations were also observed at 1 and 2 day averaging times. Associations between 37 PAI-1 and 3-day O₃ concentrations remained robust in two-pollutant models with sulfate.

1	No association was observed between O ₃ concentration and tPA, a fibrinolytic factor
2	(percent change 16.15 [95% CI: -4.62, 38.34] per 20 ppb 3-day avg O_3 concentration).
3	A study in Israel examined the association between pollutant concentrations and
4	fibrinogen among 3659 apparently healthy individuals (Steinvil et al., 2008). In single
5	pollutant models, O_3 was associated with an increase in fibrinogen at a 4-day lag among
6	men and a same-day O_3 concentration among women but results for other lags (0 through
7	7 days) were mixed (i.e., some positive and some negative; none statistically significant).

Inflammatory markers

- 8 Potential associations between short-term exposures to air pollution and inflammatory
 9 markers (C-reactive protein [CRP], white blood cell [WBC] count, albumin, and
 10 Interleukin-6 [IL-6]) were also examined in several studies.
- 11The ARIC study cohort, which included men and women aged 45-64 years old at the start12of the study, was utilized to assess the association between O3 concentrations and13markers of inflammation, albumin and WBC count (Liao et al., 2005). No association14was observed between O3 concentrations and albumin or WBC count.
- 15 Thompson et al. (2010) assessed ambient air pollution exposures and IL-6. This 16 retrospective repeated measures analysis was conducted among 45 adults (18-40 years of 17 age) in Toronto, Canada between the years of 1999 and 2006. Single pollutant models 18 were used to analyze the repeated-measures data using moving averages up to 7 days. A 19 positive association was observed between IL-6 and short-term $1-h O_3$ exposure with the 20 strongest effects observed for the average of lags 0-3 days (quantitative results not 21 provided). No association was observed for shorter averaging times (average lags of 22 <1 day). When examined by season using 2-day moving averages, the association 23 between short-term O₃ exposure and IL-6 was positive during only the spring and 24 summer.
- 25In Rotterdam, the Netherlands, a repeated measures study of healthy individuals living or26working in the city center reported no association between short-term O3 exposure and27CRP (Rudez et al., 2009). Each of the 40 participants provided between 11 and 13 blood28samples throughout a 1-year period (498 samples on 197 days). No consistent evidence of29an association was observed between O3 concentration and CRP (percent change: -0.4830[95% CI: -14.05, 13.10] per 20 ppb increase in 24-h avg O3 concentration at lag 1-day).31Additionally, no association was observed with 2 or 3 day lags.
- 32The relationship between pollutant concentrations and one-time measures of33inflammatory biomarkers was assessed in sex-stratified analyses among 3,659 apparently

healthy individuals in Tel Aviv, Israel (Steinvil et al., 2008). No evidence of an
association was observed between O_3 concentration and CRP or WBC for men and
women.

4	A panel study of healthy individuals $(n = 76)$ was conducted in Taiwan to assess the
5	relationship between air pollutants and inflammation (Chuang et al., 2007a). Health
6	endpoints were measured three times from April to June in 2004 or 2005. Ozone effects
7	were assessed in statistical models using the average of the 24 hours (1 day), 48 hours
8	(2 days), and 72 hours (3 days) before the hour of each blood sampling. Increases in CRP
9	were associated with increases in O ₃ concentrations in single-pollutant models (percent
10	change in CRP: 244.38 [95% CI: 4.54, 585.15] per 20 ppb 3-day avg O_3 concentration).
11	The association was also observed using a 2-day cumulative averaging time, but no
12	association was present with a 1-day averaging time.

Oxidative stress markers

1 2 3

13 A few studies have reported on the relationships between short-term O_3 exposure and 14 increases in markers of oxidative stress. The association between O_3 concentration and 15 markers of lipid peroxidation and antioxidant capacity was examined among 120 nonsmoking healthy college students, aged 18-22 years, from the University of 16 California, Berkeley (February-June 2002) (Chen et al., 2007a). By design, students were 17 18 chosen that had experienced different geographic concentrations of O_3 over their lifetimes 19 and during recent summer vacation in either greater Los Angeles (LA) or the 20 San Francisco Bay Area (SF). Long-term (based on lifetime residential history) and 21 shorter-term (based on the moving averages of 8-h max concentrations 1-30 days prior to 22 the day of blood collection) O_3 concentration were estimated (lifetime exposure results 23 are presented in Chapter 7). A marker of lipid peroxidation, 8-isoprostane (8-iso-PGF), 24 was assessed. This marker is formed continuously under normal physiological conditions 25 but has been found at elevated concentrations in response to environmental exposures. A 26 marker of overall antioxidant capacity, ferric reducing ability of plasma (FRAP), was also 27 measured. Levels of 8-iso-PGF were associated with 2-week ($\beta = 0.035$ 28 [pg/mL]/8-hour ppb O₃, p = 0.007) and 1-month (β = 0.031 [pg/mL]/8-hour ppb O₃, 29 p = 0.006) estimated O₃ concentrations. No evidence of association was observed 30 between short-term O_3 exposure and increases in FRAP. A chamber study performed 31 among a subset of study participants supported the primary study results. The 32 concentrations of 8-iso-PGF increased immediately after the 4-hour controlled O₃ 33 exposure ended (p = 0.10). However, levels returned to near baseline by 18 hours without 34 further exposure. The authors note that O₃ was highly correlated with PM_{10-2.5} and NO₂ in 35 this study population; however, O₃ associations remained robust in copollutant models.

1	Using blood samples collected between April and June of 2004 or 2005 in Taiwan, the
2	association between short-term O3 exposure and a marker of oxidative stress (i.e., 8-
3	hydroxy-2'-deoxyguanosine (8-OHdG)) was studied among healthy individuals (n = 76)
4	(<u>Chuang et al., 2007a</u>). Increases in 8-OHdG were associated with increases in O_3
5	concentrations in single-pollutant models (percent change in 8-OHdG: 2.46 [95% CI:
6	1.01, 3.92] per 20 ppb increase in 24-h avg O ₃). The association did not persist with 2- or
7	3-day cumulative averaging times.

Markers of overall cardiovascular health

8 Multiple studies used markers that assess overall cardiovascular well-being. Wellenius et 9 al. (2007) examined B-type natriuretic peptide (BNP), a marker of heart failure, in a 10 repeated-measures study conducted in Boston among 28 patients with congestive heart 11 failure and impaired systolic function. The authors found no evidence of an association 12 between BNP and short-term O_3 exposures at lags 0-3 days (quantitative results not 13 provided). BNP was chosen because it is directly associated with cardiac hemodynamics 14 and symptom severity among those with heart failure and is considered a marker of 15 functional status. However, the authors conclude that the use of BNP may not be useful 16 in studies of the health effects of ambient air pollutants due to the large amount of within-17 person variability in BNP levels observed in this population.

- 18 The relationship between air pollution and oxygen saturation and pulse rate, markers of 19 physiological well-being, was examined in a 2-month panel study among 31 congestive 20 heart failure patients (aged 50-85 years) in Montreal, Canada from July 2002 to October 21 2003 (Goldberg et al., 2008). All participants had limited physical functioning 22 (New York Heart Association Classification \geq II) and an ejection fraction (the fraction of 23 blood pumped out of the heart per beat) less than or equal to 35% (normal is above 55%). 24 Daily mean O_3 concentrations were calculated based on hourly measures at 10 monitoring 25 stations. There was an inverse association between O_3 concentration (lag-0) and oxygen 26 saturation when adjustment was made for temporal trends. In the models incorporating 27 personal covariates and weather factors, the association remained but was not statistically 28 significant. The associations of O₃ concentration with a lag of 1 day or a 3-day mean 29 were not statistically significant. No evidence of association was observed between O_3 30 concentration and pulse rate.
- 31Total homocysteine (tHcy) is an independent risk factor for vascular disease and32measurement of this marker after oral methionine load is used to identify individuals with33mild impairment of homocysteine metabolism. The effects of air pollution on fasting and34postmethionine-load tHcy levels were assessed among 1,213 apparently healthy35individuals from Lombardia, Italy from January 1995 to September 2005 (Baccarelli et

1	al., 2007). A 20-ppb increase in the 24-h avg O ₃ concentrations was associated with an
2	increase in fasting tHcy (percent change 6.25 [95% CI: 0.84, 11.91]) but no association
3	was observed with postmethionine-load tHcy (percent change 3.36 [95% CI: -1.30,
4	8.39]). In addition, no evidence of an association was observed between 7-day
5	cumulative averaged O_3 concentrations and tHcy (percent change for fasting tHcy 4.16
6	[95% CI: -1.76, 10.42] and percent change for postmethionine-load tHcy -0.65 [95% CI:
7	-5.66, 4.71] per 20 ppb increase in 24-h avg O ₃ concentrations). No evidence of effect
8	modification by smoking was observed.

Blood lipids and glucose metabolism markers

9 Chuang et al. (2010) conducted a population-based cross-sectional analysis of data 10 collected on 7,778 participants during the Taiwanese Survey on Prevalence of 11 Hyperglycemia, Hyperlipidemia, and Hypertension in 2001. Apolipoprotein B (ApoB), 12 the primary apolipoprotein among low-density lipoproteins, was associated with 3-day 13 avg O_3 concentration at the p <0.10 level. The 5-day mean O_3 concentration was 14 associated with an increase in triglycerides at p <0.10. In addition, the 1-, 3-, and 5-day 15 mean O₃ concentrations were associated with increased HbA1c levels (a marker used to 16 monitor the degree of control of glucose metabolism) at the p <0.05 level. The 5-day 17 mean O_3 concentration was associated with increased fasting glucose levels (p <0.10). No 18 association was observed between O₃ concentration and ApoA1.

6.3.2.5 Myocardial Infarction (MI)

19	The 2006 O ₃ AQCD did not report consistent results indicating an association between
20	short-term O ₃ exposure and MI. One study reported a positive association between
21	current day O_3 concentration and acute MI, especially among the oldest age group (55 to
22	64 year-olds) (Ruidavets et al., 2005b). No association was observed in a case-crossover
23	study of O_3 concentration during the surrounding hours and MI (<u>Peters et al., 2001</u>). Since
24	the 2006 O ₃ AQCD, a few recent epidemiologic studies have examined the association
25	between O_3 concentration and MI (<u>Henrotin et al., 2010</u> ; <u>Rich et al., 2010</u>), arterial
26	stiffness (Wu et al., 2010) and ST-segment depression (Delfino et al., 2011).
27	One of the studies conducted in the U.S. examined hospital admissions for first MI and
28	reported no association with O_3 concentration (<u>Rich et al., 2010</u>). More details on this
29	study are reported in the section on hospital admissions (Section $6.3.2.7$). A study
30	performed in Dijon, France examined the association between O ₃ concentration and
31	incident and recurrent MI (<u>Henrotin et al., 2010</u>). The mean 8-hour O_3 concentration was
32	19.1 ppb (SD 12.2 ppb). Odds ratios for the association between cumulative O_3

1	concentrations and recurrent MIs were elevated but none of the results were statistically
2	significant (OR: 1.71 [95% CI: 0.91, 3.20] per 20 ppb increase in 24-h avg O ₃
3	concentration for a cumulative lag of 1-3 days). No association was observed for incident
4	MIs. In analyses stratified by vascular risk factors, positive associations were observed
5	between 1-day lagged O ₃ concentration and MIs (incident and recurrent combined)
6	among those who reported having hypercholesterolaemia (OR: 1.52 [95% CI: 1.08, 2.15]
7	per 20 ppb increase in 24-h avg O ₃ concentration) and a slight inverse association was
8	observed among those who reported not having hypercholesterolaemia (OR: 0.69
9	[95% CI: 0.50, 0.94] per 20 ppb increase in 24-h avg O3 concentration). In other stratified
10	analyses combining different vascular factors, only those containing individuals with
11	hypercholesterolaemia demonstrated a positive association; none were inverse
12	associations.
13	Wu et al. (2010) examined mail carriers aged 25-46 years and measured exposure to O_3
14	concentrations through personal monitors [mean O_3 24.9 (SD 14.0) ppb]. Ozone
15	concentration was positively associated with arterial stiffness (percent change 11.24%
16	[95% CI: 3.67, 19.62] per 40 ppb O_3) and was robust to adjustment for ultrafine PM.
17	A study performed in the Los Angeles basin reported on the association between O ₃
18	concentration and ST-segment depression, a measure representing cardiac ischemia
19	(Delfino et al., 2011). Study participants were nonsmokers, at least 65 years old, had a
20	history of coronary artery disease, and were living in a retirement community. Study
21	periods included five consecutive days in both July to mid-October and mid-October to
22	February. Mean 24-hour O ₃ concentrations were 27.1 ppb (SD 11.5 ppb). No association
23	was observed between O_3 concentration and ST-segment depression of at least 1.0 mm
24	during any of the exposure periods (i.e., 1-h, 8-h, 1-day, 2-day avg, 3-day avg,
25	4-day avg).

6.3.2.6 Blood Pressure

26In the 2006 O3 AQCD, no epidemiologic studies examined O3-related effects on blood27pressure (BP). Recent studies have been conducted to evaluate this relationship and28overall the findings are inconsistent. The O3 concentrations for these studies are listed in29Table 6-33.

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
Zanobetti et al. (2004)	Boston,	1-h	20	
	Massachusetts	5-days	24	
Delfino et al. (2010b)	Los Angeles, California	24-h	27.1 (11.5)	Max: 60.7
<u>Choi et al. (2007)</u>	Incheon,	8-h	26.6 (11.8)	75th: 34.8
	South Korea	(warm season)		Max: 62.4
		8-h	17.5 (7.3)	75th: 22.9
		(cold season)		Max: 33.9
Chuang et al. (2010)	Taiwan		26.83 (9.7)	Max: 62.1

Table 6-33Characterization of ozone concentrations (in ppb) from studies of
blood pressure.

*Note: Studies presented in order of first appearance in the text of this section.

1	Zanobetti et al. (2004) examined the relationship between air pollutants and BP from
2	May 1999 to January 2001 for 631 repeat visits among 62 Boston residents with CVD. In
3	single-pollutant models, higher resting diastolic blood pressure (DBP) was associated
4	with the 5-day (0-4 days) averages of O ₃ concentration (RR: 1.03 [95% CI: 1.00, 1.05]
5	per 20 ppb increase in 24-hour O ₃ concentrations). However, this effect was no longer
6	apparent when $PM_{2.5}$ was included in the model (data were not presented) (Zanobetti et
7	al., 2004). Delfino et al. (2010b) examined 64 subjects 65 years and older with coronary
8	artery disease, no tobacco smoke exposure, and living in retirement communities in the
9	Los Angeles air basin with hourly (up to 14-h/day) ambulatory BP monitoring for 5 days
10	during a warm period (July-mid-October) and 5 days during a cool period (mid-October-
11	February). Investigators assessed lags of 1, 4, and 8 hours, 1 day, and up to 9 days before
12	each BP measure; no evidence of an association was observed for O_3 (change in BP
13	associated with a 20 ppb increase in 24-h avg O_3 concentration was 0.67 [95% CI: -1.16,
14	2.51 for systolic BP [SBP] and -0.25 [95% CI: -1.25, 0.75] for DBP) (Delfino et al.,
15	2010b). Choi et al. (2007) conducted a cross-sectional study to investigate the
16	relationship between air pollutants and BP among 10,459 participants of the Inha
17	University Hospital health examination from 2001 to 2003. These individuals had no
18	medical history of cardiovascular disease or hypertension. O3 concentration was
19	associated with an increase in SBP for 1-day lag in the warm season and similar effect
20	estimates were observed during the cold season but were not statistically significant
21	(quantitative results not provided). Associations between O ₃ concentration and DBP were
22	present in the cold season but not the warm season (quantitative results not provided).
23	Chuang et al. (2010) conducted a similar type of study among 7,578 participants of the
24	Taiwanese Survey on Prevalence of Hyperglycemia, Hyperlipidemia, and Hypertension
25	in 2001. Investigators examined 1-, 3-, and 5-day avg O_3 concentrations. An increase in

1	DBP was associated with the 3-day mean O_3 concentration (change in BP for a 20 ppb
2	increase in 24-h avg O ₃ concentration was 0.61 [95% CI: 0.07, 1.14]) (Chuang et al.,
3	2010). Associations were not observed for other days or with SBP.

6.3.2.7 Hospital Admissions and Emergency Department Visits

4	Upon evaluating the collective evidence for O3-related cardiovascular hospital admissions
5	and emergency department (ED) visits, the 2006 O3 AQCD concluded that "a few studies
6	observed positive O3 associations, largely in the warm season. Overall, however, the
7	currently available evidence is inconclusive regarding any association between ambient
8	O_3 exposure on cardiovascular hospitalizations" (U.S. EPA, 2006b). Table 6-34 below
9	provides information on the O_3 concentrations reported in each of the recent hospital
10	admission and ED visit studies evaluated.

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
Peel et al. (2007)	Atlanta, GA	8-h max warm season	55.6 (23.8)	
Tolbert et al. (2007)	Atlanta, GA	8-h max warm season	53.0	75th: 67.0 Max: 147.5
Katsouyanni et al. (2009)	12 Canadian cities	1-h	6.7-8.3*	75th: 8.4-12.4
	8 European cities	1-h	11.0-38.1*	75th: 15.3-49.4
	14 United Statescities	1-h	34.9-60.0*	75th: 46.8-68.8
<u>Rich et al. (2010)</u>	New Jersey	24-h	NR	
Cakmak et al. (2006a)	10 Canadian cities	1-h max	17.4	
<u>Stieb et al. (2009)</u>	7 Canadian cities	24-h	18.4	
Szyszkowicz (2008)	Edmonton, Canada	24-h	18.6 (9.3)	
Villeneuve et al. (2006a)	Edmonton, Canada	24-h	17 (9.1)	75th: 23.5
		24-h warm season	21.8 (8)	75th: 27.0
		24-h cold season	12.2 (7.4)	75th: 17.0
<u>Symons et al. (2006</u>)	Baltimore, MD	8-h warm season	31.0 (20.0)	Max: 120.0
Wellenius et al. (2005)	Allegheny County, PA	24-h	24.3 (12.2)	75th: 32.0
Zanobetti and Schwartz (2006)	Boston, MA	24-h	22.4*	75th: 31.0
Yang (2008)	Taipei, Taiwan	24-h	21.0	75th: 26.3 Max: 62.8
<u>Lee et al. (2007)</u>	Kaohsiung, Taiwan	24-h	26.5	75th: 35.5 Max: 83.0
<u>Chan et al. (2006</u>)	Taipei, Taiwan	1-h max	50.9 (26.4)	Max: 150.3
Chiu and Yang (2009)	Taipei, Taiwan	24-h	23.0	75th: 28.7 Max: 62.8
Lee et al. (2008a)	Taipei, Taiwan	24-h	21.0	75th: 26.4 Max: 62.8
Wong et al. (2009)	Hong Kong	8-h	18.5 (11.5)	75 th : 25.4 Max: 48.3
Bell et al. (2008)	Taipei, Taiwan	24-h	21.4	Max: 48.3
Buadong et al. (2009)	Bangkok, Thailand	1-h	14.4 (3.2)	Max: 33.4 Max: 41.9
Lee et al. (2003b)	Seoul, Korea	1-h max	36.0 (18.6)	75th: 44.9
Azevedo et al. (2011)	Portugal	1-h	NR	, ош. тт.0
Linares and Diaz (2010)	Madrid,Spain	24-h	17.4 (8.9)	
Middleton et al. (2008)	Nicosia, Cyprus	8-h max	28.7 - 54.9	

Table 6-34Characterization of ozone concentrations (in ppb) from studies of
hospital admissions and ED visits.

Study*	Location	Averaging Time	Mean Concentration (Standard Deviation)	Upper Range of Concentration
<u>Turner et al. (2007)</u>	Sydney, Australia	24-h	28	75th: 33
Ballester et al. (2006)	14 Spanish cities	8-h warm season	24.2 - 44.3	
DePablo et al. (2006)	Castilla-Leon, Spain	24-h	23.2-33.6	
VonKlot et al. (2005)	5 European cities	8 h max warm season	16.4 - 28.0	
<u>Oudin et al. (2010</u>)	Scania, Sweden	24-h	30.5	
<u>Halonen et al. (2009</u>)	Helsinki, Finland	8-h max warm season	35.7*	75th: 42.1 Max: 79.6
Larrieu et al. (2007)	8 French cities	8-h max warm season	34.2 - 53.1	
Barnett et al. (2006)	4 Australian cities	8-h	19.0-28.5	Max: 58.4-86.8
<u>Hinwood et al. (2006</u>)	Perth, Australia	8-h max	25.9 (6.5)	
Lanki et al. (2006)	5 European cities	8-h max warm season	31.7 - 57.2*	
Hosseinpoor et al. (2005)	Tehran, Iran	8-h max	4.9 (4.8)	75th: 7.2 Max: 99.0
Simpson et al. (2005)	4 Australian cities	1-h max	24.4-33.8	Max: 96.0-111.5
Dennekamp et al. (2010)	Melbourne, Australia	24-h	13.34	75th: 16.93
<u>Silverman et al. (2010)</u>	New York City, NY	8-h max	28*	75th: 40

*Notes: Median presented (information on mean not given); NR: Not reported; studies presented in order of first appearance in the text of this section.

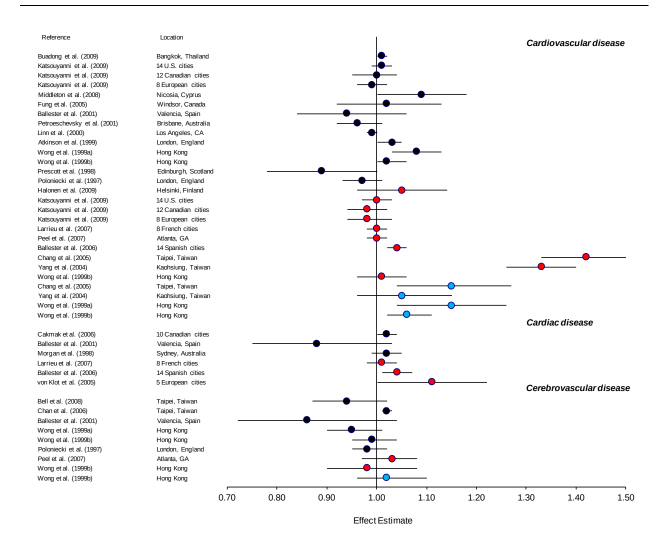
1	Multiple recent studies of O_3 concentration and cardiovascular hospital admissions and
2	ED visits have been conducted in the U.S. and Canada. Peel et al. (2007) used a case-
3	crossover framework (using a time-stratified approach matching on day of the week in
4	the calendar month of the event) to assess the relationship between air pollutants and
5	cardiovascular disease ED visits among those with and without secondary comorbid
6	conditions (hypertension, diabetes, chronic obstructive pulmonary disease [COPD],
7	congestive heart failure [CHF], and dysrhythmia). Data on over 4 million ED visits from
8	31 hospitals were collected from January 1993 to August 2000. Ozone was monitored
9	from March to October. This study was a re-analysis of a time series study conducted to
10	assess the main effects of air pollutants on cardiovascular ED visits in Atlanta (Tolbert et
11	al., 2007; Metzger et al., 2004). In the initial study, no evidence of associations was
12	observed between O ₃ concentration and all CVD visits or visits for CVD subgroups, such
13	as dysrhythmia, CHF, ischemic heart disease (IHD), and peripheral vascular and
14	cerebrovascular disease. The relative risk for all CVD visits was 1.01 (95% CI: 0.98,
15	1.04) for a 30 ppb increase in the 3-day moving avg (lags 0-2 days) of 8-hour O_3
16	concentration (Metzger et al., 2004). Similar to the initial investigation using a time-
17	series analysis, no evidence of an association was observed between short-term O_3

1	exposure and CVD visits at lag 0-2 among the entire population using the case-crossover
2	design (Peel et al., 2007). However, the relationship between O_3 concentration and
3	peripheral and cerebrovascular disease visits was stronger among patients with comorbid
4	COPD (OR: 1.29 [95% CI: 1.05-1.59] per 30 ppb, lag 0-2 days) as compared to patients
5	without COPD (OR: 1.01 [95% CI: 0.96-1.06] per 30 ppb, lag 0-2 days). The same
6	research group expanded upon the number of Atlanta hospitals providing ED visit data
7	(41 hospitals) as well as the length of the study period (1993-2004) (Tolbert et al., 2007).
8	Again, models assessing the health effects of O_3 concentration utilized data collected
9	from March through October. Similar to the results presented by Metzger et al. (2004)
10	and <u>Peel et al. (2007</u>) among the entire study population, no evidence of associations was
11	observed for O_3 concentration and CVD visits (<u>Tolbert et al., 2007</u>).
12	Existing multicity studies in North America and Europe were evaluated under a common
13	framework in the Air Pollution and Health: A European and North American Approach
14	(APHENA) study (Katsouyanni et al., 2009). One component of the study examined the
15	relationship between short-term O ₃ exposure and CVD hospital admissions among
16	individuals 65 years of age and older. The study presented multiple models but this
17	section focuses on the results for the models that used 8 df to account for temporal trends
18	and natural splines (see Section $6.2.7.2$ for additional explanation). Across the study
19	locations, no associations were observed between O3 concentration and CVD hospital
20	admissions at lags 0-1, lag 1, or a distributed lag of 0-2. Additionally, there was no
21	evidence of an association when restricting the analysis to the summer months.
22	A study of hospital admissions for MI was performed using a statewide registry from
23	New Jersey between January 2004 and December 2006 (Rich et al., 2010). Using a case-
24	crossover design, the association between the previous 24-h O_3 concentration and
25	transmural infarction ($n = 1,003$) was examined. No association was observed (OR: 0.94
26	[95% CI: 0.79, 1.13] per 20 ppb increase in 24-h avg O ₃ concentration) and this did not
27	change with the inclusion of $PM_{2.5}$ in the model.
28	Cakmak et al. (2006a) investigated the relationship between gaseous air pollutants and
29	cardiac hospitalizations in 10 large Canadian cities using a time-series approach. A total
30	of 316,234 hospital discharge records for primary diagnosis of congestive heart failure,
31	ischemic heart disease, or dysrhythmia were obtained from April 1993 through March
32	2000. Correlations between pollutants varied substantially across cities, which could
33	partially explain discrepancies in effect estimates observed across the cities. In addition,
34	pollutant lags differed across cities; the average lag for O_3 was 2.9 days. The pooled
35	effect estimate for a 20 ppb increase in the daily 1-h max O_3 concentration and the
36	percent change in hospitalizations among all 10 cities was 2.3 (95% CI: 0.11, 4.50) in an
37	all-year analysis. The authors reported no evidence of effect modification by sex,

- 1 neighborhood-level education, or neighborhood-level income. A similar multicity time-2 series study was conducted using nearly 400,000 ED visits to 14 hospitals in seven 3 Canadian cities from 1992 to 2003 (Stieb et al., 2009). Primary analyses considered daily 4 O_3 single day lags of 0-2 days; in addition, sub-daily lags of 3-h avg concentrations up to 5 12 hours before presentation to the ED were considered. Seasonal variation was assessed 6 by stratifying analyses by warm and cold seasons. No evidence of associations between 7 short-term O₃ exposure and CVD ED visits was observed. One negative, statistically 8 significant association was reported between a 1-day lag of O₃ concentration and visits 9 for angina/myocardial infarction. Ozone concentration was negatively correlated with 10 many of the other pollutants, particularly during the cold season.
- 11 The effect of air pollution on daily ED visits for ischemic stroke (n = 10,881 visits) in 12 Edmonton, Canada was assessed from April 1992 through March 2002 (Szyszkowicz, 13 2008). A 26.4% (95% CI: 3.16-54.5) increase in stroke ED visits was associated with a 14 20 ppb increase in 24-hour average O_3 concentration at lag 1 among men aged 20-15 64 years in the warm season. No associations were present among women or among men 16 age 65 and older. In addition, no associations were observed for the cold season or for 17 other lags (lag 0 or lag 2). A similar investigation over the same time period in 18 Edmonton, Canada, assessed the relationship between air pollutants and ED visits for 19 stroke (ischemic stroke, hemorrhagic stroke, and transient ischemic attack) among those 20 65 years of age and older using a case-crossover framework (Villeneuve et al., 2006a). 21 No evidence of association was reported for O₃ concentration and stroke hospitalization 22 in single or co-pollutant models (Villeneuve et al., 2006a).
- 23 Additional studies in the U.S. reported no evidence of an association between O_3 24 concentrations and ED visits, hospitalizations, or symptoms leading to hospitalization 25 (Symons et al., 2006; Zanobetti and Schwartz, 2006; Wellenius et al., 2005). Symons et 26 al. (2006) used a case-crossover framework to assess the relationship between air 27 pollutants and the onset of symptoms (dyspnea) severe enough to lead to hospitalization 28 (through the ED) for congestive heart failure. The study was conducted from April to 29 December of 2002 in Baltimore, Maryland. Exposures were assigned using 3 index times: 30 8-hour and 24-hour periods prior to symptom onset and date of hospital admission. No 31 evidence of association was reported for O₃ concentrations. Although seasonal variation 32 was not assessed, the time frame for the study did not involve an entire year (April to 33 December). Wellenius et al. (2005) investigated the association between air pollutants 34 and congestive heart failure hospitalization among Medicare beneficiaries in Pittsburgh, 35 Pennsylvania from 1987 to 1999 utilizing a case-crossover framework. A total of 55,019 36 admissions from the emergency room with a primary discharge diagnosis of CHF were 37 collected. No evidence of an association was reported for O₃ concentration and CHF 38 hospitalization (Wellenius et al., 2005). Finally, Zanobetti and Schwartz (2006) assessed

1	
1	the relationship between air pollutants and hospital admissions through the ED for MI
2	and pneumonia among patients aged 65 and older residing in the greater Boston area
3	(1995-1999) using a case-crossover framework with control days in the same month
4	matched on temperature. Pollution exposures were assigned for the same day and for the
5	mean of the exposure the day of and the day before the admission. Ozone concentration
6	was not associated with MI admissions in all-year and seasonal analyses.
7	Several recent studies have examined the relationship between air pollution and CVD
8	hospital admissions and/or emergency department visits in Asia. Of note, some areas of
9	Asia have a more tropical climate than the U.S. and do not experience similar seasonal
10	changes. In Taiwan, fairly consistent positive associations have been reported for O_3
11	concentration and congestive heart failure hospital admissions (for single- and
12	copollutant models) in Taipei on warm days (Yang, 2008) and in Kaohsiung (Lee et al.,
13	2007); cerebrovascular disease ED visits (for lag 0 single- and two-pollutant models but
14	not other lags) in Taipei (Chan et al., 2006); and arrhythmia ED visits in Taipei among
15	those without comorbid conditions (Chiu et al., 2009; Lee et al., 2008a) and in Taipei on
16	warm days among those with and without comorbid conditions (Lee et al., 2008a).
17	However, one study in Taiwan did not show an association. Bell et al. (2008) reported no
18	evidence of an association between O ₃ concentration and hospital admissions for
19	ischemic heart disease or cerebrovascular disease. Studies based in Asia but outside
20	Taiwan were also performed. A Hong Kong-based investigation (Wong et al., 2009)
21	reported no consistent evidence of a modifying effect of influenza on the relationship
22	between O ₃ concentration and CVD admissions. Among elderly populations in Thailand,
23	O ₃ concentration was associated with CVD visits, but this association was not detected
24	among younger age groups (15-64) (<u>Buadong et al., 2009</u>). Also, a study performed in
25	Seoul, Korea reported a positive association between O ₃ concentration and hospital
26	admissions for ischemic heart disease; the association was slightly greater among those
27	over 64 years of age (Lee et al., 2003b).
28	Positive associations between short-term O ₃ exposure and CVD hospital admissions
29	and/or ED visits have been reported in other areas of the world as well (Azevedo et al.,
30	2011; Linares and Diaz, 2010; Middleton et al., 2008; Turner et al., 2007; Ballester et al.,
31	2006; DePablo et al., 2006; VonKlot et al., 2005), although not consistently as some
32	studies reported no association (<u>Oudin et al., 2010; Halonen et al., 2009; Larrieu et al.</u> ,
33	2007; Barnett et al., 2006; Hinwood et al., 2006; Lanki et al., 2006; Hosseinpoor et al.,
34	<u>2005; Simpson et al., 2005</u>).
35	A couple of studies (U.S. and Australia) have examined cardiac arrests where emergency
36	services attempted treatment/resuscitation. No evidence of an association between O_3
50	services anompted rearment resuscitation. To evidence of an association between O3

1 2	concentration and out-of-hospital cardiac arrest was observed (<u>Dennekamp et al., 2010</u> ; <u>Silverman et al., 2010</u>).
3	An increasing number of air pollution studies have investigated the relationship between
4	O3 concentrations and CVD hospital admissions and/or ED visits. As summarized in the
5	2006 O ₃ AQCD, some, especially those reporting results stratified by season (or
6	temperature) or comorbid conditions have reported positive associations. However, even
7	studies performing these stratified analyses are not consistent and the overall evidence
8	remains inconclusive regarding the association between short-term O3 exposure and CVD
9	hospital admissions and ED visits. The Hospital Admission (HA) and ED visit studies
10	evaluated in this section are summarized in Figure 6-21 through Figure 6-25, which
11	depict the associations for studies in which quantitative data were presented. Table 6-35
12	through <u>Table 6-39</u> provide the numerical results displayed in the figures.



Note: Change in O_3 standardized to 20 ppb for 24-h avg period, 30 ppb for 8-h avg period, and 40 ppb for 1-h avg period (see Section 2.2). Ozone concentrations in ppb. Seasons depicted by colors – black: all year; red: warm season; light blue: cold season. Age groups of study populations were not specified or were adults with the exception of <u>Katsouyanni et al. (2009</u>), <u>Fung et al.</u> (2005), <u>Wong et al. (1999b</u>), and <u>Prescott et al. (1998</u>), which included only individuals aged 65+. Studies organized by outcome and season and then listed in descending order of publication date.

Figure 6-21 Effect estimate (95% CI) per increment ppb increase in ozone for over all cardiovascular ED visits or hospital admissions.

Table 6-35Effect estimate (95% CI) per increment ppb increase in ozone for
overall cardiovascular ED visits or hospital admissions in studies
presented inFigure 6-21.

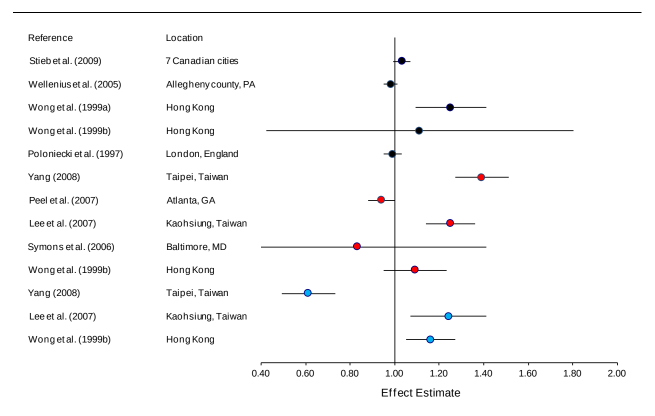
Study*	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
<u>Atkinson et al. (1999</u>)	London, England	Cardiovascular disease	8-h	1.03 (1.00, 1.05)
Ballester et al. (2006)	14 Spanish cities	Cardiovascular disease	8-h warm season	1.04 (1.02 , 1.06)
		Cardiac disease	8-h warm season	1.04 (1.01, 1.07)
Ballester et al. (2006)	Valencia, Spain	Cardiovascular disease	8-h	0.94 (0.84, 1.06)
		Cardiac disease	8-h	0.88 (0.75, 1.03)
		Cerebrovascular disease	8-h	0.86 (0.72, 1.04)
<u>Bell et al. (2008</u>)	Taipei, Taiwan	Cerebrovascular disease	24-h	0.94 (0.87, 1.02)
Buadong et al. (2009)	Bangkok, Thailand	Cardiovascular disease	1-h	1.01 (1.00, 1.02)
<u>Cakmak et al. (2006a</u>)	10 Canadian cities	Cardiac disease	1-h max	1.02 (1.00, 1.04)
<u>Chan et al. (2006</u>)	Taipei, Taiwan	Cerebrovascular disease	1-h max	1.02 (1.01, 1.03)
<u>Chang et al. (2005</u>)	Taipei, Taiwan	Cardiovascular disease	24-h warm season	1.42 (1.33 , 1.50)
			24-h cold season	1.15 (1.04, 1.27)
Fung et al. (2005)	Windsor, Canada	Cardiovascular disease	1-h	1.02 (0.92, 1.13)
Halonen et al. (2009)	Helsinki, Finland	Cardiovascular disease	8-h max warm season	1.05 (0.96, 1.14)
Katsouyanni et al.	14 U.S. cities	Cardiovascular disease	1-h max	1.01 (0.99, 1.03)
<u>(2009</u>)			1-h max warm season	1.00 (0.97, 1.03)
	12 Canadian	Cardiovascular	1-h max	1.00 (0.95, 1.04)
	cities	disease	1-h max warm season	0.98 (0.94, 1.02)
	8 European cities		1-h max	0.99 (0.96, 1.02)
		disease	1-h max warm season	0.98 (0.94, 1.03)
Larrieu et al. (2007)	8 French cities	Cardiac disease	8-h max warm season	1.01 (0.98, 1.04)
Linn et al. (2000)	Los Angeles, California	Cardiovascular disease	24-h	0.99 (0.98, 1.00)
Middleton et al. (2008)	Nicosia, Cyprus	Cardiovascular disease	8-h max	1.09 (1.00, 1.18)

Study*	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
<u>Morgan et al. (1998)</u>	Sydney, Australia	Cardiac disease	1-h max	1.02 (0.99, 1.05)
<u>Peel et al. (2007</u>)	Atlanta, GA	Cardiovascular disease	8-h warm season	1.00 (0.98, 1.02)
		Cerebrovascular disease	8-h warm season	1.03 (0.97, 1.08)
Petroeschevsky et al. (2001)	Brisbane, Australia	Cardiovascular disease	8-h	0.96 (0.92, 1.01)
Poloniecki et al. (1997)	London, England	Cardiovascular disease	8-h	0.97 (0.93, 1.01)
		Cerebrovascular disease	8-h	0.98 (0.95, 1.02)
Prescott et al. (1998)	Edinburgh, Scotland	Cardiovascular disease	24-h	0.89 (0.78, 1.00)
VonKlot et al. (2005)	5 European cities	Cardiac disease	8-h max warm season	1.11 (1.00, 1.22)
<u>Wong et al. (1999b</u>)	Hong Kong	Cardiovascular disease	24-h	1.08 (1.03 , 1.13)
			24-h cold season	1.15 (1.04, 1.26)
		Cerebrovascular disease	24-h	0.95 (0.90, 1.01)
<u>Wong et al. (1999a</u>)	Hong Kong	Cardiovascular	24-h	1.02 (1.03 , 1.06)
		disease	24-h warm season	1.01 (0.96, 1.06)
			24-h cold season	1.06 (1.02, 1.11)
		Cerebrovascular	24-h	0.99 (0.95, 1.04)
		disease	24-h warm season	0.98 (0.90, 1.08)
			24-h cold season	1.02 (0.96, 1.10)
Yang et al. (2004)	Kaohsiung,	Cardiovascular disease	24-h warm season	1.33 (1.26 , 1.40)
	Taiwan		24-h cold season	1.05 (0.96, 1.15)

*Studies included in Figure 6-21..

Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Age groups of study populations were not specified or were adults with the exception of <u>Katsouyanni et al. (2009</u>), <u>Fung et al. (2005</u>), <u>Wong et al.</u> (1999a), and <u>Prescott et al. (1998</u>), which included only individuals aged 65+. Studies listed in alphabetical order.

Warm season defined as: March-October (<u>Peel et al., 2007</u>), May-October (<u>Ballester et al., 2005</u>; <u>Wong et al., 1999a</u>), May-September (<u>Halonen et al., 2009</u>), April-September (<u>Larrieu et al., 2007</u>; <u>VonKlot et al., 2005</u>) <u>Katsouyanni et al. (2009</u>), ≥ 20°C (<u>Chang et al., 2005</u>) and ≥ 25°C (<u>Yang et al., 2004</u>). Cold season defined as: November-April (<u>Wong et al., 1999a</u>), <20°C (<u>Chang et al., 2005</u>) and <25°C (<u>Yang et al., 2004</u>), December-March (Wong et al., 1999b)



Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Seasons depicted by colors: black: all year; red: warm season; light blue: cold season. Outcomes were all congestive heart failure, with the exception of <u>Symons et al. (2006</u>), which examined onset of congestive heart failure symptoms leading to a heart attack. Age groups of study populations were not specified or were adults with the exception of <u>Wellenius et al. (2005</u>) and <u>Wong et al. (1999a</u>), which included only individuals aged 65+. Studies organized by outcome and season and then listed in descending order of publication date.

Figure 6-22 Effect estimate (95% CI) per increment ppb increase in ozone for congestive heart failure ED visits or hospital admissions.

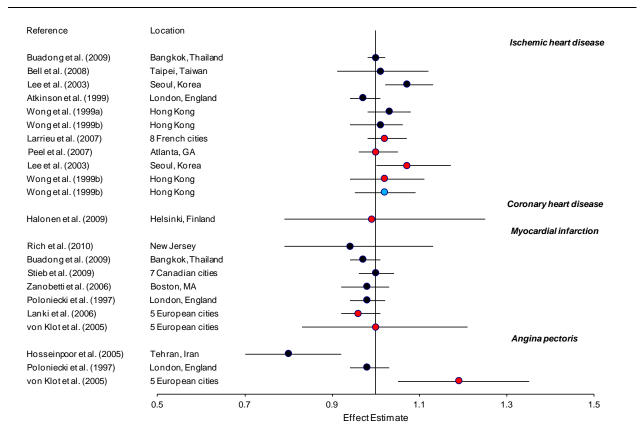
Table 6-36Effect estimate (95% CI) per increment ppb increase in ozone for
congestive heart failure ED visits or hospital admissions for
studies in Figure 6-22.

Study*	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
<u>Lee et al. (2007</u>)	Kaohsiung, Taiwan	Congestive heart failure	24-h warm season	1.25 (1.15, 1.36)
		Congestive heart failure	24-h cold season	1.24 (1.09, 1.41)
Peel et al. (2007)	Atlanta, GA	Congestive heart failure	8-h warm season	0.94 (0.89, 1.00)
<u>Poloniecki et al.</u> (1997)	London, England	Congestive heart failure	8-h	0.99 (0.95, 1.03)
Stieb et al. (2009)	7 Canadian cities	Congestive heart failure	24-h	1.03 (0.98, 1.07)
<u>Symons et al.</u> (2006)	Baltimore, MD	Onset of congestive heart failure symptoms leading to heart attack	8-h warm season	0.83 (0.49, 1.41)
<u>Wellenius et al.</u> (2005)	Allegheny county, PA	Congestive heart failure	24-h	0.98 (0.96, 1.01)
Wong et al.	Hong Kong	Congestive heart failure	24-h	1.11 (1.04, 1.80)
<u>(1999a</u>)			24-h warm season	1.09 (0.96, 1.23)
			24-hcold season	1.16 (1.06, 1.27)
<u>Yang (2008</u>)	Taipei, Taiwan	Congestive heart failure	24-h warm season	1.39 (1.27, 1.51)
		Congestive heart failure	24-h cold season	0.61 (0.52, 0.73)
<u>Wong et al.</u> (1999b)	Hong Kong	Congestive heart failure	24-h	1.25 (1.11, 1.41)

*Studies include those from Figure 6-22.

Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Outcomes were all congestive heart failure, with the exception of <u>Symons et al. (2006</u>), which examined onset of congestive heart failure symptoms leading to a heart attack. Age groups of study populations were not specified or were adults with the exception of <u>Wellenius et al. (2005</u>) and <u>Wong et al. (1999a</u>), which included only individuals aged 65+. Studies listed in alphabetical order.

Warm season defined as: March-October (<u>Peel et al., 2007</u>), April-November (<u>Symons et al., 2006</u>), May-October (<u>Wong et al., 1999a</u>) \geq 20°C (<u>Yang, 2008</u>), and \geq 25°C (<u>Lee et al., 2007</u>). Cold season defined as: November-April (<u>Wong et al., 1999a</u>), <20°C (<u>Yang, 2008</u>), and <25°C (<u>Lee et al., 2007</u>).



Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Seasons depicted by colors: black: all year; red: warm season; light blue: cold season. Age groups of study populations were not specified or were adults with the exception of <u>Wong et al. (1999a)</u> and <u>Atkinson et al. (1999</u>), which included only individuals aged 65+. Studies organized by outcome and season and then listed in descending order of publication date.

Figure 6-23 Effect estimate (95% CI) per increment ppb increase in ozone for ischemic heart disease, coronary heart disease, myocardial infarction, and angina pectoris ED visits or hospital admissions.

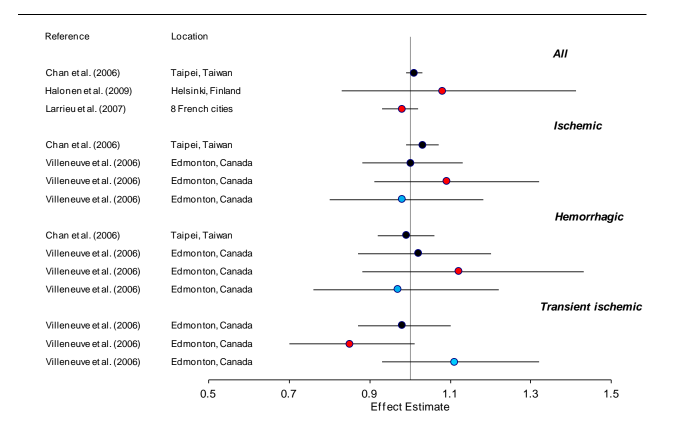
Table 6-37Effect estimate (95% CI) per increment ppb increase in ozone for
ischemic heart disease, coronary heart disease, myocardial
infarction, and angina pectoris Evisits or hospital admissions for
studies presented in Figure 6-23.

Study*	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
Atkinson et al. (1999)	London, England	Ischemic heart disease	8-h	0.97 (0.94, 1.01)
<u>Bell et al. (2008)</u>	Taipei, Taiwan	Ischemic heart disease	24-h	1.01 (0.91, 1.12)
Buadong et al. (2009)	Bangkok,	Ischemic heart disease	1-h	1.00 (0.98, 1.02)
	Thailand	Myocardial infarction	1-h	0.97 (0.94, 1.01)
<u>Halonen et al. (2009</u>)	Helsinki, Finland	Coronary heart disease	8-h max warm season	0.99 (0.79, 1.25)
Hosseinpoor et al. (2005)	Tehran, Iran	Angina	8-h max	0.80 (0.70, 0.92)
Lanki et al. (2006)	5 European cities	Myocardial infarction	8-h max warm season	0.96 (0.92, 1.01)
Larrieu et al. (2007)	8 French cities	Ischemic heart disease	8-h max warm season	1.02 (0.98, 1.07)
Lee et al. (2003b)	Seoul, Korea	Ischemic heart disease	1-h max	1.07 (1.02, 1.13)
		Ischemic heart disease	1-h max warm season	1.07 (1.00, 1.17)
Peel et al. (2007)	Atlanta, GA	Ischemic heart disease	8-h warm season	1.00 (0.96, 1.05)
Poloniecki et al. (1997)	London, England	Myocardial infarction	8-h	0.98 (0.94, 1.02)
		Angina	8-h	0.98 (0.94, 1.03)
<u>Rich et al. (2010</u>)	New Jersey	Myocardial infarction	24-h	0.94 (0.79, 1.13)
<u>Stieb et al. (2009</u>)	7 Canadian cities	Myocardial infarction	2-h	1.00 (0.96, 1.04)
VonKlot et al. (2005)	5 European cities	Myocardial infarction	8-h max warm season	1.00 (0.83, 1.21)
		Angina	8-h max warm season	1.19 (1.05, 1.35)
Wong et al. (1999a)	Hong Kong	Ischemic heart disease	24-h	1.01 (0.94, 1.06)
			24-h warm season	1.02 (0.94, 1.11)
			24-h cold season	1.02 (0.95, 1.09)
Wong et al. (1999b)	Hong Kong	Ischemic heart disease	24-h	1.03 (0.98, 1.08)
Zanobetti and Schwartz (2006)	Boston, MA	Myocardial infarction	24-h	0.98 (0.92, 1.03)

*Sudies included from <u>Figure 6-23</u>.

Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Age groups of study populations were not specified or were adults with the exception of <u>Wong et al. (1999a</u>) and <u>Atkinson et al. (1999</u>), which included only individuals aged 65+. Studies listed in alphabetical order.

Warm season defined as: March-October (<u>Peel et al., 2007</u>), June-August (<u>Lee et al., 2003b</u>), May-September (<u>Halonen et al., 2009</u>), May-October (<u>Buadong et al., 2009</u>), and April-September (<u>Larrieu et al., 2007</u>; <u>Lanki et al., 2006</u>; <u>VonKlot et al., 2005</u>). Cold season defined as: November-April (<u>Buadong et al., 2009</u>).



Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Seasons depicted by colors: black: all year; red: warm season; light blue: cold season. Age groups of study populations were not specified or were adults with the exception of <u>Villeneuve et al.</u> (2006a), which included only individuals aged 65+, and.<u>Chan et al.</u> (2006), which included only individuals aged 50+. Studies organized by outcome and season and then listed in descending order of publication date.

Figure 6-24 Effect estimate (95% CI) per increment ppb increase in ozone for stroke ED visits or hospital admissions.

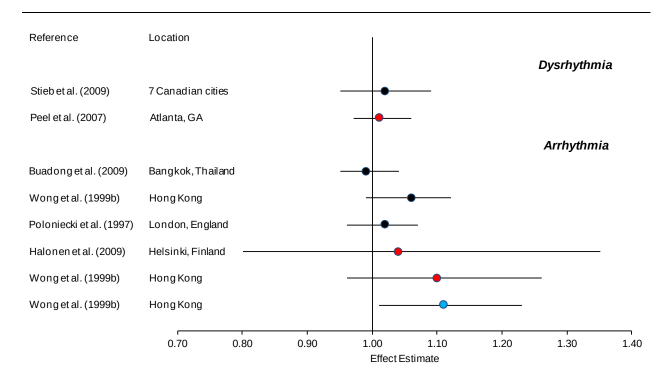
Table 6-38Effect estimate (95% CI) per increment ppb increase in ozone for
stroke ED visits or hospital admissions for studies presented in
Figure 6-24.

Study*	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
<u>Chan et al. (2006)</u>	Taipei, Taiwan	All/non-specified stoke	1-h max	1.01 (0.99 ,1.03)
		Ischemic stroke	1-h max	1.03 (0.99, 1.07)
		Hemorrhagic stroke	1-h max	0.99 (0.92, 1.06)
<u>Halonen et al. (2009</u>)	Helsinki, Finland	All/non-specified stoke	8-h max warm season	1.08 (0.83, 1.41)
<u>Larrieu et al. (2007</u>)	8 French cities	All/non-specified stoke	8-h max warm season	0.98 (0.93 , 1.02)
Villeneuve et al. (2006a)	Edmonton, Canada	Ischemic stroke	24-h	1.00 (0.88, 1.13)
		Ischemic stroke	24-h warm season	1.09 (0.91, 1.32)
		Ischemic stroke	24-h cold season	0.98 (0.80, 1.18)
		Hemorrhagic stroke	24-h	1.02 (0.87, 1.20)
		Hemorrhagic stroke	24-h warm season	1.12 (0.88, 1.43)
		Hemorrhagic stroke	24-h cold season	0.97 (0.76, 1.22)
		Transient ischemic stroke	24-h	0.98 (0.87, 1.10)
		Transient ischemic stroke	24-h warm season	0.85 (0.70, 1.01)
		Transient ischemic stroke	24-h cold season	1.11 (0.93, 1.32)

*Studies included from Figure 6-24..

Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Age groups of study populations were not specified or were adults with the exception of <u>Villeneuve et al. (2006a</u>), which included only individuals aged 65+, and <u>Chan et al. (2006</u>), which included only individuals aged 50+. Studies listed in alphabetical order.

Warm season defined as: May-September (<u>Halonen et al., 2009</u>), and April-September (<u>Larrieu et al., 2007</u>; <u>Villeneuve et al., 2006a</u>). Cold season defined as: October-March (<u>Villeneuve et al., 2006a</u>).



Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Seasons depicted by colors: black: all year; red: warm season; light blue: cold season. Age groups of study populations were not specified or were adults with the exception of <u>Wong et al. (1999a</u>), which included only individuals aged 65+. Studies organized by outcome and season and then listed in descending order of publication date.

Figure 6-25 Effect estimate (95% CI) per increment ppb increase in ozone for arrhythmia and dysrhythmia ED visits or hospital admissions.

Table 6-39Effect estimate (95% CI) per increment ppb increase in ozone for
arrhythmia and dysrhythmia ED visits or hospital admissions for
studies presented in Figure 6-25.

Study	Location	Outcome	Averaging Time	Effect Estimate (95% CI)
Buadong et al. (2009)	Bangkok, Thailand	Arrhythmia	1-h	0.99 (0.95, 1.04)
Halonen et al. (2009)	Helsinki, Finland	Arrhythmia	8-h max warm season	1.04 (0.80, 1.35)
Peel et al. (2007)	Atlanta, GA	Dysrhythmia	8-h warm season	1.01 (0.97, 1.06)
Poloniecki et al. (1997)	London, England	Arrhythmia	8-h	1.02 (0.96, 1.07)
<u>Stieb et al. (2009</u>)	7 Canadian cities	Dysrhythmia	24-h	1.02 (0.95, 1.09)
<u>Wong et al. (1999a</u>)	Hong Kong	Arrhythmia	24-h	1.06 (0.99, 1.12)
			24-h warm season	1.10 (0.96, 1.26)
			24-h cold season	1.11 (1.01, 1.23)

*Studies included from Figure 6-25...

Note: Change in O_3 standardized to 20 ppb for 24-hour averaging period, 30 ppb for 8-hour averaging period, and 40 ppb for 1-hour averaging period (see Section 2.2). Ozone concentrations in ppb. Age groups of study populations were not specified or were adults with the exception of (<u>Wong et al., 1999a</u>), which included only individuals aged 65+. Studies listed in alphabetical order. Warm season defined as: March-October (<u>Peel et al., 2007</u>), May-October (<u>Wong et al., 1999a</u>) and May-September (<u>Halonen et al., 2009</u>). Cold season defined as: November-April (<u>Wong et al., 1999a</u>).

6.3.2.8 Cardiovascular Mortality

1	As discussed within this section (Section 6.3), epidemiologic studies provide inconsistent
2	evidence of an association between short-term O ₃ exposure and cardiovascular effects.
3	However, toxicological studies have demonstrated O ₃ -induced cardiovascular effects,
4	specifically enhanced atherosclerosis and ischemia, which could lead to death. The 2006
5	O3 AQCD provided evidence, primarily from single-city studies, of consistent positive
6	associations between short-term O3 exposure and cardiovascular mortality. Recent
7	multicity studies conducted in the U.S., Canada, and Europe further support the
8	association between short-term O_3 exposure and cardiovascular mortality.
9	As discussed in Section 6.2.7.2, the APHENA study (Katsouyanni et al., 2009) also
10	examined associations between short-term O3 exposure and mortality and found
11	consistent positive associations for cardiovascular mortality in all-year analyses.
12	However, in analyses restricted to the summer season, results were more variable with no
13	evidence of an association in the Canadian dataset in the population <75 years of age, and
14	evidence of associations persisting or increasing in magnitude in the Canadian
15	(population \geq 75 years of age), U.S., and European datasets. Additional multicity studies
16	from the U.S. (Zanobetti and Schwartz, 2008b), Europe (Samoli et al., 2009), Italy
17	(Stafoggia et al., 2010), and Asia (Wong et al., 2010) that conducted summer season

and/or all-year analyses provide additional support for an association between short-term O_3 exposure and cardiovascular mortality (Figure 6-36).

3 Of the studies evaluated, only the APHENA study (Katsouyanni et al., 2009) and the 4 Italian multicity study (Stafoggia et al., 2010) conducted an analysis of the potential for 5 copollutant confounding of the O₃-cardiovascular mortality relationship. In the European 6 dataset, when focusing on the natural spline model with 8 df/year (Section 6.2.7.2) and 7 lag 1 results in order to compare results across study locations (Section 6.6.2.1), 8 cardiovascular mortality risk estimates were robust to the inclusion of PM₁₀ in 9 copollutant models in all-year analyses with more variability in the Canadian and U.S. 10 datasets (i.e., cardiovascular O₃ mortality risk estimates were reduced or increased in 11 copollutant models). In summer season analyses, cardiovascular O₃ mortality risk 12 estimates were robust in the European dataset and attenuated but remained positive in the 13 U.S. dataset. Similarly, in the Italian multicity study (Stafoggia et al., 2010), which was 14 limited to the summer season, cardiovascular mortality risk estimates were robust to the 15 inclusion of PM_{10} in copollutant models. Based on the APHENA and Italian multicity 16 results, O_3 cardiovascular mortality risk estimates appear to be robust to inclusion of 17 PM₁₀ in copollutant models. However, in the U.S. and Canadian datasets there was 18 evidence that O_3 cardiovascular mortality risk estimates are moderately to substantially 19 sensitive (e.g., increased or attenuated) to PM₁₀. The mostly every-6th-day sampling 20 schedule for PM₁₀ in the Canadian and U.S. datasets greatly reduced their sample size 21 and limits the interpretation of these results.

6.3.2.9 Summary of Epidemiologic Studies

22	Overall, the available body of evidence examining the relationship between short-term
23	exposures to O ₃ concentrations and cardiovascular morbidity is inconsistent. Across
24	studies, different definitions, i.e., ICD-9 diagostic codes were used for both all-cause and
25	cause-specific cardiovascular morbidity (Table 6-35, Table 6-36, Table 6-37, Table 6-38,
26	and Table 6-39), which may contribute to inconsistency in results. However, within
27	diagnostic categories, no consistent pattern of association was found with O3. Generally,
28	the studies summarized in this section used nearest air monitors to assess O ₃
29	concentrations, with a few exceptions that used modeling or personal exposure monitors
30	(these exceptions were noted throughout the previous sections). The inconsistencies in
31	the associations observed between short-term O_3 and CVD morbidities are unlikely to be
32	explained by the different exposure assignment methods used (see Section 4.6). The wide
33	variety of biomarkers considered and the lack of consistency among definitions used for
34	specific cardiovascular disease endpoints (e.g., arrhythmias, HRV) make comparisons
35	across studies difficult. Despite the inconsistent evidence for an association between O_3

1

2

1	concentration and CVD morbidity, mortality studies indicate a consistent positive
2	association between short-term O3 exposure and cardiovascular mortality in multicity
3	studies and a multicontinent study.

6.3.3 Toxicology: Cardiovascular Effects

- 4In the previous O_3 AQCDs (U.S. EPA, 2006b, 1996a) experimental animal studies have5reported relatively few cardiovascular system alterations after exposure to O_3 and other6photochemical oxidants. The limited amount of research directed at examining7 O_3 -induced cardiovascular effects has primarily found alterations in heart rate (HR), heart8rhythm, and BP after O_3 exposure. Although O_3 induced changes in HR and core9temperature (T_{CO}) in a number of rat studies, these responses have not been reported or10extensively studied in humans exposed to O_3 and may be unique to rodents.
- 11 According to recent animal toxicology studies, short-term O₃ exposure induces vascular 12 oxidative stress and proinflammatory mediators, alters HR and HRV, and disrupts the 13 regulation of the pulmonary endothelin system (study details are provided in Table 6-40). 14 A number of these effects were variable between strains examined, suggesting a genetic 15 component to development of O₃ induced cardiovascular effects. Further, recent studies 16 provide evidence that extended O₃ exposure enhances the risk of ischemia-reperfusion 17 (I/R) injury and atherosclerotic lesion development. Still, few studies have investigated 18 the role of O₃ reaction products in these processes, but more evidence is provided for 19 elevated inflammatory and reduction-oxidation (redox) cascades known to initiate these 20 cardiovascular pathologies.

Heart Rate, Rhythm, and Heart Rate Variability

21	Studies (Arito et al., 1992; Arito et al., 1990; Uchiyama and Yokoyama, 1989;
22	Yokoyama et al., 1989; Uchiyama et al., 1986) report O ₃ exposure (0.2-1.0 ppm, 3 hours
23	to 3 days) in rats decreased T_{CO} , HR, and mean arterial pressure (MAP). In addition, O_3
24	exposure $(0.1 - 1.0 \text{ ppm}, 3 \text{ hours to } 3 \text{ days})$ in rats induced arrhythmias, including
25	increased PR interval and QRS complex, premature atrial contraction, and incomplete
26	A-V block (Arito et al., 1990; Yokoyama et al., 1989; Uchiyama et al., 1986). The effects
27	were more pronounced in adult and awake rats than in younger or sleeping animals,
28	whereas no sex-related differences were noted in these O_3 induced outcomes (Uchiyama
29	et al., 1986). However, these cardiovascular responses to O_3 , including decreased T_{CO} and
30	HR, could be attenuated by increased ambient temperatures and environmental stress and
31	exhibited adaptation (Watkinson et al., 2003; Watkinson et al., 1993). These studies
32	suggest that these responses to O_3 were the result of the rodent hypothermic response,

1 2 3 4 5	which serves as a physiological and behavioral defense mechanism to minimize the irritant effects of O_3 inhalation, (Iwasaki et al., 1998; Arito et al., 1997). As humans do not appear to exhibit decreased HR, MAP, and T_{CO} with routine environmental (Section <u>6.3.2</u>) or controlled laboratory (Section <u>6.3.1</u>) exposures to O_3 , caution must be used in extrapolating the results of these animal studies to humans.
6	Other studies have shown that O ₃ can increase BP in animal models. Rats exposed to
7	0.6 ppm O_3 for 33 days had increased systolic pressure and HR (<u>Revis et al., 1981</u>).
8	Increased BP triggers the release of atrial natriuretic factor (ANF), which has been found
9	in increased levels in the heart, lungs, and circulation of O_3 exposed (0.5 ppm) rats
10	(Vesely et al., 1994a, b, c). Exposures to high concentrations of O ₃ (1.0 ppm) have also
11	been found to lead to heart and lung edema (Friedman et al., 1983), which could be the
12	result of increased ANF levels. Thus, O3 may increase blood pressure and HR, leading to
13	increased ANF and tissue edema.
14	Recent studies report strain differences in HR and HRV in response to a 2-hour O ₃
15	pretreatment followed by exposure to carbon black (CB) in mice (C3H/HeJ [HeJ],
16	C57BL/6J [B6], and C3H/HeOuJ [OuJ]) (Hamade and Tankersley, 2009; Hamade et al.,
17	2008). These mice strains were chosen from prior studies on lung inflammatory and
18	hyperpermeability responses to be at increased risk (B6 and OuJ) or resistant (HeJ) to
19	O ₃ -induced health effects (<u>Kleeberger et al., 2000</u>). HR decreased during O ₃ pre-exposure
20	for all strains, but recovered during the CB exposure (Hamade et al., 2008). Percent
21	change in HRV parameters, SDNN (indicating total HRV) and rMSSD (indicating beat-
22	to-beat HRV), were increased in both C3H mice strains, but not B6 mice, during O_3
23	pre-exposure and recovered during CB exposure when compared to the filtered air group.
24	The two C3H strains differ by a mutation in the Toll-like receptor 4 (TLR4) gene, but
25	these effects did not seem to be related to this mutation since similar responses were
26	observed. Hamade et al. (2008) speculate that the B6 and C3H strains differ in
27	mechanisms of HR response after O_3 exposure between withdrawal of sympathetic tone
28	and increase of parasympathetic tone; however, no direct evidence for this conclusion
29	was reported. The strain differences observed in HR and HRV suggest that genetic
30	variability affects cardiac responses after acute air pollutant exposures.
31	Hamade and Tankersley (2009) continued this investigation of gene-environment
32	interactions on cardiopulmonary adaptation of O3 and CB induced changes in HR and
33	HRV using the previously described (Hamade et al., 2008) daily exposure scheme for 3
34	consecutive days. By comparing day-1 interim values it is possible to observe that O_3
35	exposure increased SDNN and rMSSD, but decreased HR in all strains. Measures of HR
36	and HRV in B6 and HeJ mice recovered to levels consistent with filtered air treated mice
37	by day 3; however, these responses in OuJ mice remained suppressed. B6 mice had no

- 1 change in respiratory rate (RR) after O₃ treatment, whereas HeJ mice on days 1 and 2 had 2 increased RR and OuJ mice on days 2 and 3 exhibited increased RR. V_T did not change 3 with treatment among the strains. Overall, B6 mice were mildly responsive with rapid 4 adaptation, whereas C3 mice were highly responsive with adaptation only in HeJ mice 5 with regards to changes in cardiac and respiratory responses. HR and HRV parameters 6 were not equally correlated with V_T and RR between the three mice strains, which 7 suggest that strains vary in the integration of the cardiac and respiratory systems. These 8 complex interactions could help explain variability in interindividual responses to air 9 pollution.
- 10 Hamade et al. (2010) expanded their investigation to explore the variation of these strain 11 dependent cardiopulmonary responses with age. As was observed previously, all 12 experimental mouse strains (B6, HeJ, and OuJ) exhibited decreased HR and increased 13 HRV after O_3 exposure. Younger O_3 -exposed mice had a significantly lower HR 14 compared to older exposed mice, indicating an attenuation of the bradycardic effect of O_3 15 with age. Younger mice also had a greater increase in rMSSD in HeJ and OuJ strains and 16 SDNN in HeJ mice. Conversely, B6 mice had a slightly greater increase in SDNN in 17 aged mice compared to the young mice. No change was observed in the magnitude of the O3 induced increase of SDNN in OuJ mice or rMSSD in B6 mice. The B6 and HeJ mice 18 19 genetically vary in respect to the nuclear factor erythroid 2-related factor 2 (Nrf-2). The 20 authors propose that the genetic differences between the mice strains could be altering the 21 formation of ROS, which tends to increase with age, thus modulating the changes in 22 cardiopulmonary physiology after O₃ exposure.
- 23 Strain and age differences in HR and heart function were further investigated in B6 and 24 129S1/SvlmJ (129) mice in response to a sequential O_3 and filtered air or CB exposure 25 (Tankersley et al., 2010). Young 129 mice showed a decrease in HR after O_3 or O_3 and 26 CB exposure. This bradycardia was not observed in B6 or older animals in this study, 27 suggesting a possible alteration or adaptation of the autonomic nervous system activity 28 with age. However, these authors did previously report bradycardia in similarly aged 29 young B6 mice (Hamade et al., 2010; Hamade and Tankersley, 2009; Hamade et al., 30 2008). Ozone exposure in 129 mice also resulted in an increase in left ventricular 31 chamber dimensions at end diastole (LVEDD) in young and old mice and a decrease in 32 left ventricular posterior wall thickness at end systole (PWTES) in older mice. The 33 increase in LVEDD caused a decrease in fractional shortening, which can be used as a 34 rough indicator of left ventricular function. Regression analysis revealed a significant 35 interaction between age and strain on HR and PWTES, which implies that aging affects 36 HR and heart function in response to O_3 differently between mouse strains.

Vascular Disease and Injury

1	A recent study in young mice (C57Bl/6) and rhesus monkeys examined the effects of
2	short-term O ₃ exposure (0.5 ppm, 1 or 5 days) on a number of cardiovascular endpoints
3	(Chuang et al., 2009). Mice exposed to O_3 for 5 days had increased HR as well as mean
4	and diastolic blood pressure. This is in contrast to the bradycardia that was reported in
5	18-20 week-old B6 mice treated with O ₃ , as described above (Hamade and Tankersley,
6	2009; Hamade et al., 2008). Increased blood pressure could be explained by the inhibition
7	in endothelial-dependent (acetylcholine) vasorelaxation from decreased bioavailability of
8	aortic nitric oxide (\cdot NO). Ozone caused a decrease in aortic NO _X (nitrite and nitrate
9	levels) and a decrease in total, but not phosphorylated, endothelial nitric oxide synthase
10	(eNOS). Ozone also increased vascular oxidative stress in the form of increased aortic
11	and lung lipid peroxidation (F2-isoprostane), increased aortic protein nitration (3-
12	nitrotyrosine), decreased aortic superoxide dismutase (SOD2) protein and activity, and
13	decreased aortic aconitase activity, indicating specific inactivation by O_2^- and ONOO ⁻ .
14	Mitochondrial DNA (mtDNA) damage was also used as a measure of oxidative and
15	nitrative stress in mice and infant rhesus monkeys exposed to O ₃ . Chuang et al. (2009)
16	observed that mtDNA damage accumulated in the lung and aorta of mice after 1 and
17	5 days of O_3 exposure and in the proximal and distal aorta of O_3 treated nonhuman
18	primates. Additionally, genetically hyperlipidemic mice exposed to O_3 (0.5 ppm) for
19	8 weeks had increased a ortic atherosclerotic lesion area (Section $7.3.1$), which may be
20	associated with the short-term exposure changes discussed. Overall, this study suggests
21	that O_3 initiates an oxidative environment by increasing O_2^- production, which leads to
22	mtDNA damage and \cdot NO consumption, known to perturb endothelial function (<u>Chuang</u>
23	et al., 2009). Endothelial dysfunction is characteristic of early and advanced
24	atherosclerosis and coincides with impaired vasodilation and blood pressure regulation.
25	Vascular occlusion resulting from atherosclerosis can block blood flow causing ischemia.
26	The restoration of blood flow in the vessel or reperfusion can cause injury to the tissue
27	from subsequent inflammation and oxidative damage. Perepu et al. (2010) observed that
28	O_3 exposure (0.8 ppm, 28 or 56 days) enhanced the sensitivity to myocardial I/R injury in
29	Sprague-Dawley rats while increasing oxidative stress levels and pro-inflammatory
30	mediators and decreasing production of anti-inflammatory proteins. Ozone was also
31	found to decrease the left ventricular developed pressure, rate of change of pressure
32	development, and rate of change of pressure decay while increasing left ventricular end
33	diastolic pressure in isolated perfused hearts. In this ex vivo heart model, O3 induced
34	oxidative stress by decreasing SOD enzyme activity and increasing malondialdehyde
35	levels. Ozone also elicited a proinflammatory state which was evident by an increase in
36	TNF- α and a decrease in the anti-inflammatory cytokine IL-10. <u>Perepu et al. (2010</u>)
37	concluded that O_3 exposure may result in a greater I/R injury.

Effects on Cardiovascular-Related Proteins

1	Increased BP, changes in HRV, and increased atherosclerosis may be related to increases
2	in the vasoconstrictor peptide, endothelin-1 (amino acids 1-21, ET-1[1-21]). Regulation of
3	the pulmonary endothelin system can be affected in rats by inhalation of PM (0, 5,
4	50 mg/m ³ , EHC-93) and O ₃ (Thomson et al., 2006; Thomson et al., 2005). Exposure to
5	either O_3 (0.8 ppm) or PM increased plasma ET-1 _[1-21] , ET-3 _[1-21] , and the ET-1 precursor
6	peptide, bigET-1. Increases in circulating $ET-1_{[1-21]}$ could be a result of a transient
7	increase in the gene expression of lung preproET-1 and endothelin converting enzyme-1
8	(ECE-1) immediately following inhalation of O ₃ or PM. These latter gene expression
9	changes (e.g., preproET-1 and ECE-1) were additive with co-exposure to O_3 and PM.
10	Conversely, preproET-3 decreased immediately after O_3 exposure, suggesting the
11	increase in ET-3[1-21] was not through de novo production. A recent study also found
12	increased ET-1 gene expression in the aorta of O ₃ -exposed rats (Kodavanti et al., 2011).
13	These rats also exhibited an increase in ET_BR after O_3 exposure; however, they did not
14	demonstrate increased biomarkers for vascular inflammation, thrombosis, or oxidation.
15	O_3 can oxidize protein functional groups and disturb the affected protein. For example,
15 16	O_3 can oxidize protein functional groups and disturb the affected protein. For example, the soluble plasma protein fibrinogen is oxidized by O_3 (0.01-0.03 ppm) in vitro, creating
16	the soluble plasma protein fibrinogen is oxidized by O_3 (0.01-0.03 ppm) in vitro, creating
16 17	the soluble plasma protein fibrinogen is oxidized by O_3 (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen
16 17 18	the soluble plasma protein fibrinogen is oxidized by O ₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (<u>Rosenfeld et al., 2009</u> ; <u>Rozenfeld et al., 2008</u>). In these studies, oxidized fibrinogen
16 17 18 19	the soluble plasma protein fibrinogen is oxidized by O_3 (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the
16 17 18 19 20	the soluble plasma protein fibrinogen is oxidized by O_3 (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers.
16 17 18 19 20 21	the soluble plasma protein fibrinogen is oxidized by O ₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers. Oxidized fibrinogen also developed the ability to self assemble creating fibrinogen
16 17 18 19 20 21 22	 the soluble plasma protein fibrinogen is oxidized by O₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers. Oxidized fibrinogen also developed the ability to self assemble creating fibrinogen aggregates that may play a role in thrombosis. Since O₃ does not readily translocate past
16 17 18 19 20 21 22 23	the soluble plasma protein fibrinogen is oxidized by O ₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers. Oxidized fibrinogen also developed the ability to self assemble creating fibrinogen aggregates that may play a role in thrombosis. Since O ₃ does not readily translocate past the ELF and pulmonary epithelium and fibrinogen is primarily a plasma protein, it is
 16 17 18 19 20 21 22 23 24 	the soluble plasma protein fibrinogen is oxidized by O ₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers. Oxidized fibrinogen also developed the ability to self assemble creating fibrinogen aggregates that may play a role in thrombosis. Since O ₃ does not readily translocate past the ELF and pulmonary epithelium and fibrinogen is primarily a plasma protein, it is uncertain if O ₃ would have the opportunity to react with plasma fibrinogen. However,
 16 17 18 19 20 21 22 23 24 25 	the soluble plasma protein fibrinogen is oxidized by O ₃ (0.01-0.03 ppm) in vitro, creating fibrinogen and fibrin aggregates, characteristically similar to defective fibrinogen (Rosenfeld et al., 2009; Rozenfeld et al., 2008). In these studies, oxidized fibrinogen retained the ability to form fibrin gels that are involved in coagulation, however the aggregation time increased and the gels were rougher than normal with thicker fibers. Oxidized fibrinogen also developed the ability to self assemble creating fibrinogen aggregates that may play a role in thrombosis. Since O ₃ does not readily translocate past the ELF and pulmonary epithelium and fibrinogen is primarily a plasma protein, it is uncertain if O ₃ would have the opportunity to react with plasma fibrinogen. However, fibrinogen can be released from the basolateral face of pulmonary epithelial cells during

Studies on Ozone Reaction Products

28Although toxicological studies have demonstrated O_3 -induced effects on the29cardiovascular system, it remains unclear if the mechanism is through a reflex response30or the result of effects from O_3 reaction products (U.S. EPA, 2006b, 1996a). Oxysterols31derived from cholesterol ozonation, such as β -epoxide and $5\beta, 6\beta$ -epoxycholesterol (and32its metabolite cholestan-6-oxo-3,5-diol), have been implicated in inflammation associated33with cardiovascular disease (Pulfer et al., 2005; Pulfer and Murphy, 2004). Two other34cholesterol ozonolysis products, atheronal-A and -B (e.g., cholesterol secoaldehyde),

1	have been found in human atherosclerotic plaques and shown in vitro to induce foam cell
2	formation and induce cardiomyocyte apoptosis and necrosis (<u>Sathishkumar et al., 2005;</u>
3	Wentworth et al., 2003); however, these products have not been found in the lung
4	compartment or systemically after O_3 exposure. The ability to form these cholesterol
5	ozonation products in the circulation in the absence of O_3 exposure complicates their
6	implication in O_3 induced cardiovascular disease.
0	implication in O3 induced cardiovascular disease.
7	Although it has been proposed that O_3 reaction products released after the interaction of
8	O_3 with ELF constituents (see Section <u>5.2.3</u>) on O_3 interaction with ELF) are responsible
9	for systemic effects, it is not known whether they gain access to the vascular space.
10	Alternatively, extrapulmonary release of diffusible mediators, such as cytokines or
11	endothelins, may initiate or propagate inflammatory responses in the vascular or systemic
12	compartments (Cole and Freeman, 2009) (Section 5.3.8). Ozone reacts within the lung to
13	amplify ROS production, induce pulmonary inflammation, and activate inflammatory
14	cells, resulting in a cascading proinflammatory state and extrapulmonary release of
15	diffusible mediators that could lead to cardiovascular injury.
16	A recent study that examined Ω reaction hyperaducts has shown that shalestered
10	A recent study that examined O_3 reaction byproducts has shown that cholesterol
	secoaldehyde (e.g., atheronal A) induces apoptosis in vitro in mouse macrophages (Gao
18	et al., 2009b) and cardiomyocytes (<u>Sathishkumar et al., 2009</u>). Additionally, atheronal-A
19	and -B has been found to induce in vitro macrophage and endothelial cell
20	proinflammatory events involved in the initiation of atherosclerosis (Takeuchi et al.,
21	<u>2006</u>). These O_3 reaction products when complexed with low density lipoprotein
22	upregulate scavenger receptor class A and induce dose-dependent macrophage
23	chemotaxis. Atheronal-A increases expression of the adhesion molecule, E-selectin, in
24	endothelial cells, while atheronal-B induces monocyte differentiation. These events
25	contribute to both monocyte recruitment and foam cell formation in atherosclerotic
26	vessels. It is unknown whether these O_3 reaction products gain access to the vascular
27	space from the lungs. Alternative explanations include the extrapulmonary release of
28	diffusible mediators that may initiate or propagate inflammatory responses in the vascular
29	or systemic compartments.

Study ^a *	Model	O₃ (ppm)	Exposure Duration	Effects
<u>Chuang et al. (2009)</u>	Mice; C57Bl/6; M; 6 weeks	0.5	1 or 5 days, 8-h/day	Increased HR and blood pressure. Initiated an oxidative environment by increasing vascular O_2^- production, which lead to mtDNA damage and \cdot NO consumption, known to perturb endothelial function.
	Monkey; rhesus <i>Macaca mulatta</i> ; M; Infant (180 days old)	0.5	5 days, 8-h/day	Increased aortic mtDNA damage.
<u>Perepu et al. (2010)</u>	Rat; Sprague-Dawley; 50-75 g	0.8	28 days, 8-h/day	Enhanced the sensitivity to myocardial I/R injury while increasing oxidative stress and pro-inflammatory mediators and decreasing production of anti-inflammatory proteins.
Hamade et al. (2008)	Mice; C57BI/6J, C3H/HeJ, and C3H/HeOuJ; M; 18-20 weeks	0.6 (subsequent CB exposure, 536 µg/m ³)	2-h followed by 3 h of CB	Decreased HR. Strain differences observed in HRV suggest that genetic variability affects cardiac responses.
<u>Hamade and</u> <u>Tankersley (2009</u>)	Mice; C57BI/6J, C3H/HeJ, and C3H/HeOuJ; M; 18-20 weeks	0.6 (subsequent CB exposure, 536 μg/m ³)	3 days, 2-h/day followed by 3-h of CB	Strains varied in integration of the cardiac and respiratory systems, implications in interindividual variability. B6 mice were mildly responsive with rapid adaptation, whereas C3 mice were highly responsive with adaptation only in HeJ mice with regards to changes in cardiac and respiratory responses.
<u>Hamade et al. (2010)</u>	Mice; C57BI/6J, C3H/HeJ, and C3H/HeOuJ; M; 5 or 12 mo old	0.6 (subsequent CB exposure, 536 µg/m ³)	2-h followed by 3-h of CB	Aged mice exhibited attenuated changes in cardiopulmonary physiology after O_3 exposure. Genetic differences between mice strains could be altering formation of ROS, which tends to increase with age, thus modulating O_3 induced effects.
<u>Tankersley et al.</u> (2010)	Mice; C57Bl/6J, 129S1/SvlmJ; M/F; 5 or 18 mo old	0.6 (subsequent CB exposure, 556 µg/m ³)	2-h followed by 3-h of CB	Significant interaction between age and strain on HR and PWTES, which implies that aging affects the HR and function in response to O_3 differently between mouse strains.
<u>Thomson et al.</u> (2005)	Rat; Fischer-344; M; 200-250 g	0.4 or 0.8	4-h	Activation of the vasoconstricting ET system. Increased plasma ET-1 through higher production and slower clearance.
<u>Thomson et al.</u> (2006)	Rat; Fischer-344; M; 200-250 g	0.8	4-h	Increased plasma ET-3 not due to de novo synthesis, unlike ET-1.
<u>Kodavanti et al.</u> (2011)	Rat; Wistar; M; 10-12 weeks	0.5 or 1.0	2 days, 5-h/day	No changes to aortic genes of thrombosis, inflammation, or proteolysis, except ET-1 and ETBR (1.0 ppm).

Table 6-40Characterization of study details for Section 6.3.3.

^aResults from previous studies are presented in Table AX5-14 of the 2006 O_3 AQCD and Table 6-23 of the 1996 O_3 AQCD. *Study details for Section <u>6.3.3</u>^a

Summary of Toxicological Studies

1	Overall, animal studies suggest that O ₃ exposure may result in O ₃ induced cardiovascular
2	effects. Studies provide evidence for both increased and decreased HR, however it is
3	uncertain if O ₃ -induced bradycardia would also occur in humans or if it is due solely to a
4	rodent hypothermic response. Animal studies also provide evidence for increased HRV,
5	arrhythmias, vascular disease, and injury following short-term O3 exposure. In addition, a
6	series of studies highlight the role of gene-environment interactions and age in the
7	induction of effects and attenuation of responses to O ₃ exposure.
8	Biologically plausible mechanisms are present for the cardiovascular effects observed in
9	animal exposure studies, however there is a lack of coherence with controlled human
10	exposure and epidemiologic studies. Further discussion of the modes of action that may
11	lead to cardiovascular effects can be found in Section 5.3.8. Recent studies suggest that
12	O_3 exposure may disrupt both the NO ^{\cdot} and endothelin systems, which can result in an
13	increase in HR, HRV, and ANF. The observed bradycardia following O_3 exposure may
14	be the result of reflex reactions, including the trigeminocardiac reflex, evoked following
15	the stimulation of sensory receptors lining the nose and RT. These mechanisms of
16	parasympathetically-derived cardiac effects are described in more detail in Section 5.3.2.
17	Additionally, O ₃ may increase oxidative stress and vascular inflammation promoting the
18	progression of atherosclerosis and leading to increased susceptibility to I/R injury. As O ₃
19	reacts quickly with the ELF and does not translocate to the heart and large vessels,
20	studies suggest that the cardiovascular effects exhibited could be caused by reaction
21	byproducts of O3 exposure. However, direct evidence of translocation of O3 reaction
22	products to the cardiovascular system has not been demonstrated in vivo. Alternatively,
23	extrapulmonary release of diffusible mediators, such as cytokines or endothelins, may
24	initiate or propagate inflammatory responses in the vascular or systemic compartments
25	leading to the reported cardiovascular pathologies.

6.3.4 Summary and Causal Determination

26	In previous O ₃ reviews (U.S. EPA, 2006b, 1996a) very few studies examined the effect of
27	short-term O_3 exposure on the cardiovascular system. More recently, the body of
28	scientific evidence available that has examined the effect of O_3 on the cardiovascular
29	system has advanced, but overall still remains small.
30	Although limited in number, toxicological studies have provided evidence of Ω induced
30	Although limited in number, toxicological studies have provided evidence of O ₃ -induced
30 31	Although limited in number, toxicological studies have provided evidence of O ₃ -induced cardiovascular effects. Animal toxicological studies have reported enhanced I/R injury,

- 1 HRV is supported by a recent controlled human exposure study that also found increased 2 high frequency HRV, but not altered blood pressure, following O₃ exposure (Fakhri et al., 3 2009). Toxicological studies investigating the role of O_3 in heart rate regulation are 4 mixed with both bradycardic and tachycardic responses observed. However, these 5 changes in cardiac function provide preliminary evidence for O₃-induced modulation of 6 the autonomic nervous system leading to cardiovascular complications. It is still 7 uncertain how O_3 inhalation may cause systemic toxicity; however the cardiovascular 8 effects of O₃ found in animals correspond to the development and maintenance of an 9 extrapulmonary oxidative, proinflammatory environment that may result from pulmonary 10 inflammation.
- 11 The epidemiologic studies evaluated do not support the evidence of O_3 -induced 12 cardiovascular effects observed in the toxicological studies. This is highlighted by the 13 multiple studies that examined the association between short-term O_3 exposure and 14 cardiovascular-related hospital admissions and ED visits and other various cardiovascular 15 effects and found no evidence of a consistent relationship with O₃ exposure. Although 16 there is inconsistent evidence for O₃-induced cardiovascular morbidity in the 17 epidemiologic literature, single-city studies reviewed in the 2006 O_3 AOCD, and recent 18 multicity studies, and a multicontinent study demonstrate consistent positive associations 19 between short-term O_3 exposure and cardiovascular mortality. Additionally, O_3 mortality 20 associations were found to remain robust in copollutant models with PM. However, the 21 lack of coherence between the results from studies that examined associations between
- 21lack of coherence between the results from studies that examined associations between22short-term O3 exposure and cardiovascular morbidity and subsequently cardiovascular23mortality complicate the interpretation of the overall evidence for O3-induced24cardiovascular effects.
- 25 In conclusion, animal toxicological studies provide some evidence for O_3 -induced 26 cardiovascular effects, but the effects observed were not consistently supported by 27 controlled human exposure studies or epidemiologic studies. Although the toxicological 28 evidence provides initial support to the relatively strong body of evidence indicating 29 O₃-induced cardiovascular mortality, there is a lack of coherence with controlled human 30 exposure and epidemiologic studies of cardiovascular morbidity which together do not 31 support O_3 -induced cardiovascular effects. Thus, the overall body of evidence across 32 disciplines is suggestive of a causal relationship between relevant short-term 33 exposures to O₃ and cardiovascular effects.

6.4 Central Nervous System Effects

1	
1	The 2006 O_3 AQCD included toxicological evidence that acute exposures to O_3 are
2	associated with alterations in neurotransmitters, motor activity, short and long term
3	memory, and sleep patterns. Additionally, histological signs of neurodegeneration have
4	been observed. Reports of headache, dizziness, and irritation of the nose with O_3
5	exposure are common complaints in humans, and some behavioral changes in animals
6	may be related to these symptoms rather than indicative of neurotoxicity. Peterson and
7	Andrews (1963) and Tepper et al. (1983) showed that mice would alter their behavior to
8	avoid O_3 exposure. Murphy et al. (1964) and Tepper et al. (1982) showed that running-
9	wheel behavior was suppressed, and Tepper et al. (1985) subsequently demonstrated the
10	effects of a 6-hour exposure to O_3 on the suppression of running-wheel behavior in rats
11	and mice, with the lowest effective concentration being about $0.12 \text{ ppm } O_3$ in the rat and
12	about 0.2 ppm in the mouse. The suppression of active behavior by 6 hours of exposure
13	to 0.12 ppm O_3 has recently been confirmed by Martrette et al. (2011) in juvenile female
14	rats, and the suppression of three different active behavior parameters was found to
15	become more pronounced after 15 days of exposure. A table of studies examining the
16	effects of O_3 on behavior can be found on p 6-128 of the 1996 O_3 AQCD. Generally
17	speaking, transient changes in behavior in rodent models appear to be dependent on a
18	complex interaction of factors such as (1) the type of behavior being measured, with
19	some behaviors increased and others suppressed; (2) the factors motivating that behavior
20	(differences in reinforcement); and (3) the sensitivity of the particular behavior
21	(e.g., active behaviors are more affected than more sedentary behaviors). Many
22	behavioral changes are likely to result from avoidance of irritation, but more recent
23	studies indicate that O ₃ also directly affects the CNS.
24	Research in the area of O ₃ -induced neurotoxicity has notably increased over the past few
25	years, with the majority of the evidence coming from toxicological studies that examined
26	the association between O3 exposure, neuropathology, and neurobehavioral effects, and
27	more limited evidence from epidemiologic studies. In an epidemiologic study conducted
28	by Chen and Schwartz (2009), data from the NHANES III cohort was utilized to study
29	the relationship between long-term O ₃ exposure (mean annual O ₃ concentration of
30	26.5 ppb) and neurobehavioral effects among adults aged 20-59 years. The authors
31	observed an association between annual exposure to O_3 and tests measuring coding
32	ability and attention/short-term memory. Each 10-ppb increase in annual O ₃ levels
33	corresponded to an aging-related cognitive performance decline of 3.5 years for coding
34	ability and 5.3 years for attention/short-term memory. These associations persisted in
35	both crude and adjusted models. There was no association between annual O_3
36	concentrations and reaction time tests. The authors conclude that overall there is a
37	positive association between O ₃ exposure and reduced performance on neurobehavioral

- 1 tests. Although Chen and Schwartz (2009) is a long-term exposure study, it is included in 2 this section because it is the first epidemiologic study to demonstrate that exposure to 3 ambient O_3 is associated with decrements in neurocognitive tests related to memory and 4 attention in humans. This epidemiologic evidence of an effect on the CNS due to 5 exposure to ambient concentrations of O_3 is coherent with animal studies demonstrating 6 that exposure to O_3 can produce a variety of CNS effects including behavioral deficits, 7 morphological changes, and oxidative stress in the brains of rodents. In these rodent 8 studies, interestingly, CNS effects were reported at O_3 concentrations that were generally 9 lower than those concentrations commonly observed to produce pulmonary or cardiac 10 effects in rats.
- 11 A number of new studies demonstrate various perturbations in neurologic function or 12 histology, including changes similar to those observed with Parkinson's and Alzheimer's 13 disease pathologies occurring in similar regions of the brain (Table 6-41). Many of these 14 include exposure durations ranging from short-term to long-term, and as such are 15 discussed here and in Chapter 7 with emphasis on the effects resulting from exposure 16 durations relevant to the respective chapter. Several studies assess short- and long-term 17 memory acquisition via passive avoidance behavioral testing and find decrements in test 18 performance after O_3 exposure, consistent with the aforementioned observation made in 19 humans by Chen and Schwartz (2009). Impairment of long-term memory has been 20 previously described in rats exposed to 0.2 ppm O_3 for 4 hours (Rivas-Arancibia et al., 21 1998) and in other studies of 4-hour exposures at concentrations of 0.7 to 1 ppm (Dorado-22 Martinez et al., 2001; Rivas-Arancibia et al., 2000; Avila-Costa et al., 1999). More 23 recently, statistically significant decreases in both short and long-term memory were

observed in rats after 15 days of exposure to 0.25 ppm O₃ (Rivas-Arancibia et al., 2010).

25 The central nervous system is very sensitive to oxidative stress, due in part to its high 26 content of polyunsaturated fatty acids, high rate of oxygen consumption, and low 27 antioxidant enzyme capacity. Oxidative stress has been identified as one of the 28 pathophysiological mechanisms underlying neurodegenerative disorders such as 29 Parkinson's and Alzheimer's disease, among others (Simonian and Coyle, 1996). It is 30 also believed to play a role in altering hippocampal function, which causes cognitive 31 deficits with aging (Vanguilder and Freeman, 2011). A particularly common finding in 32 studies of O₃-exposed rats is lipid peroxidation in the brain, especially in the 33 hippocampus, which is important for higher cognitive function including contextual 34 memory acquisition. Performance in passive avoidance learning tests is impaired when 35 the hippocampus is injured, and the observed behavioral effects are well correlated with 36 histological and biochemical changes in the hippocampus, including reduction in spine 37 density in the pyramidal neurons (Avila-Costa et al., 1999), lipoperoxidation (Rivas-38 Arancibia et al., 2010; Dorado-Martinez et al., 2001), progressive neurodegeneration, and

24

- 1 activated and phagocytic microglia (Rivas-Arancibia et al., 2010). The hippocampus is 2 also one of the main regions affected by age-related neurodegenerative diseases, 3 including Alzheimer's disease, and it may be more sensitive to oxidative damage in aged 4 rats. In a study of young (47 days) and aged (900 days) rats exposed to 1 ppm O_3 for 4 h, 5 O₃-induced lipid peroxidation occurred to a greater extent in the striatum of young rats, 6 whereas it was highest in the hippocampus in aged rats (Rivas-Arancibia et al., 2000). 7 Martínez-Canabal and Angora-Perez (2008) showed exposure of rats to 0.25 ppm, 8 4h/day, for 7, 15, or 30 days increased lipoperoxides in the hippocampus. This effect was 9 observed at day 7 and continued to increase with time, indicating cumulative oxidative damage. O₃-induced changes in lipid peroxidation, neuronal death, and COX-2 positive 10 11 cells in the hippocampus could be significantly inhibited by daily treatment with growth 12 hormone (GH), which declines with age in most species. The protective effect of GH on 13 -induced oxidative stress was greatest at 15 days of exposure and was non-significant at 14 day 30. Consistent with these findings, lipid peroxidation in the hippocampus of rats was 15 observed to increase significantly after a 30-day exposure to 0.25 ppm, but not after a 16 single 4-hour exposure to the same concentration (Mokoena et al., 2010). However, 17 4 hours of exposure was sufficient to cause significant increases in lipid peroxidation 18 when the concentration was increased to 0.7 ppm, and another study observed lipid 19 peroxidation after a 4-hour exposure to 0.4 ppm (Dorado-Martinez et al., 2001). 20 Other commonly affected areas of the brain include the striatum, substantia nigra, 21 cerebellum, olfactory bulb, and frontal/prefrontal cortex. The striatum and substantia nigra are particularly sensitive to oxidative stress because the metabolism of dopamine,
- 22 23 central to their function, is an oxidative process perturbed by redox imbalance. Oxidative 24 stress has been implicated in the premature death of substantia nigra dopamine neurons in 25 Parkinson's disease. Angoa-Pérez et al. (2006) have shown progressive lipoperoxidation 26 in the substantia nigra and a decrease in nigral dopamine neurons in ovariectomized 27 female rats exposed to 0.25 ppm O_3 , 4h/day, for 7, 15, or 30 days. Estradiol, an 28 antioxidant, attenuated O₃-induced oxidative stress and nigral neuronal death, and the 29 authors note that in humans, estrogen therapy can ameliorate symptoms of Parkinson's 30 disease, which is more prevalent in men. Progressive oxidative stress has also been 31 observed in the striatum and substantia nigra of rats after 15 and 30 days of exposure to 32 0.25 ppm O₃ for 4 h/day, along with a loss of dopaminergic neurons from the substantia 33 nigra (Pereyra-Muñoz et al., 2006). Decreases in motor activity were also observed at 15 34 and 30 days of exposure, consistent with other reports (Martrette et al., 2011; Dorado-35 Martinez et al., 2001). Using a similar O_3 exposure protocol, Santiago-López et al. (2010) 36 also observed a progressive loss of dopaminergic neurons within the substantia nigra, 37 accompanied by alterations in the morphology of remaining cells and an increase in p53 38 levels and nuclear translocation.

1	The olfactory bulb also undergoes oxidative damage in O ₃ exposed animals, in some
2	cases altering olfactory-dependent behavior. Lipid peroxidation was observed in the
3	olfactory bulbs of ovariectomized female rats exposed to 0.25 ppm O ₃ (4 h/day) for 30 or
4	60 days (Guevara-Guzmán et al., 2009). O ₃ also induced decrements in a selective
5	olfactory recognition memory test, and the authors note that early deficits in odor
6	perception and memory are components of human neurodegenerative diseases. The
7	decrements in olfactory memory were not due to damaged olfactory perception based on
8	other tests. However, deficits in olfactory perception emerged with longer exposures
9	(discussed in Chapter 7). As with the study by Angoa-Pérez et al. (2006) described
10	above, a protective effect for estradiol was demonstrated for both lipid peroxidation and
11	olfactory memory defects. The role of oxidative stress in memory deficits and associated
12	morphological changes has also been demonstrated via attenuation by other antioxidants
13	as well, such as α -tocopherol (<u>Guerrero et al., 1999</u>) and taurine (<u>Rivas-Arancibia et al.</u> ,
14	2000). It is unclear how persistent these effects might be. One study of acute exposure,
15	using 1 ppm O_3 for 4 hours, observed morphological changes in the olfactory bulb of rats
16	at 2 hours, and 1 and 10 days, but not 15 days, after exposure (Colín-Barenque et al.,
17	<u>2005</u>).
18	Other acute studies also report changes in the CNS. Lipid peroxidation was observed in
19	multiple regions of the brain after a 1- to 9-hour exposure to 1 ppm O_3 (Escalante-
20	Membrillo et al., 2005). Ozone has also been shown to alter gene expression of
21	endothelin-1 (pituitary) and inducible nitric oxide synthase (cerebral hemisphere) after a
22	single 4-hour exposure to 0.8 ppm O_3 , indicating potential cerebrovascular effects. This
23	concentration-dependent effect was not observed at 0.4 ppm O_3 (Thomson et al., 2007).
24	Vascular endothelial growth factor was upregulated in astroglial cells in the central
25	respiratory areas of the brain of rats exposed to 0.5 ppm O_3 for 3 hours (Araneda et al.,
26	2008). The persistence of CNS changes after a single exposure was also examined and
27	the increase in vascular endothelial growth factor was present after a short (3 hours)
28	recovery period. Thus, there is evidence that O ₃ -induced CNS effects are both
29	concentration- and time-dependent.
30	Because O_3 can produce a disruption of the sleep-wake cycle (U.S. EPA, 2006b), Alfaro-
31	Rodríguez and González-Piña (2005) examined whether acetylcholine in a region of the
32	brain involved in sleep regulation was altered by O ₃ . After a 24-hour exposure to 0.5 ppm
33	O_3 , the acetylcholine concentration in the medial preoptic area was decreased by 58% and
34	strongly correlated with a disruption in paradoxical sleep. Such behavioral-biochemical
35	effects of O ₃ are confirmed by a number of studies which have demonstrated
36	morphological and biochemical changes in rats.

1	CNS effects have also been demonstrated in newborn and adult rats whose only exposure
2	to O_3 occurred in utero. Several neurotransmitters were assessed in male offspring of
3	dams exposed to 1 ppm O_3 during the entire pregnancy (<u>Gonzalez-Pina et al., 2008</u>). The
4	data showed that catecholamine neurotransmitters were affected to a greater degree than
5	indole-amine neurotransmitters in the cerebellum. CNS changes, including behavioral,
6	cellular, and biochemical effects, have also been observed after in utero exposure to
7	0.5 ppm O ₃ for 12 h/day from gestational days 5-20 (Boussouar et al., 2009). Tyrosine
8	hydroxylase labeling in the nucleus tractus solatarius was increased after in utero
9	exposure to O_3 whereas Fos protein labeling did not change. When these offspring were
10	challenged by immobilization stress, neuroplasticity pathways, which were activated in
11	air-exposed offspring, were inhibited in O_3 -exposed offspring. Although an O_3 exposure
12	C-R was not studied in these two in utero studies, it has been examined in one study.
13	Santucci et al. (2006) investigated behavioral effects and gene expression after in utero
14	exposure of mice to as little as 0.3 ppm O_3 . Increased defensive/submissive behavior and
15	reduced social investigation were observed in both the 0.3 and 0.6 ppm O_3 groups.
16	Changes in gene expression of brain-derived neurotrophic factor (BDNF, increased in
17	striatum) and nerve growth factor (NGF, decreased in hippocampus) accompanied these
18	behavioral changes. Thus, these three studies demonstrate that CNS effects can occur as a
19	result of in utero exposure to O_3 , and although the mode of action of these effects is not
20	known, it has been suggested that circulating lipid peroxidation products may play a role
21	(Boussouar et al., 2009). Importantly, these CNS effects occurred in rodent models after
22	in utero only exposure to relevant concentrations of O_3 .

Study	Model	O₃ (ppm)	Exposure Duration	Effects
Martrette et al. (2011)	Rat; Wistar; F; Weight: 152g; 7 weeks old	0.12	1-15 days, 6 h/day	Significant decrease in rearing, locomotor activity, and jumping activity at day 1, with a further decrease in these activities by day 15.
Angoa-Pérez et al. (2006)	Rat; Wistar; F; Weight: 300g; ovariectomized	0.25	7 to 60 days, 4-h/day, 5 days/week	Progressive lipid peroxidation and loss of tyrosine hydrolase-immunopositive neurons in the substantia nigra starting at 7 days.
<u>Guevara-Guzmán et al.</u> (2009)	Rat; Wistar; F; 264g; ovariectomized	0.25	30 and 60 days, 4h/day	Estradiol treatment protected against lipid peroxidation and decreases in estrogen receptors and dopamine β -hydroxylase in olfactory bulbs along with deficits in olfactory recognition memory.
Martínez-Canabal and Angora-Perez (2008)	Rat; Wistar; M; Weight: 300g	0.25	7 to 30 days, 4-h/day	Growth hormone inhibited O ₃ -induced increases in lipoperoxidation and COX-2 positive cells in the hippocampus.
Pereyra-Muñoz et al. (2006)	Rat; Wistar; M; 250-300g	0.25	15 and 30 days, 4-h/day	Decreased motor activity, increased lipid peroxidation, altered morphology, and loss of dopamine neurons in substantia nigra and striatum, increased expression of DARPP- 32, iNOS, and SOD.
Rivas-Arancibia et al. (2010)	Rat; Wistar; M; 250-300g	0.25	15 to 90 days, 4-h/ day	Ozone produced significant increases in lipid peroxidation in the hippocampus, and altered the number of p53 positive immunoreactive cells, activated and phagocytic microglia cells, GFAP immunoreactive cells, and doublecortine cells, and short- and long-term memory- retention latency.
Santiago-López et al. (2010)	Rat; Wistar; M; 250-300g	0.25	15, 30, and 60 days, 4-h/day	Progressive loss of dopamine reactivity in the substantia nigra, along with morphological changes. Increased p53 levels and nuclear translocation.
Thomson et al. (2007)	Rat; Fischer-344; M; 200-250g	0.4; 0.8	4-h; assays at 0 and 24 h postexposure	At 0.8 ppm, O_3 produced rapid perturbations in the ET-NO pathway gene expression in the brain. Ozone induced a small but significant time- and concentration- dependent increase in prepro-endothelin-1 mRNA levels in the cerebral hemisphere and pituitary, whereas TNFa and iNOS mRNA levels were decreased at 0 h and unchanged or increased, respectively, at 24 h.
<u>Alfaro-Rodríguez and</u> <u>González-Piña (2005</u>)	Rat; Wistar; M; 292g	0.5	24-h	During the light phase, O_3 caused a significant decrease in paradoxical sleep accompanied by a significant decrease in Ach levels in the hypothalamic medial preoptic area. The same effects occurred during the dark phase exposure to O_3 in addition to a significant increase in slowwave sleep and decrease in wakefulness.
Araneda et al. (2008)	Rats; Sprague- Dawley; M; 280- 320g	0.5	3-h (measurements taken at 0 h and 3 h after exposure)	Ozone upregulated VEGF in astroglial cells located in the respiratory center of the brain. VEGF co-located with IL-6 and TNF in cells near blood vessel walls, and blood vessel area was markedly increased.

Table 6-41Central nervous system and behavioral effects of short-term ozone
exposure in rats

Study	Model	O₃ (ppm)	Exposure Duration	Effects
Boussouar et al. (2009)	Rat; Sprague- Dawley; M; adult offspring of prenatally exposed dams; 403-414g	0.5	From embryonic day E5 to E20 for 1-h/day; immobilization stress	Prenatal O_3 exposure had a long term impact on the nucleus tractus solitarius of adult rats, as revealed during immobilization stress.
Soulage et al. (2004)	Rat; Sprague- Dawley; M; Approx. 7 weeks old	0.7	5-h	Ozone produced differential effects on peripheral and central components of the sympatho-adrenal system. While catecholamine biosynthesis was increased in portions of the brain, the catecholamine turnover rate was significantly increased in the heart and cerebral cortex and inhibited in the lung and striatum.
Calderón Guzmán et al. (2006); (2005)	Rat; Wistar; M; 21 days old; well-nourished and malnourished groups	0.75	15 successive days for 4-h/day	A significant decrease in body weight was observed in both well nourished (WN) and malnourished (MN) rats after O ₃ exposure. Localized ATPase, TBARS, and GSH levels changed in response to O ₃ in certain brain areas and the O ₃ -induced changes were dependent on nutritional condition.
Colín-Barenque et al. (2005)	Rats; Wistar; M; 250-300g	1.0	4-h; assays at 2-h, 24-h, 10 days, and 15 days after exposure	A significant loss of dendritic spines in granule cells of the olfactory bulb occurred at 2 hrs to 10 days after exposure. Cytological and ultrastructural changes returned towards normal morphology by 15 days.
Escalante-Membrillo et al. (2005)	Rats; Wistar; M; 280-320g	1.0	1-, 3-, 6-, or 9-h	Significant increases in TBARS occurred in hypothalamus, cortex, striatum, midbrain, thalamus, and pons. Partial but significant recovery was observed by 3 h after the 9 h exposure.
Gonzalez-Pina et al. (2008)	Rat; Wistar; M;	1	12-h/day, 21 days of gestation; assays at 0, 5, & 10 days postnatal	Prenatal O_3 exposure produced significant decreases in cerebellar monoamine but not indolamine content at 0 and 5 days after birth with a partial recovery by 10 days. 5- hydroxy-indole-acetic acid levels were significantly increased at 10 days.

6.4.1 Neuroendocrine Effects

1	According to the 2006 O ₃ AQCD, early studies suggested an interaction of O ₃ with the
2	pituitary-thyroid-adrenal axis, because thyroidectomy, hypophysectomy, and
3	adrenalectomy protected against the lethal effects of O ₃ . Concentrations of 0.7-1.0 ppm
4	O_3 for a 1-day exposure in male rats caused changes in the parathyroid, thymic atrophy,
5	decreased serum levels of thyroid hormones and protein binding, and increased prolactin.
6	Increased toxicity to O ₃ was reported in hyperthyroid rats and T3 supplementation was
7	shown to increase metabolic rate and pulmonary injury in the lungs of O ₃ -treated animals.
8	The mechanisms by which O ₃ affects neuroendocrine function are not well understood,
9	but previous work suggests that high ambient levels of O ₃ can produce marked neural
10	disturbances in structures involved in the integration of chemosensory inputs, arousal,

1	and motor control, effects that may be responsible for some of the behavioral effects seen
2	with O_3 exposure. A more recent study exposing immature female rats to 0.12 ppm O_3
3	demonstrated significantly increased serum levels of the thyroid hormone free T ₃ after
4	15 days of exposure, whereas free T_4 was unchanged (<u>Martrette et al., 2011</u>). These
5	results are in contrast to those previously presented whereby 1 ppm O_3 for 1 day
6	significantly decreased T_3 and T_4 (<u>Clemons and Garcia, 1980</u>), although comparisons are
7	made difficult by highly disparate exposure regimens along with sex differences.
8	Martrette et al. (2011) also demonstrated significantly increased corticosterone levels
9	after 15 days of exposure, suggesting a stress related response.

6.4.2 Summary and Causal Determination

10	In rodents, O_3 exposure has been shown to cause physicochemical changes in the brain
11	indicative of oxidative stress and inflammation. Newer toxicological studies add to earlier
12	evidence that acute exposures to O_3 can produce a range of effects on the central nervous
12	system and behavior. Previously observed effects, including neurodegeneration,
13	alterations in neurotransmitters, short and long term memory, and sleep patterns, have
15	
	been further supported by recent studies. In instances where pathology and behavior are
16	both examined, animals exhibit decrements in behaviors tied to the brain regions or
17	chemicals found to be affected or damaged. For example, damage in the hippocampus,
18	which is important for memory acquisition, was correlated with impaired performance in
19	tests designed to assess memory. Thus the brain is functionally affected by O ₃ exposure.
20	The single epidemiologic study conducted showed an association between O_3 exposure
21	and memory deficits in humans as well, albeit on a long-term exposure basis. Notably,
22	exposure to O_3 levels as low as 0.25 ppm for 7 days has resulted in progressive
23	neurodegeneration and deficits in both short and long-term memory in rodents.
24	Examination of changes in the brain at lower exposure concentrations or at 0.25 ppm for
25	shorter durations has not been reported, but 0.12 ppm O_3 has been shown to alter
26	behavior. It is possible that some behavioral changes may reflect avoidance of irritation
27	as opposed to functional changes in brain morphology or chemistry, but in many cases
28	functional changes are related to oxidative stress and damage. In some instances, changes
29	were dependent on the nutritional status of the rats (high versus low protein diet). For
30	example, O ₃ produced an increase in glutathione in the brains of rats fed the high protein
31	diet but decreases in glutathione in rats fed low protein chow (Calderón Guzmán et al.,
32	2006). The hippocampus, one of the main regions affected by age-related
33	neurodegenerative diseases, appears to be more sensitive to oxidative damage in aged rats
34	(Rivas-Arancibia et al., 2000), and growth hormone, which declines with age in most
35	species, may be protective (Martínez-Canabal and Angora-Perez, 2008). Developing

1	animals may also be sensitive, as changes in the CNS, including biochemical, cellular,
2	and behavioral effects, have been observed in juvenile and adult animals whose sole
3	exposure occurred in utero, at levels as a low as 0.3 ppm. A number of studies
4	demonstrate O ₃ -induced changes that are also observed in human neurodegenerative
5	disorders such as Alzheimer's and Parkinson's disease, including signs of oxidative
6	stress, loss of neurons/neuronal death, reductions in dopamine levels, increased COX-2
7	expression, and increases in activated microglia in important regions of the brain
8	(hippocampus, substantia nigra).
9	Thus, evidence for neurological effects from epidemiologic and controlled human
10	exposure studies is lacking. However, the toxicological evidence for the impact of O_3 on
11	the brain and behavior is strong, and suggestive of a causal relationship between O_3
12	exposure and effects on the central nervous system.

6.5 Effects on Other Organ Systems

6.5.1 Effects on the Liver and Xenobiotic Metabolism

13	Early investigations of the effects of O_3 on the liver centered on xenobiotic metabolism,
14	and the prolongation of drug-induced sleeping time, which was observed at 0.1 ppm O_3
15	(Graham et al., 1981). In some species, only adults and especially females were affected.
16	In rats, high (1.0-2.0 ppm for 3 hours) acute O ₃ exposures caused increased production of
17	NO by hepatocytes and enhanced protein synthesis (Laskin et al., 1996; Laskin et al.,
18	1994). Except for the earlier work on xenobiotic metabolism, the responses occurred only
19	after very high acute O_3 exposures. One study, conducted at 1 ppm O_3 exposure, has been
20	identified (Last et al., 2005) in which alterations in gene expression underlying
21	O3-induced cachexia and downregulation of xenobiotic metabolism were examined. A
22	number of the downregulated genes are known to be interferon (IFN) dependent,
23	suggesting a role for circulating IFN. A more recent study by Aibo et al. (2010)
24	demonstrates exacerbation of acetaminophen-induced liver injury in mice after a single
25	6-hour exposure to 0.25 or 0.5 ppm O_3 . Data indicate that O_3 may worsen drug-induced
26	liver injury by inhibiting hepatic repair. The O3-associated effects shown in the liver are
27	thought to be mediated by inflammatory cytokines or other cytotoxic mediators released
28	by activated macrophages or other cells in the lungs (Laskin and Laskin, 2001; Laskin et
29	al., 1998; Vincent et al., 1996a). Recently, increased peroxidated lipids were detected in
30	the plasma of O_3 exposed animals (<u>Santiago-López et al., 2010</u>).
31	In summary, mediators generated by O_3 exposure may cause effects on the liver in
32	laboratory rodents. Ozone exposures as low as 0.1 ppm have been shown to affect

1	drug-induced sleeping time, and exposure to 0.25 ppm can exacerbate liver injury
2	induced by a common analgesic. However, very few studies at relevant concentrations
3	have been conducted, and no data from controlled human exposure or epidemiologic
4	studies are currently available. Therefore the collective evidence is inadequate to
5	determine if a causal relationship exists between short-term O_3 exposure and
6	effects on the liver and metabolism.

6.5.2 Effects on Cutaneous and Ocular Tissues

7	In addition to the lungs, the skin is highly exposed to O_3 and contains O_3 reactive targets
8	(polyunsaturated fatty acids) that can produce lipid peroxides. The 2006 O_3 AQCD (U.S.
9	EPA, 2006b) reported that although there is evidence of oxidative stress at near ambient
10	O3 concentrations, skin and eyes are only affected at high concentrations (greater than
11	1-5 ppm). Ozone exposure (0.8 ppm for 7 days) induces oxidative stress in the skin of
12	hairless mice, along with proinflammatory cytokines (Valacchi et al., 2009). A recent
13	study demonstrated that 0.25 ppm O_3 differentially alters expression of
14	metalloproteinases in the skin of young and aged mice, indicating that age may
15	potentially increase risk of oxidative stress (Fortino et al., 2007). In young mice, healing
16	of skin wounds is not significantly affected by O ₃ exposure (Lim et al., 2006). However,
17	exposure to 0.5 ppm O_3 for 6 h/day significantly delays wound closure in aged mice. As
18	with effects on the liver described above, the effects of O_3 on the skin and eyes have not
19	been widely studied, and information from controlled human exposure or epidemiologic
20	studies is not currently available. Therefore the collective evidence is inadequate to
21	determine if a causal relationship exists between short-term O_3 exposure and
22	effects on cutaneous and ocular tissues.

6.6 Mortality

6.6.1 Summary of Findings from 2006 Ozone AQCD

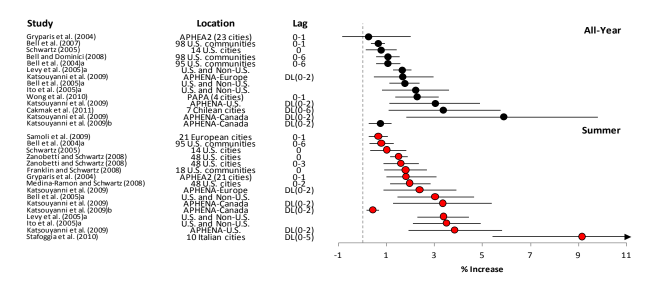
23	The 2006 O_3 AQCD reviewed a large number of time-series studies consisting of single-
24	and multicity studies, and meta-analyses. In the large U.S. multicity studies that
25	examined all-year data, summary effect estimates corresponding to single-day lags
26	ranged from a 0.5-1% increase in all-cause (nonaccidental) mortality per a standardized
27	unit increase in O_3 of 20 ppb for 24-h avg, 30 ppb for 8-h max, and 40 ppb for 1-h max as
28	discussed in Section 2.2. The association between short-term O_3 exposure and mortality
29	was substantiated by a collection of meta-analyses and international multicity studies.

1	The studies evaluated found some evidence for heterogeneity in O_3 mortality risk
2	estimates across cities and studies. Studies that conducted seasonal analyses, although
3	more limited in number, reported larger O3 mortality risk estimates during the warm or
4	summer season. Overall, the 2006 O_3 AQCD identified robust associations between
5	various measures of daily ambient O_3 concentrations and all-cause mortality, with
6	additional evidence for associations with cardiovascular mortality, which could not be
7	readily explained by confounding due to time, weather, or copollutants. However, it was
8	noted that multiple uncertainties remain regarding the O3-mortality relationship
9	including: the extent of residual confounding by copollutants; factors that modify the
10	O ₃ -mortality association; the appropriate lag structure for identifying O ₃ -mortality effects
11	(e.g., single-day lags versus distributed lag model); the shape of the O_3 -mortality C-R
12	function and whether a threshold exists; and the identification of susceptible populations.
13	Collectively, the 2006 O_3 AQCD concluded that "the overall body of evidence is highly
14	suggestive that O ₃ directly or indirectly contributes to non-accidental and
15	cardiopulmonary-related mortality."

6.6.2 Associations of Mortality and Short-Term Ozone Exposure

10	
17	

16 Recent studies that examined the association between short-term O₃ exposure and mortality further confirmed the associations reported in the 2006 O₃ AQCD. New 18 multicontinent and multicity studies reported consistent positive associations between 19 short-term O₃ exposure and all-cause mortality in all-year analyses, with additional 20 evidence for larger mortality risk estimates during the warm or summer months 21 (Figure 6-26; Table 6-42). These associations were reported across a range of ambient O_3 22 concentrations that were in some cases quite low (Table 6-43).



Note: Effect estimates are for a 40 ppb increase in 1-h max, 30 ppb increase in 8-h max, and 20 ppb increase in 24-h avg O_3 concentrations. An "a" represent multicity studies and meta-analyses from the 2006 O_3 AQCD. <u>Bell et al. (2005)</u>, <u>Ito et al. (2005)</u>, and <u>Levy et al. (2005)</u> used a range of lag days in the meta-analysis: Lag 0, 1, 2, or average 0-1 or 1-2; single-day lags from 0 to 3; and lag 0 and 1-2; respectively. A "b" represents risk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations (see explanation in Section <u>6.2.7.2</u>).

Figure 6-26 Summary of mortality risk estimates for short-term ozone exposure and all-cause (nonaccidental) mortality from all-year and summer season analyses.

Study*	Location	Lag	Avg Time	% Increase (95% CI)	
All-year					
Gryparis et al. (2004)	APHEA2 (23 cities)	0-1	1-h max	0.24 (-0.86, 1.98)	
<u>Bell et al. (2007)</u>	98 U.S. communities	0-1	24-h avg	0.64 (0.34, 0.92)	
Schwartz (2005a)	14 U.S. cities	0	1-h max	0.76 (0.13, 1.40)	
Bell and Dominici (2008)	98 U.S. communities	0-6	24-h avg	1.04 (0.56, 1.55)	
<u>Bell et al. (2004)</u> ^a	95 U.S. communities	0-6	24-h avg	1.04 (0.54, 1.55)	
<u>Levy et al. (2005</u>) ^a	U.S. and Non-U.S.		24-h avg	1.64 (1.25, 2.03)	
Katsouyanni et al. (2009)	APHENA-europe	DL(0-2)	1-h max	1.66 (0.47, 2.94)	
<u>Bell et al. (2005)</u> ^a	U.S. and Non-U.S.		24-h avg	1.75 (1.10, 2.37)	
(<u>Ito et al., 2005</u>) ^a	U.S. and Non-U.S.		24-h avg	2.20 (0.80, 3.60)	
(<u>Wong et al., 2010</u>)	PAPA (4 cities)	0-1	8-h avg	2.26 (1.36, 3.16)	
<u>Katsouyanni et al. (2009)</u>	APHENA-U.S.	DL(0-2)	1-h max	3.02 (1.10, 4.89)	
<u>Cakmak et al. (2011)</u>	7 Chilean cities	DL(0-6)	8-h max	3.35 (1.07, 5.75)	
Katsouyanni et al. (2009)	APHENA-Canada	DL(0-2)	1-h max	5.87 (1.82, 9.81)	
Katsouyanni et al. (2009) ^b	APHENA-Canada	DL(0-2)	1-h max	0.73 (0.23, 1.20)	
Summer					
<u>Samoli et al. (2009)</u>	21 European cities	0-1	8-h max	0.66 (0.24, 1.05)	
Bell et al. (2004) ^a	95 U.S. communities	0-6	24-h avg	0.78 (0.26, 1.30)	
<u>Schwartz (2005a)</u>	14 U.S. cities	0	1-h max	1.00 (0.30, 1.80)	
Zanobetti and Schwartz (2008a)	48 U.S. cities	0	8-h max	1.51 (1.14, 1.87)	
Zanobetti and Schwartz (2008b)	48 U.S. cities	0-3	8-h max	1.60 (0.84, 2.33)	
Franklin and Schwartz (2008)	18 U.S. communities	0	24-h avg	1.79 (0.90, 2.68)	
<u>Gryparis et al. (2004)</u>	APHEA2 (21 cities)	0-1	8-h max	1.80 (0.99, 3.06)	
Medina-Ramón and Schwartz (2008)	48 U.S. cities	0-2	8-h max	1.96 (1.14, 2.82)	
<u>Katsouyanni et al. (2009)</u>	APHENA-europe	DL(0-2)	1-h max	2.38 (0.87, 3.91)	
<u>Bell et al. (2005</u>) ^a	U.S. and Non-U.S.		24-h avg	3.02 (1.45, 4.63)	
Katsouyanni et al. (2009)	APHENA-Canada	DL(0-2)	1-h max	3.34 (1.26, 5.38)	
Katsouyanni et al. (2009)	APHENA-Canada	DL(0-2)	1-h max	0.42 (0.16, 0.67)	
Levy et al. (2005) ^a	U.S. and Non-U.S.		24-h avg	3.38 (2.27, 4.42)	
lto et al. (2005) ^a	U.S. and Non-U.S.		24-h avg	3.50 (2.10, 4.90)	
Katsouyanni et al. (2009)	APHENA-U.S.	DL(0-2)	1-h max	3.83 (1.90, 5.79)	
<u>Stafoggia et al. (2010)</u>	10 Italian cities	DL(0-5)	8-h max	9.15 (5.41, 13.0)	

Table 6-42Corresponding effect estimates for Figure 6-26.

*Studies included from Figure 6-26..

^aMulticity studies and meta-analyses from the 2006 O_3 AQCD. <u>Bell et al. (2005</u>)^a, <u>Ito et al. (2005</u>)^a, and <u>Levy et al. (2005</u>)^a used a range of lag days in the meta-analysis: Lag 0, 1, 2, or average 0-1 or 1-2; Single-day lags from 0-3; and Lag 0 and 1-2; respectively. ^bRisk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations (see explanation in Section <u>6.2.7.2</u>).

Table 6-43Range of mean and upper percentile ozone concentrations in
previous and recent multicity studies.

Study	Location	Years	Averaging Time	Mean Concentration (ppb) ^a	Upper Percentile Concentrations (ppb) ^a
<u>Gryparis et al.</u> (2004) ^b	23 European cities (APHEA2)	1990-1997	1-h max 8-h max	Summer: 1-h max: 44-117 8-h max: 30-99 Winter: 1-h max: 11-57 8-h max: 8-49	Summer: 1-h max: 62-173 8-h max: 57-154 Winter: 1-h max: 40-88 8-h max: 25-78
<u>Schwartz</u> (2005a) ^b	14 U.S. cities	1986-1993	1-h max	35.1-60	25th: 26.5-52 75th: 46.3-69
<u>Bell et al. (2004</u>)	95 U.S. communities (NMMAPS)	1987-2000	24-h avg	26.0	NR
<u>Bell et al. (2007</u>)	98 U.S. communities (NMMAPS)	1987-2000	24-h avg	26.0 ^d	NR
<u>Bell and</u> Dominici (2008)	98 U.S. communities (NMMAPS)	1987-2000 (All year and May-September)	24-h avg	All year: 26.8 May-September: 30.0	Maximum: All year: 37.3 May-September: 47.2
<u>Franklin and</u> Schwartz (2008)	18 U.S. communities	2000-2005 (May-September)	24-h avg	21.4-48.7	NR
<u>Katsouyanni et</u> al. (2009) ^{b,e}	NMMAPS 12 Canadian cities (APHEA2)	1987-1996 (Canada and U.S.) varied by city for Europe	1-h max	U.S.: 13.3-38.4 Canada: 6.7-8.4 Europe:18.3-41.9	75th: U.S.: 21.0-52.0 Canada: 8.7-12.5 Europe: 24.0-65.8
Medina-Ramón and Schwartz (2008) ⁶	48 U.S. cities	1989-2000 (May-September)	8-h max	16.1-58.8	NR
<u>Samoli et al.</u> (2009) [♭]	21 European cities (APHEA2)	1990-1997 (June-August)	8-h max	20.0-62.8	75th: 27.2-74.8
<u>Stafoggia et al.</u> (<u>2010</u>)	10 Italian cities	2001-2005 (April-September)	8-h max	41.2-58.9	75th: 47.0-71.6
<u>Cakmak et al.</u> (2011)	7 Chilean cities	1997-2007	8-h max	59.0-87.6	NR
<u>Wong et al.</u> (2010)	PAPA (4 cities)	1999-2003 (Bangkok) 1996-2002 (Hong Kong) 2001-2004 (Shanghai) 2001-2004 (Wuhan)	8-h avg	18.7-43.7	75th: 38.4 - 60.4 Max: 92.1 - 131.8
Zanobetti and Schwartz (2008b)	48 U.S. cities	1989-2000 (June-August)	8-h max	15.1-62.8	Max: 34.3-146.2 75th: 19.8-75.9

Study	Location	Years	Averaging Time	Mean Concentration (ppb) ^a	Upper Percentile Concentrations (ppb) ^a
Zanobetti and	48 U.S. citiesc	1989-2000	8-h max		Max:
Schwartz (2008a)		(Winter: Dec-Feb)		Winter: 16.5	Winter: 40.6
<u>(2008a</u>)		(Spring: March-May)		Spring: 41.6	Spring: 91.4
		(Summer: June-Aug)		Summer: 47.8	Summer: 103.0
		(Autumn: Sept-Nov)		Autumn: 33.5	Autumn: 91.2

 $^{a}O_{3}$ concentrations were converted to ppb if the study presented them as $\mu g/m^{3}$ by using the conversion factor of 0.51 assuming standard temperature (25° C) and pressure (1 atm).

^bStudy only reported median O₃ concentrations.

°Cities with less than 75% observations in a season excluded. As a result, 29 cities examined in winter, 32 in spring, 33 in autumn, and all 48 in the summer.

^dBell et al. (2007) did not report mean O₃ concentrations, however, it used a similar dataset as Bell et al. (2004) which consisted of 95 U.S. communities for 1987-2000. For comparison purposes the 24-h avg O₃ concentrations for the 95 communities from Bell et al. (2004) are reported here. ^eStudy did not present air quality data for the summer months.

1	In addition to examining the relationship between short-term O ₃ exposure and all-cause
2	mortality, recent studies attempted to address the uncertainties that remained upon the
3	completion of the 2006 O ₃ AQCD. As a result, given the robust associations between
4	short-term O_3 exposure and mortality presented across studies in the 2006 O_3 AQCD and
5	supported in the new multicity studies (Figure 6-26), the following sections primarily
6	focus on the examination of previously identified uncertainties in the O ₃ -mortality
7	relationship, specifically: O3 associations with cause-specific mortality, confounding, lag
8	structure (e.g., multiday effects and mortality displacement), effect modification
9	(i.e., sources of heterogeneity in risk estimates across cities); and the O3-mortality C-R
10	relationship. Focusing specifically on these uncertainties allows for a more detailed
11	characterization of the relationship between short-term O ₃ exposure and mortality.

6.6.2.1 Confounding

12	Recent epidemiologic studies examined potential confounders of the O ₃ -mortality
13	relationship. These studies specifically focused on whether PM and its constituents or
14	seasonal trends confounded the association between short-term O ₃ exposure and
15	mortality.

Confounding by PM and PM Constituents

16 An important question in the evaluation of the association between short-term O₃ 17 exposure and mortality is whether the relationship is confounded by particulate matter, 18 particularly the PM chemical components that are found in the "summer haze" mixture 19 which also contains O_3 . However, because of the temporal correlation among these PM 20 components and O₃, and their possible interactions, the interpretation of results from

copollutant models that attempt to disentangle the health effects associated with each
 pollutant is challenging. Further complicating the interpretation of copollutant results, at
 times, is the every-3rd or -6th day PM sampling schedule employed in most locations,
 which limits the number of days where both PM and O₃ data is available.

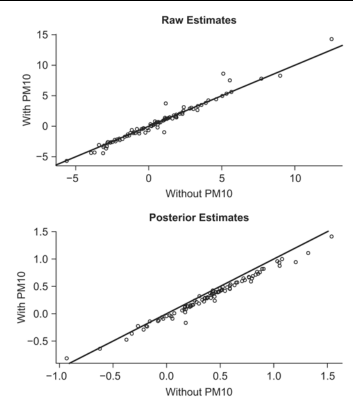
5 The potential confounding effects of PM₁₀ and PM_{2.5} on the O₃-mortality relationship 6 were examined by Bell et al. (2007) using data on 98 U.S. urban communities for the 7 years 1987-2000 from the National Morbidity, Mortality, and Air Pollution Study 8 (NMMAPS). In this analysis the authors included PM as a covariate in time-series 9 models, and also examined O3-mortality associations on days when O3 concentrations 10 were below a specified value. This analysis was limited by the small fraction of days when both PM and O₃ data were available, due to the every-3rd - or 6th -day sampling 11 12 schedule for the PM indices, and the limited amount of city-specific data for PM_{25} 13 because it was only collected in most cities since 1999. As a result, of the 91 14 communities with $PM_{2.5}$ data, only 9.2% of days in the study period had data for both O_3 15 and PM_{2.5}, resulting in the use of only 62 communities in the PM_{2.5} analysis. An 16 examination of the correlation between PM (PM_{10} and $PM_{2.5}$) and O_3 across various strata 17 of daily PM₁₀ and PM_{2.5} concentrations found that neither PM size fraction was highly 18 correlated with daily O₃ concentrations across any of the strata examined. These results 19 were also observed when using 8-h max and 1-h max O₃ exposure metrics. National and 20 community-specific effect estimates of the association between short-term O_3 exposure 21 and mortality were robust to inclusion of PM_{10} or $PM_{2.5}$ in time-series models through the 22 range of O_3 concentrations (i.e., <10 ppb, 10-20, 20-40, 40-60, 60-80, and >80 ppb). 23 Even with the small number of days in which both $PM_{2.5}$ and O_3 data was available, the 24 percent increases in nonaccidental deaths per 10 ppb increase 24-h avg O₃ concentrations 25 at lag 0-1 day were 0.22% (95% CI: -0.22, 0.65) without PM_{2.5} and 0.21% (95% CI: 26 -0.22, 0.64) with $PM_{2.5}$ in 62 communities.

27 Although strong correlations between PM and O₃ were not reported by Bell et al. (2007) 28 the patterns observed suggest regional differences in their correlation (Table 6-44). Both 29 PM_{10} and $PM_{2.5}$ show positive correlations with O₃ in the Industrial Midwest, Northeast, 30 Urban Midwest, and Southeast, especially in the summer months, presumably, because of 31 the summer peaking sulfate. However, the mostly negative or weak correlations between 32 PM and O_3 in the summer in the Southwest, Northwest, and southern California could be 33 due to winter-peaking nitrate. Thus, the potential confounding effect of PM on the 34 O₃-mortality relationship could be influenced by the relative contribution of sulfate and 35 nitrate, which varies regionally and seasonally.

	No. of Communities	Winter	Spring	Summer	Fall	Yearly
PM ₁₀						
Industrial Midwest	19	0.37	0.44	0.44	0.39	0.41
Northeast	15	0.34	0.44	0.36	0.44	0.40
Urban Midwest	6	0.24	0.25	0.22	0.26	0.24
Southwest	9	0.00	0.02	-0.02	0.10	0.03
Northwest	11	-0.17	-0.20	-0.13	-0.11	-0.16
Southern California	7	0.19	0.08	0.12	0.19	0.14
Southeast	25	0.33	0.35	0.31	0.31	0.32
U.S.	93	0.23	0.26	0.24	0.26	0.25
PM _{2.5}						
Industrial Midwest	19	0.18	0.39	0.43	0.44	0.36
Northeast	13	0.05	0.26	0.16	0.43	0.25
Urban Midwest	4	0.22	0.31	0.15	0.32	0.20
Southwest	9	-0.15	-0.08	-0.17	-0.15	-0.14
Northwest	11	-0.32	-0.34	-0.39	-0.24	-0.31
Southern California	7	-0.25	-0.22	-0.25	-0.15	-0.23
Southeast	26	0.38	0.47	0.30	0.37	0.39
U.S.	90	0.09	0.21	0.12	0.22	0.16

Table 6-44Correlations between PM and ozone by season and region.

Source: Bell et al. (2007).



Note: The diagonal line indicates 1:1 ratio. Source: Reprinted with permission of Informa UK Ltd, (<u>Smith et al., 2009b</u>).

Figure 6-27 Scatter plots of ozone mortality risk estimates with versus without adjustment for PM₁₀ in NMMAPS cities.

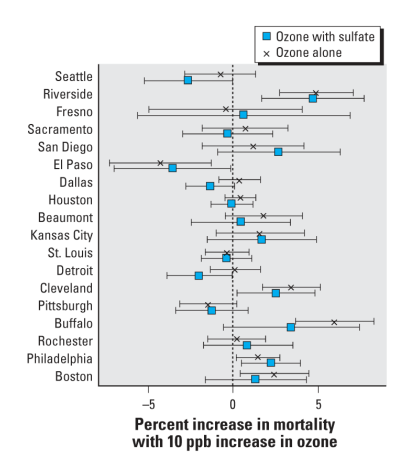
1 2	In an attempt to reassess a number of issues associated with the O_3 -mortality relationship, including confounding, <u>Smith et al. (2009b</u>) re-analyzed the publicly available NMMAPS
3	database for the years 1987-2000. Similar to <u>Bell et al. (2007</u>), the PM ₁₀ data used in the
4	Smith et al. (2009b) analysis consisted primarily of every-6th day data. In analyses
5	conducted to examine the potential confounding effects of PM_{10} , the authors reported
6	that, in most cases, O_3 mortality risk estimates were reduced by between 22% and 33% in
7	copollutant models. This is further highlighted in Figure 6-27, which shows scatter plots
8	of O3-mortality risk estimates with adjustment for PM10 versus without adjustment for
9	PM ₁₀ . Smith et al. (2009b) point out that a larger fraction (89 out of 93) of the posterior
10	estimates lie below the diagonal line (i.e., estimates are smaller with PM ₁₀ adjustment)
11	compared to the raw estimates (56 out of 93). This observation could be attributed to both
12	sets of posterior estimates being calculated by "shrinking towards the mean" along with
13	the small number of days where both PM_{10} and O_3 data was available. However, the most

prominent feature of these plots is that the variation of O_3 -mortality risk estimates across cities is much larger than the impact of PM_{10} adjustment on the O_3 -mortality relationship.

- 3 Franklin and Schwartz (2008) examined the sensitivity of O_3 mortality risk estimates to 4 the inclusion of $PM_{2.5}$ or PM chemical components associated with secondary aerosols (e.g., sulfate $[SO_4^{2-}]$, organic carbon [OC], and nitrate $[NO_3-]$) in copollutant models. 5 6 This analysis consisted of between 3 and 6 years of data from May through September 7 2000-2005 from 18 U.S. communities. The association between O_3 and non-accidental 8 mortality was examined in single-pollutant models and after adjustment for PM_{25} , 9 sulfate, organic carbon, or nitrate concentrations. The single-city effect estimates were 10 combined into an overall estimate using a random-effects model. In the single-pollutant 11 model, the authors found a 0.89% (95% CI: 0.45, 1.33%) increase in nonaccidental 12 mortality with a 10 ppb increase in same-day 24-hour summertime O_3 concentrations 13 across the 18 U.S. communities. Adjustment for $PM_{2.5}$ mass, which was available for 14 84% of the days, decreased the O₃-mortality risk estimate only slightly (from 0.88% to 15 0.79%), but the inclusion of sulfate in the model reduced the risk estimate by 31% (from 16 0.85% to 0.58%). However, sulfate data were only available for 18% of the days. 17 Therefore, a limitation of this study is the limited amount of data for $PM_{2.5}$ chemical 18 components due to the every-3rd-day or every-6th-day sampling schedule. For example, 19 when using a subset of days when organic carbon measurements were available (i.e., 17% 20 of the available days), O_3 mortality risk estimates were reduced to 0.51% (95% CI: -0.36 21 to 1.36) in a single-pollutant model.
- 22 Consistent with the studies previously discussed, the results from Franklin and Schwartz 23 (2008) also demonstrate that the interpretation of the potential confounding effects of 24 copollutants on O₃ mortality risk estimates is not straightforward as a result of the PM 25 sampling schedule employed in most cities. However, Franklin and Schwartz (2008) find 26 that O_3 -mortality risk estimates, although attenuated in some cases (i.e., sulfate), remain 27 positive. As presented in Figure 6-28, the regional and city-to-city variations in O_3 28 mortality risk estimates appear greater than the impact of adjusting for copollutants. In 29 addition, in some cases, a negative O₃ mortality risk estimate becomes even more 30 negative with the inclusion of sulfate (e.g., Seattle) in a copollutant model, or a null O_3 31 mortality risk estimate becomes negative when sulfate is included (e.g., Dallas and 32 Detroit). Thus, the reduction in the overall O_3 mortality risk estimate (i.e., across cities) 33 needs to be assessed in the context of the heterogeneity in the single-city estimates.

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Source: Franklin and Schwartz (2008).

Figure 6-28 Community-specific ozone-mortality risk estimates for nonaccidental mortality per 10 ppb increase in same-day 24-h average summertime ozone concentrations in single-pollutant models and copollutant models with sulfate.

1	In the APHENA study, the investigators from the U.S. (NMMAPS), Canadian, and
2	European (APHEA2) multicity studies collaborated and conducted a joint analysis of
3	PM_{10} and O_3 using each of these datasets (Katsouyanni et al., 2009). For mortality, each
4	dataset consisted of a different number of cities and years of air quality data: U.S.
5	encompassed 90 cities with daily O_3 data from 1987-1996 of which 36 cities had summer
6	only O_3 measurements; Europe included 23 cities with 3-7 years of daily O_3 data during
7	1990-1997; and Canada consisted of 12 cities with daily O_3 data from 1987 to 1996. As
8	discussed in Section 6.2.7.2, the APHENA study conducted extensive sensitivity
9	analyses, of which the 8 df/year results for both the penalized spline (PS) and natural
10	spline (NS) models are presented in the text for comparison purposes, but only the NS

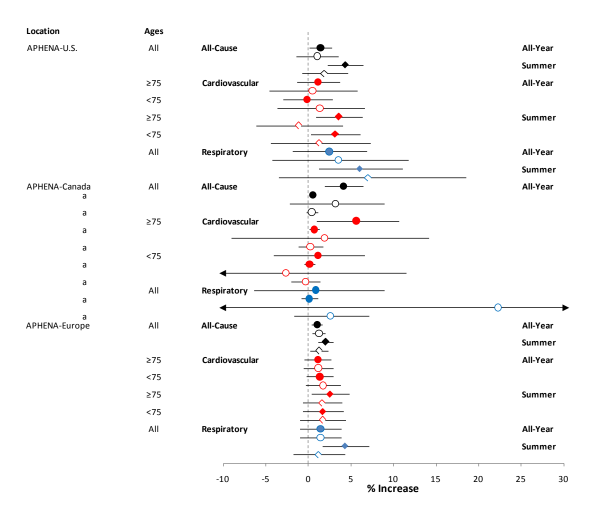
1	results are presented in figures because alternative spline models have previously been
2	shown to result in similar effect estimates (HEI, 2003). Additionally, for the Canadian
3	results, figures contain risk estimates standardized to both a 40 ppb increment for
4	1-h max O ₃ concentrations, consistent with the rest of the ISA, but also the approximate
5	IQR across the Canadian cities as discussed previously (Section 6.2.7.2).
6	In the three detects, the outhous found concrelly positive associations between short term
0 7	In the three datasets, the authors found generally positive associations between short-term
8	O_3 exposure and all-cause, cardiovascular, and respiratory mortality. The estimated
	excess risks for O_3 were larger for the Canadian cities than for the U.S. and European sitis when even intro the net of the conformation of DM and O_1 modulity risk
9	cities. When examining the potential confounding effects of PM_{10} on O_3 mortality risk
10	estimates, the sensitivity of the estimates varied across the data sets and age groups. In
11	the Canadian dataset, O_3 risk estimates were modestly reduced, but remained positive,
12	when adjusting for PM ₁₀ for all-cause mortality for all ages in the PS (4.5% [95% CI: 2.2,
13	6.7%]) and NS (4.2% [95% CI: 1.9, 6.5%]) models to 3.8% (95% CI: -1.4, 9.8%) and
14	3.2% (95% CI: -2.2, 9.0%), respectively, at lag 1 for a 40 ppb increase in 1-h max O_3
15	concentrations (Figure 6-29; Table 6-45). However, adjusting for PM_{10} reduced O_3
16	mortality risk estimates in the \geq 75-year age group, but increased the risk estimates in the
17	<75-year age group. For cardiovascular and respiratory mortality more variable results
18	were observed with O_3 risk estimates being reduced and increased, respectively, in
19	copollutant models with PM_{10} (Figure 6-29; Table 6-45). Unlike the European and U.S.
20	datasets, the Canadian dataset only conducted copollutant analyses at lag 1; as a result, to
21	provide a comparison across study locations only the lag 1 results are presented for the
22	European and U.S. datasets in this section.
23	In the European data, O_3 risk estimates were robust when adjusting for PM ₁₀ in the year-
24	round data for all-cause, cardiovascular and respiratory mortality. When restricting the
25	analysis to the summer months moderate reductions were observed in O_3 risk estimates
26	for all-cause mortality with more pronounced reductions in respiratory mortality. In the
20	U.S. data, adjusting for PM_{10} moderately reduced O ₃ risk estimates for all-cause mortality
28	in a year-round analysis at lag 1 (e.g., both the PS and NS models were reduced from
29	0.18% to $0.13%$) (Figure 6-29; Table 6-45). Similar to the European data, when
30	restricting the analysis to the summer months, in the U.S. O_3 mortality risk estimates
31	were moderately reduced, but remained positive, when adjusting for PM_{10} for all-cause
32	mortality. However, when examining cause-specific mortality risk estimates, consistent
33	with the results from the Canadian dataset, which employed a similar PM sampling
55	with the results from the Canadian dataset, which employed a similar PWI sampling

strategy (i.e., every-6th-day sampling), O_3 risk estimates for cardiovascular and respiratory mortality were more variable (i.e., reduced or increased in all-year and summer analyses). Overall, the estimated O_3 risks appeared to be moderately to substantially sensitive to inclusion of PM₁₀ in copollutant models. Despite the multicity

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Note: Effect estimates are for a 40 ppb increase in 1-h max O_3 concentrations at lag 1. All estimates are for the 8 df/year model with natural splines. Circles represent all-year analysis results while diamonds represent summer season analysis results. Open circles and diamonds represent copollutant models with PM_{10} . Black = all-cause mortality; red = cardiovascular mortality; and blue = respiratory mortality.

^aRisk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O₃ concentrations (see explanation in Section <u>6.2.7.2</u>).

Figure 6-29 Percent increase in all-cause (nonaccidental) and cause-specific mortality from natural spline models with 8 df/yr from the APHENA study for single- and copollutant models.

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Location*	Mortality	Ages	Season	Copollutant	% Increase (95% CI)
APHENA-U.S.	All-Cause	All	All-year		1.42 (0.08, 2.78)
				PM ₁₀	1.02 (-1.40, 3.50)
			Summer		4.31 (2.22, 6.45)
				PM ₁₀	1.90 (-0.78, 4.64)
	Cardiovascular	≥ 75	All-year		1.10 (-1.33, 3.67)
				PM ₁₀	0.47 (-4.61, 5.79)
		<75	All-year		-0.16 (-3.02, 2.86)
				PM ₁₀	1.34 (-3.63, 6.61)
		≥ 75	Summer		3.58 (0.87, 6.37)
				PM ₁₀	-1.17 (-6.18, 4.07)
		<75	Summer		3.18 (0.31, 6.12)
				PM ₁₀	1.26 (-4.46, 7.28)
	Respiratory	All	All-year		2.46 (-1.87, 6.86)
				PM ₁₀	3.50 (-4.23, 11.8)
			Summer		6.04 (1.18, 11.1)
				PM ₁₀	7.03 (-3.48, 18.5)
APHENA-Canada	All-Cause	All	All-year		4.15 (1.90, 6.45)
					0.52 (0.24, 0.80) ^a
				PM ₁₀	3.18 (-2.18, 8.96)
				PM ₁₀	0.40 (-0.28, 1.10) ^a
	Cardiovascular	≥ 75	All-year		5.62 (0.95, 10.7)
					0.70 (0.12, 1.30)a
				PM ₁₀	1.90 (-9.03, 14.1)
				PM ₁₀	0.24 (-1.20, 1.70) ^a
		<75	All-year		1.10 (-4.08, 6.61)
					0.14 (-0.53, 0.82) ^a
				PM ₁₀	-2.64 (-14.7, 11.5)
				PM ₁₀	-0.34 (-2.00, 1.40) ^a
	Respiratory	All	All-year		0.87 (-6.40, 8.96)
					0.11 (-0.84, 1.10) ^a
				PM ₁₀	22.3 (-12.6, 71.3)
				PM ₁₀	2.60 (-1.70, 7.10) ^a

Table 6-45Corresponding effect estimates for Figure 6-29.

Location*	Mortality	Ages	Season	Copollutant	% Increase (95% CI)
APHENA-Europe	All-Cause	All	All-year		1.02 (0.39, 1.66)
				PM ₁₀	1.26 (0.47, 1.98)
			Summer		2.06 (1.10, 2.94)
				PM ₁₀	1.26 (0.16, 2.30)
	Cardiovascular	≥ 75	All-year		1.10 (-0.47, 2.70)
				PM ₁₀	1.18 (-0.55, 2.94)
		<75	All-year		1.34 (-0.24, 2.94)
				PM ₁₀	1.74 (-0.31, 3.75)
		≥ 75	Summer		2.54 (0.39, 4.80)
				PM ₁₀	1.58 (-0.70, 3.99)
		<75	Summer		1.66 (-0.70, 4.15)
				PM ₁₀	1.66 (-1.02, 4.40)
	Respiratory	All	All-year		1.42 (-1.02, 3.83)
				PM ₁₀	1.42 (-1.02, 3.83)
			Summer		4.31 (1.66, 7.11)
				PM ₁₀	1.18 (-1.79, 4.31)

*Effect estimates from Figure 6-29.

^aRisk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O₃ concentrations (see explanation in Section <u>6.2.7.2</u>).

1	Stafoggia et al. (2010) examined the potential confounding effects of PM_{10} on the
2	O ₃ -mortality relationship in individuals 35 years of age and older in 10 Italian cities from
3	2001 to 2005. In a time-stratified case-crossover analysis, using data for the summer
4	months (i.e., April-September), the authors examined O ₃ -mortality associations across
5	each city, and then obtained a pooled estimate through a random-effects meta-analysis.
6	Stafoggia et al. (2010) found a strong association with nonaccidental mortality (9.2%
7	[95% CI: 5.4, 13.0%] for a 30 ppb increase in 8-h max O ₃ concentrations) in an
8	unconstrained distributed lag model (lag 0-5) that persisted in copollutant models with
9	PM_{10} (9.2% [95% CI: 5.4, 13.7%]). Additionally, when examining cause-specific
10	mortality, the authors found positive associations between short-term O ₃ exposure and
11	cardiovascular (14.3% [95% CI: 6.7, 22.4%]), cerebrovascular (8.5% [95% CI: 0.1,
12	16.3%]), and respiratory (17.6% [95% CI: 1.8, 35.6%]) mortality in single-pollutant
13	models. In copollutant models, O_3 -mortality effect estimates for cardiovascular and
14	cerebrovascular mortality were robust to the inclusion of PM_{10} (9.2% [95% CI: 5.4,
15	13.7%]) and 7.3% [95% CI: -1.2, 16.3%], respectively), and attenuated, but remained
16	positive, for respiratory mortality (9.2% [95% CI: -6.9, 28.8%]). Of note, the correlations
17	between O_3 and PM_{10} across cities were found to be generally low, ranging from (-0.03 to
18	0.49). The authors do not specify the sampling strategy used for PM_{10} in this analysis.

Confounding by Seasonal Trend

1	The APHENA study (Katsouyanni et al., 2009), mentioned above, also conducted
2	extensive sensitivity analyses to identify the appropriate: (1) smoothing method and basis
3	functions to estimate smooth functions of time in city-specific models; and (2) degrees of
4	freedom to be used in the smooth functions of time, to adjust for seasonal trends. Because
5	O ₃ peaks in the summer and mortality peaks in the winter, not adjusting or not
6	sufficiently adjusting for the seasonal trend would result in an apparent negative
7	association between the O ₃ and mortality time-series. Katsouyanni et al. (2009) examined
8	the effect of the extent of smoothing for seasonal trends by using models with 3 df/year,
9	8 df/year (the choice for their main model), 12 df/year, and df/year selected using the sum
10	of absolute values of partial autocorrelation function of the model residuals (PACF)
11	(i.e., choosing the degrees of freedom that minimizes positive and negative
12	autocorrelations in the residuals). Table 6-46 presents the results of the degrees of
13	freedom analysis using alternative methods to calculate a combined estimate: the Berkey
14	et al. (1998) meta-regression and the two-level normal independent sampling estimation
15	(TLNISE) hierarchical method. The results show that the methods used to combine
16	single-city estimates did not influence the overall results, and that neither 3 df/year nor
17	choosing the df/year by minimizing the sum of absolute values of PACF of regression
18	residuals was sufficient to adjust for the seasonal negative relationship between O ₃ and
19	mortality. However, it should be noted, the majority of studies in the literature that
20	examined the mortality effects of short-term O ₃ exposure, particularly the multicity
21	studies, used 7 or 8 df/year to adjust for seasonal trends, and in both methods a positive
22	association was observed between O ₃ exposure and mortality.

Table 6-46Sensitivity of ozone risk estimates per 10 µg/m³ increase in
24-h average ozone concentrations at lag 0-1 to alternative
methods for adjustment of seasonal trend, for all-cause mortality
using Berkey MLE and TLNISE Hierarchical Models.

Seasonality Control	Berkey	TLNISE
3 df/year	-0.54 (-0.88, 0.20)	-0.55 (-0.88, -0.22)
8 df/year	0.30 (0.11, 0.50)	0.31 (0.09, 0.52)
12 df/year	0.34 (0.15, 0.53)	0.33 (0.12, 0.54)
PACF	-0.62 (-1.01, -0.22)	-0.62 (-0.98, -0.27)

Source: Reprinted with permission of Health Effects Institute (Katsouyanni et al., 2009).

6.6.2.2 Effect Modification

1	There have been several multicity studies that examined potential effect modifiers, or
2	time-invariant factors, which may modify O_3 mortality risk estimates. These effect
3	modifiers can be categorized into either individual-level or community-level
4	characteristics, which are traditionally examined in second stage regression models. The
5	results from these analyses also inform upon whether certain populations are greater risk
6	of an O_3 -related health effects (Chapter 8). In addition to potentially modifying the
7	association between short-term O_3 exposure and mortality, both individual-level and
8	community-level characteristics may contribute to the geographic pattern of spatial
9	heterogeneity in O_3 mortality risk estimates. As a result, the geographic pattern of O_3
10	mortality risk estimates is also evaluated in this section.

Individual-Level Characteristics

11	Medina-Ramón and Schwartz (2008) conducted a case-only study in 48 U.S. cities to
12	identify populations potentially at increased risk to O3-related mortality for the period
13	1989-2000 (May through September of each year [i.e., warm season]). A case-only
14	design predicts the occurrence of time-invariant characteristics among cases as a function
15	of the exposure level (Armstrong, 2003). For each potential effect modifier
16	(time-invariant individual-level characteristics), city-specific logistic regression models
17	were fitted, and the estimates were pooled across all cities. Furthermore, the authors
18	examined potential differences in individual effect modifiers according to several city
19	characteristics (e.g., mean O ₃ level, mean temperature, households with central air
20	conditioning, and population density) in a meta-regression. Across cities, the authors
21	found a 1.96% (95% CI: 1.14-2.82%) increase in mortality at lag 0-2 for a 30 ppb
22	increase in 8-h max O3 concentrations. Additionally, Medina-Ramón and Schwartz

- 1 (2008) examined a number of individual-level characteristics (e.g., age, race) and chronic 2 conditions (e.g., secondary causes of death) as effect modifiers of the association between 3 short-term O_3 exposure and mortality. The authors found that older adults (i.e., ≥ 65), 4 women >60 years of age, black race, and secondary atrial fibrillation showed the greatest 5 additional percent change in O_3 -related mortality (Table 6-47). When examining city-6 level characteristics, the authors found that older adults, black race, and secondary atrial 7 fibrillation had a larger effect on O_3 mortality risk estimates in cities with lower mean O_3 8 concentrations. Of note, a similar case-only study (Schwartz, 2005b) examined potential 9 effect modifiers of the association between temperature and mortality, which would be 10 expected to find results consistent with the Medina-Ramón and Schwartz (2008) study due to the high correlation between temperature and O₃. However, when stratifying days 11 12 by temperature Schwartz (2005b) found strong evidence that diabetes modified the 13 temperature-mortality association on hot days, which was not as evident when examining 14 the O₃-mortality association in Medina-Ramón and Schwartz (2008). This difference 15 could be due to the study design and populations included in both studies, a multicity 16 study including all ages (Medina-Ramón and Schwartz, 2008) compared to a single-city 17 study of individuals \geq 65 years of age (Schwartz, 2005b). However, when examining 18 results stratified by race, nonwhites were found to have higher mortality risks on both hot 19 and cold days, which provide some support for the additional risk found for black race in 20 Medina-Ramón and Schwartz (2008).
- 21 Individual-level factors that may result in increased risk of O_3 -related mortality were also 22 examined by Stafoggia et al. (2010). As discussed above, using a time-stratified case-23 crossover analysis, the authors found an association between short-term O_3 exposure and 24 nonaccidental mortality in an unconstrained distributed lag model in 10 Italian cities 25 (9.2% [95% CI: 5.4, 13.0%; lag 0-5 for a 30 ppb increase in 8-h max O₃ concentrations). 26 Stafoggia et al. (2010) conducted additional analyses to examine whether age, sex, 27 income level, location of death, and underlying chronic conditions increased the risk of 28 O₃-related mortality, but data were only available for nine of the cities for these analyses. 29 Of the individual-level factors examined, the authors found the strongest evidence for 30 increased risk of O₃-related mortality in individuals \geq 85 years of age (22.4% [95% CI: 31 15.0, 30.2%]), women (13.7% [95% CI: 8.5, 19.7%]), and out-of-hospital deaths (13.0% 32 [95% CI: 6.0, 20.4%]). When focusing specifically on out-of hospital deaths and the 33 subset of individuals with chronic conditions, Stafoggia et al. (2010) found the strongest 34 association for individuals with diabetes, which is consistent with the potentially 35 increased risk of diabetics on hot days observed in Schwartz (2005b).

Table 6-47Additional percent change in ozone-related mortality for individual-
level characteristics.

	Percentage	(95% CI)
Socio-demographic characteristics		
Age 65 yr or older	1.10	0.44, 1.77
Women	0.58	0.18, 0.98
Women <60 yr old ^b	-0.09	-0.76, 0.58
Women ≥ 60 yr old ^b	0.60	0.25, 0.96
Black race	0.53	0.19, 0.87
Low education	-0.29	-0.81, 0.23
Chronic conditions (listed as secondary cause)		
Respiratory system diseases		
Asthma	1.35	-0.31, 3.03
COPD	0.01	-0.49, 0.52
Circulatory system diseases		
Atherosclerosis	-0.72	-1.89, 0.45
Atherosclerotic CVD	0.74	-0.86, 2.37
Atherosclerotic heart disease	-0.38	-1.70, 0.96
Congestive heart disease	-0.04	-0.39, 0.30
Atrial fibrillation	1.66	0.03, 3.32
Stroke	0.17	-0.28, 0.62
Other diseases		
Diabetes	0.19	-0.46, 0.84
Inflammatory diseases	0.18	-1.09, 1.46

^aThese estimates represent the additional percent change in mortality for persons who had the characteristic being examined compared to persons who did not have the characteristic, when the mean O_3 level of the previous 3 days increased 10 ppb. These values were not standardized because they do not represent the actual effect estimate for the characteristic being evaluated, but instead, the difference between effect estimates for persons with versus without the condition.

^bCompared with males in the same age group.

Source: Reprinted with permission of Lippincott Williams & Wilkins (Medina-Ramón and Schwartz, 2008).

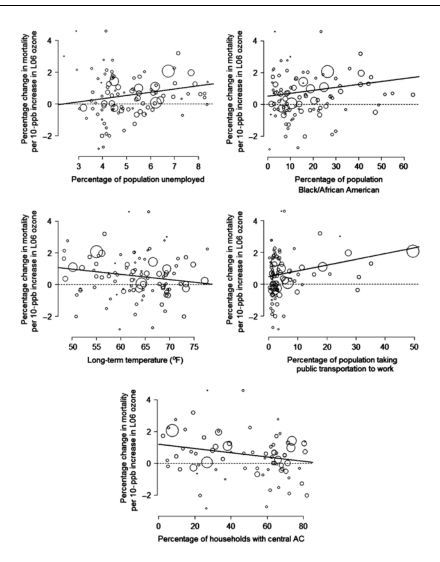
1	Additionally, Cakmak et al. (2011) examined the effect of individual-level characteristics
2	that may modify the O ₃ -mortality relationship in 7 Chilean cities. In a time-series analysis
3	using a constrained distributed lag of 0-6 days, Cakmak et al. (2011) found evidence for
4	larger O_3 mortality effects in individuals >75 years of age compared to younger ages,
5	which is similar to Medina-Ramón and Schwartz (2008) and Stafoggia et al. (2010).
6	Unlike the studies discussed above O3-mortality risk estimates were found to be slightly
7	larger in males (3.71% [95% CI: 0.79, 6.66] for a 40 ppb increase in max 8-h avg O_3
8	concentrations), but were not significantly different than those observed for females
9	(3.00% [95% CI: 0.43, 5.68]). The major focus of <u>Cakmak et al. (2011</u>) is the

- 1examination of the influence of SES indicators (i.e., educational attainment, income level,2and employment status) on the O3-mortality relationship. The authors found the largest3risk estimates in the lowest SES categories for each of the indicators examined this4includes: primary school not completed when examining educational attainment; the5lowest quartile of income level; and unemployed individuals when comparing6employment status.
- 7 Overall, uncertainties exist in the interpretation of the potential effect modifiers identified 8 in Medina-Ramón and Schwartz (2008), Stafoggia et al. (2010), and Cakmak et al. (2011) 9 of the O_3 -mortality relationship due to the heterogeneity in O_3 -mortality risk estimates 10 across cities as highlighted in Smith et al. (2009b) (Figure 6-27) and Franklin and Schwartz (2008) (Figure 6-28). In addition, it is likely that individual-level factors 11 12 identified in Medina-Ramón and Schwartz (2008), (Stafoggia et al., 2010), and Cakmak 13 et al. (2011) only modify the O₃-mortality relationship and do not entirely explain the 14 observed regional heterogeneity in O₃-mortality risk estimates.

Community-level Characteristics

- 15 Several studies also examined city-level (i.e., ecological) variables in an attempt to 16 explain the observed city-to-city variation in estimated O_3 -mortality risk estimates. Bell 17 and Dominici (2008) investigated whether community-level characteristics, such as race, 18 income, education, urbanization, transportation use, PM and O₃ concentrations, number 19 of O_3 monitors, weather, and air conditioning use could explain the heterogeneity in 20 O₃-mortality risk estimates across cities. The authors analyzed 98 U.S. urban 21 communities from NMMAPS for the period 1987-2000. In the all-year regression model 22 that included no community-level variables, a 20 ppb increase in 24-h avg O₃ 23 concentrations during the previous week was associated with a 1.04% (95% CI: 0.56, 24 1.55) increase in mortality. Bell and Dominici (2008) found that higher O₃-mortality 25 effect estimates were associated with an increase in: percent unemployment, fraction of 26 the population Black/African-American, percent of the population that take public 27 transportation to work; and with a reduction in: temperatures and percent of households 28 with central air conditioning (Figure 6-30). The modification of O₃-mortality risk 29 estimates reported for city-specific temperature and prevalence of central air conditioning 30 in this analysis confirm the result from the meta-analyses reviewed in the 2006 O_3 31 AQCD.
- 32The APHENA project (Katsouyanni et al., 2009) examined potential effect modification33of O3 risk estimates in the Canadian, European, and U.S. data sets using a consistent set34of city-specific variables. Table 6-48 presents the results from all age analyses for all-35cause mortality using all-year O3 data for the average of lag 0-1 day. While there are

1	several significant effect modifiers in the U.S. data, the results are mostly inconsistent
2	with the results from the Canadian and European data sets. The positive effect
3	modification by percentage unemployed and the negative effect modification by mean
4	temperature (i.e., a surrogate for air conditioning rate) are consistent with the results
5	reported by Bell and Dominici (2008) discussed above. However, the lack of consistency
6	across the data sets, even between the Canadian and U.S. data, makes it difficult to
7	interpret the results. Some of these associations may be due to coincidental correlations
8	with other unmeasured factors that vary regionally (e.g., mean SO_2 tend to be higher in
9	the eastern U.S.).



Note: The size of each circle corresponds to the inverse of the standard error of the community's maximum likelihood estimate. Risk estimates are for a 10 ppb increase in 24-h avg ozone concentrations during the previous week. Source: Reprinted with permission of Johns Hopkins Bloomberg School of Public Health (Bell and Dominici, 2008).

Figure 6-30 Ozone mortality risk estimates and community-specific characteristics, U.S., 1987-2000.

Table 6-48Percent change in all-cause mortality, for all ages, associated with
a 40ppb increase in 1-h max ozone concentrations at Lag 0–1 at the
25th and 75th percentile of the center-specific distribution of
selected effect modifiers.

	(Canada		I	Europe			U.S.	
Effect Modifier	25th Percentile Estimate (95% CI)	75th Percentile Estimate (95% CI)	t Value	25th Percentile Estimate (95% CI)	75th Percentile Estimate (95% CI)	t Value	25th Percentile Estimate (95% CI)	75th Percentile Estimate (95% CI)	t Value
NO ₂ CV	3.10	3.99	1.33	1.66	1.34	-0.49	1.26	0.08	-2.87
	(1.90, 4.40)	(2.38, 5.62)		(0.71, 2.62)	(-0.08, 2.78)		(0.47, 1.98)	(-0.78, 0.95)	
Mean SO ₂	2.22	4.72	2.16	1.58	1.66	0.16	0.47	1.98	2.79
	(0.71, 3.83)	(2.94, 6.61)		(0.47, 2.62)	(0.39, 2.86)		(-0.47, 1.42)	(1.10, 2.94)	
O ₃ CV	2.86	3.50	0.60	2.62	1.10	-2.65	0.16	1.50	2.68
	(0.79, 5.05)	(2.14, 4.89)		(1.50, 3.75)	(0.24, 1.98)		(-0.70, 1.10)	(0.71, 2.22)	
Mean	3.91	2.54	-1.58	1.74	1.50	-0.43	-0.08	1.26	2.64
NO ₂ /PM ₁₀	(2.54, 5.29)	(0.95, 4.15)		(0.87, 2.70)	(0.47, 2.62)		(-1.02, 0.95)	(0.47, 2.06)	
Mean	2.86	3.50	0.83	1.58	1.58	-0.04	2.14	0.00	-4.40
Temperature	(0.95, 4.72)	(2.22, 4.89)		(0.39, 2.86)	(0.31, 2.78)		(1.34, 2.94)	(-0.78, 0.79)	
% ≥ 75 yr	2.22	4.23	2.68	1.50	1.82	0.52	1.02	1.02	-0.02
-	(0.79, 3.58)	(3.02, 5.54)		(0.55, 2.46)	(0.55, 3.10)		(0.24, 1.90)	(0.31, 1.74)	
Age- standardized Mortality	2.62	4.07	1.14	1.10	1.98	1.07	0.00	1.58	3.81
	(0.79, 4.48)	(2.22, 5.87)		(-0.16, 2.38)	(0.79, 3.26)		(-0.94, 0.87)	(0.87, 2.38)	
%	2.78	3.75	1.88	1.42	1.34	-0.07	0.16	1.50	2.45
Unemployed	(1.42, 4.07)	(2.54, 4.89)		(-0.47, 3.34)	(-0.47, 3.18)		(-0.78, 1.18)	(0.71, 2.30)	

Source: Adapted with permission of Health Effects Institute Katsouyanni et al. (2009).

Regional Pattern of Ozone-Mortality Risk Estimates

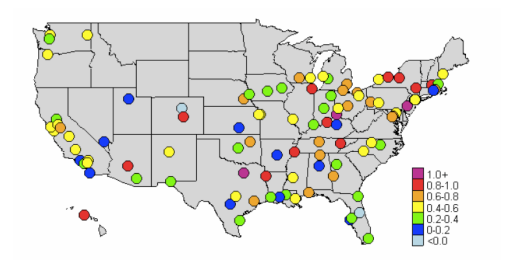
1	In addition to examining whether individual- and community-level factors modify the
2	O3-mortality association, studies have also examined whether these associations varied
3	regionally within the U.S. Bell and Dominici (2008), in the study discussed above, also
4	noted that O ₃ -mortality risk estimates were higher in the Northeast (1.44% [95% CI: 0.78,
5	2.10%]) and Industrial Midwest (0.73% [95% CI: 0.11, 1.35%]), while null associations
6	were observed in the Southwest and Urban Midwest (Table 6-49). The regional
7	heterogeneity in O ₃ -mortality risk estimates was further reflected by Bell and Dominici
8	(2008) in a map of community-specific Bayesian O_3 -mortality risk estimates

1	(Figure 6-31). It is worth noting that in the analysis of PM_{10} using the same data set, Peng
2	et al. (2005) also found that both the Northeast and Industrial Midwest showed
3	particularly elevated effects, especially during the summer months. As mentioned above,
4	although no evidence for confounding of O_3 mortality risk estimates by PM_{10} was
5	observed, <u>Bell et al. (2007</u>) did find regional differences in the correlation between O_3
6	and PM_{10} . Thus, the heterogeneity in O_3 mortality risk estimates may need to be
7	examined as a function of the correlation between PM and O ₃ .
8	Smith et al. (2009b), as discussed earlier, also examined the regional difference in O_3
9	mortality risk estimates across the same seven regions and similarly found evidence for
10	regional heterogeneity. In addition, Smith et al. (2009b) constructed spatial maps of the
11	risk estimates by an extension of a hierarchical model that allows for spatial auto-
12	correlation among the city-specific random effects. Figure 6-32 presents the spatial map
13	of O_3 mortality coefficients from the <u>Smith et al. (2009b</u>) analysis that used 8-h max O_3
14	concentrations during the summer. The results from the Bell and Dominici (2008)
15	analysis (Figure 6-31) shows much stronger apparent heterogeneity in O ₃ -mortality risk
16	estimates across cities than the smoothed map from Smith et al. (2009b) (Figure 6-32),
17	but both maps generally show larger risk estimates in the eastern region of the U.S.

Table 6-49Percentage increase in daily mortality for a 10 ppb increase in
24-h average ozone concentrations during the previous week by
geographic region in the U.S., 1987-2000.

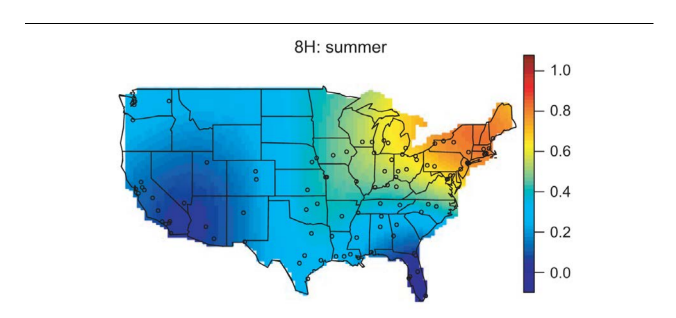
	No. of Communities	Regional Estimate	95% PI*
Regional results			
Industrial Midwest	20	0.73	0.11, 1.35
Northeast	16	1.44	0.78, 2.10
Northwest	12	0.08	-0.92, 1.09
Southern California	7	0.21	-0.46, 0.88
Southeast	26	0.38	-0.07, 0.85
Southwest	9	-0.06	-0.92, 0.81
Urban Midwest	7	-0.05	-1.28, 1.19
National results			
All continental communities	97	0.51	0.27, 076
All communities	98	0.52	0.28, 0.77

Source: Reprinted with permission of Johns Hopkins Bloomberg School of Public Health (Bell and Dominici, 2008).



Source: Reprinted with permission of Johns Hopkins Bloomberg School of Public Health, (Bell and Dominici, 2008).

Figure 6-31 Community-specific Bayesian ozone-mortality risk estimates in 98 U.S. communities.



Source: Reprinted with permission of Informa UK Ltd. (Smith et al., 2009b).

Figure 6-32 Map of spatially dependent ozone-mortality coefficients for 8-h max ozone concentrations using summer data.

6.6.2.3 Interaction

1	Interactions can lead to either antagonistic or synergistic effects; however, most studies
2	attempt to identify potential factors that interact synergistically with O ₃ to increase the
3	risk of mortality. Within this section, interactive effects are defined as time-varying
4	covariates, such as temperature and copollutants that are included in 1st stage time-series
5	regression models. To date, only a few time-series studies have investigated the potential
6	interaction between O ₃ exposure and copollutants or weather variables. This can be
7	attributed to the moderate to high correlation between O ₃ and these covariates, which
8	makes such investigations methodologically challenging.
9	<u>Ren et al. (2008)</u> examined the possible synergistic effect between O_3 and temperature on
10	mortality in the 60 largest eastern U.S. communities from the NMMAPS data during the
11	warm months (i.e., April to October) from 1987-2000. This analysis was restricted to the
12	eastern areas of the U.S. (i.e., Northeast, Industrial Midwest and Southeast) because a
13	previous study which focused specifically on the eastern U.S. found that
14	temperature-mortality patterns differ between the northeast and southeast regions
15	possibly due to climatic differences (Curriero et al., 2002). To examine possible
16	geographic differences in the interaction between temperature and O ₃ , <u>Ren et al. (2008</u>)
17	further divided the NMMAPS regions into the Northeast, which included the Northeast
18	and Industrial Midwest regions (34 cities), and the Southeast, which included the
19	Southeast region (26 cities). The potential synergistic effects between O ₃ and temperature
20	were examined using two different models. Model 1 included an interaction term in a
21	Generalized Additive Model (GAM) for O3 and maximum temperature (3-day avg values
22	were used for both terms) to examine the bivariate response surface and the pattern of
23	interaction between the two variables in each community. Model 2 consisted of a
24	Generalized Linear Model (GLM) that used interaction terms to stratify by "low,"
25	"moderate," and "high" temperature days using the first and third quartiles of temperature
26	as cut-offs to examine the percent increase in mortality in each community. Furthermore,
27	a two-stage Bayesian hierarchical model was used to estimate the overall percent increase
28	in all-cause mortality associated with short-term O3 exposure across temperature levels
29	and each region using model 2. The same covariates were used in both model 1 and 2.
30	The bivariate response surfaces from model 1 suggest possible interactive effects
31	between O ₃ and temperature although the interpretation of these results is not
32	straightforward due to the high correlation between these terms. The apparent interaction
33	between temperature and O ₃ as evaluated in model 2 varied across geographic regions. In
34	the northeast region, a 20 ppb increase in 24-h avg O3 concentrations at lag 0-2 was
35	associated with an increase of 4.49% (95% posterior interval [PI]: 2.39, 6.36%), 6.21%
36	(95% PI: 4.47, 7.66%) and 12.8% (95% PI: 9.77, 15.7%) in mortality at low, moderate
37	and high temperature levels, respectively. The corresponding percent increases in

mortality in the southeast region were 2.27% (95% PI: -2.23, 6.46%) for low temperature, 3.02% (95% PI: 0.44, 5.70%) for moderate temperature, and 2.60% (95% PI: -0.66, 6.01%) for high temperature.

4 When examining the relationship between temperature and O₃-related mortality, the 5 results reported by <u>Ren et al. (2008</u>) (i.e., higher O_3 -mortality risks on days with higher 6 temperatures) may appear to contradict the results of Bell and Dominici (2008) described 7 earlier (i.e., communities with higher temperature have lower O₃-mortality risk 8 estimates). However, the observed difference in results can be attributed to the 9 interpretation of effect modification in a second-stage regression which uses long-term 10 average temperatures, as was performed by Bell and Dominici (2008), compared to a 11 first-stage regression that examines the interaction between daily temperature and O₃-12 related mortality. In this case, the second-stage regression results from Bell and Dominici 13 (2008) indicate that a city with lower temperatures, on average, tend to show a stronger 14 O_3 mortality effect, whereas, in the first-stage regression performed by Ren et al. (2008), 15 the days with higher temperature tend to show a larger O_3 -mortality effect. This observed 16 difference may in part reflect the higher air conditioning use in communities with higher 17 long-term average temperatures. Therefore, the findings from Ren et al. (2008) indicating 18 generally lower O_3 risk estimates in the southeast region where the average temperature is 19 higher than in the northeast region is consistent with the regional results reported by Bell 20 and Dominici (2008). As demonstrated by the results from both Ren et al. (2008) and 21 Bell and Dominici (2008) caution is required when interpreting results from studies that 22 examined interactive effects using two different approaches because potential effect 23 modification as suggested in a second-stage regression generally does not provide 24 evidence for a short-term interaction examined in a first-stage regression. Overall, further 25 examination of the potential interactive (synergistic) effects of O₃ and covariates in time-26 series regression models is required to more clearly understand the factors that may 27 influence O₃ mortality risk estimates.

6.6.2.4 Evaluation of the Ozone-Mortality C-R Relationship and Related Issues

Evaluation of the O₃-mortality C-R relationship is not straightforward because the evidence from multicity studies (using log-linear models) suggests that O₃-mortality associations are highly heterogeneous across regions. In addition, there are numerous issues that may influence the shape of the O₃-mortality C-R relationship and the observed association between short-term O₃ exposure and mortality that warrant examination including: multi-day effects (distributed lags), mortality displacement (i.e., hastening of death by a short period), potential adaptation, and the exposure metric used to compute

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risks (e.g., 1-hour daily max versus 24-h avg). The following section presents the recent studies identified that conducted an initial examination of these issues.

Multiday Effects, Mortality Displacement, and Adaptation

3 The pattern of positive lagged associations followed by negative associations in a 4 distributed lag model may be considered an indication of "mortality displacement" 5 (i.e., deaths are occurring in frail individuals and exposure is only moving the day of 6 death to a day slightly earlier). Zanobetti and Schwartz (2008b) examined this issue in 48 7 U.S. cities during the warm season (i.e., June-August) for the years 1989-2000. In an 8 initial analysis, the authors applied a GLM to examine same-day O₃-mortality effects, and 9 in the model included an unconstrained distributed lag for apparent temperature to take 10 into account the effect of temperature on the day death occurred and the previous 7 days. 11 To examine mortality displacement Zanobetti and Schwartz (2008b) refit models using 12 two approaches: an unconstrained and a smooth distributed lag each with 21-day lags for 13 O₃. In this study, all-cause mortality as well as cause-specific mortality 14 (i.e., cardiovascular, respiratory, and stroke) were examined for evidence of mortality 15 displacement. The authors found a 0.96% (95% CI: 0.60, 1.30%) increase in all-cause 16 mortality across all 48 cities for a 30 ppb increase in 8-h max O_3 concentrations at lag 0 17 whereas the combined estimate of the unconstrained distributed lag model (lag 0-20) was 18 1.54% (95% CI: 0.15, 2.91%). Similarly, when examining the cause-specific mortality 19 results (Table 6-50), larger risk estimates were observed for the distributed lag model 20 compared to the lag 0 day estimates. However, for stroke a slightly larger effect was 21 observed at lags 4-20 compared to lags 0-3 suggesting a larger window for O₃-induced 22 stroke mortality. This is further supported by the sum of lags 0 through 20 days showing 23 the greatest effect. Overall, these results suggest that estimating the mortality risk using a 24 single day of O_3 exposure may underestimate the public health impact, but the extent of 25 multi-day effects appear to be limited to a few days. This is further supported by the 26 shape of the combined smooth distributed lag (Figure 6-33). It should be noted that the 27 proportion of total variation in the effect estimates due to the between-cities heterogeneity, as measured by I^2 statistic, was relatively low (4% for the lag 0 estimates 28 29 and 21% for the distributed lag), but 21 out of the 48 cities exhibited null or negative 30 estimates. As a result, the estimated shape of the distributed lag cannot be interpreted as a 31 general form of lag structure of associations applicable to all the cities included in this 32 analysis.

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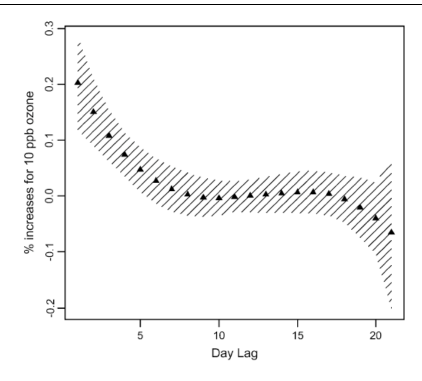
Table 6-50	Estimated effect of a 10 ppb increase in 8-h max ozone
	concentrations on mortality during the summer months for
	single-day and distributed lag models.

	% (Percentage)	95% CI
Total mortality		
Lag 0	0.32	0.20, 0.43
Sum lags 0-20	0.51	0.05, 0.96
Sum lags 0-3	0.53	0.28, 0.77
Sum lags 4-20	-0.02	-0.35, 0.31
Cardiovascular mortality		
Lag 0	0.47	0.30, 0.64
Sum lags 0-20	0.49	-0.01, 1.00
Sum lags 0-3	0.80	0.48, 1.13
Sum lags 4-20	-0.23	-0.67, 0.22
Respiratory mortality		
Lag 0	0.54	0.26, 0.81
Sum lags 0-20	0.61	-0.41, 1.65
Sum lags 0-3	0.83	0.38, 1.28
Sum lags 4-20	-0.24	-1.08, 0.60
Stroke		
Lag 0	0.37	0.01, 0.74
Sum lags 0-20	2.20	0.76, 3.67
Sum lags 0-3	0.92	0.26, 1.59
Sum lags 4-20	1.26	0.05, 2.49

Source: Reprinted with permission of American Thoracic Society, Zanobetti and Schwartz (2008b).

1	Samoli et al. (2009) also investigated the temporal pattern of mortality effects in response
2	to short-term exposure to O_3 in 21 European cities that were included in the APHEA2
3	project. Using a method similar to Zanobetti and Schwartz (2008b), the authors applied
4	unconstrained distributed lag models with lags up to 21 days in each city during the
5	summer months (i.e., June through August) to examine the effect of O ₃ on all-cause,
6	cardiovascular, and respiratory mortality. They also applied a generalized additive
7	distributed lag model to obtain smoothed distributed lag coefficients. However, unlike
8	Zanobetti and Schwartz (2008b), Samoli et al. (2009) controlled for temperature using a
9	linear term for humidity and an unconstrained distributed lag model of temperature at
10	lags 0-3 days. The choice of 0- through 3-day lags of temperature was based on a
11	previous European multicity study (Baccini et al., 2008), which suggested that summer
12	temperature effects last only a few days. Upon combining the individual city estimates
13	across cities in a second stage regression, Samoli et al. (2009) found that the estimated

1 effects on respiratory mortality were extended for a period of two weeks. However, for 2 all-cause and cardiovascular mortality, the 21-day distributed lag models yielded null or 3 (non-significant) negative estimates (Table 6-51). Figure 6-34 shows the distributed lag 4 coefficients for all-cause mortality, which exhibit a declining trend and negative 5 coefficients beyond 5-day lags. The authors' interpretation of these results was that 6 "using single-day exposures may have overestimated the effects on all-cause and 7 cardiovascular mortality, but underestimated the effects on respiratory mortality." Thus, 8 the results in part suggest evidence of mortality displacement for all-cause and 9 cardiovascular mortality.



Source: Reprinted with permission of American Thoracic Society (<u>Zanobetti and Schwartz, 2008b</u>). Note: The triangles represent the percent increase in all-cause mortality for a 10 ppb increase in 8-h max O_3 concentrations at each lag while the shaded areas are the 95% point-wise confidence intervals.

Figure 6-33 Estimated combined smooth distributed lag for 48 U.S. cities during the summer months.

Table 6-51Estimated percent increase in cause-specific mortality (and
95% CIs) for a 10-µg/m³ increase in maximum 8-hour ozone during
June-August.

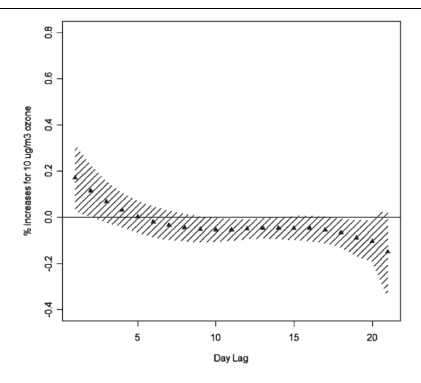
	Fixed effects % (95% CI)	Random effects % (95% CI)
Total mortality ^a		
Lag 0	0.28 (0.11, 0.45)	0.28 (0.07, 0.48)
Average lags 0-1	0.24 (0.15, 0.34)	0.22 (0.08, 0.35)
Sum lags 0-20, unconstrained	0.01 (-0.40, 0.41)	-0.54 (-1.28, 0.20)
Sum lags 0-20, penalized	0.01 (-0.41, 0.42)	-0.56 (-1.30, 0.19)
Cardiovascular mortality ^a		
Lag 0	0.43 (0.18, 0.69)	0.37 (0.05, 0.69)
Average lags 0-1	0.33 (0.19, 0.48)	0.25 (0.03, 0.47)
Sum lags 0-20, unconstrained	-0.33 (-0.93, 0.29)	-0.62 (-1.47, 0.24)
Sum lags 0-20, penalized	-0.32 (-0.92, 0.28)	-0.57 (-1.39, 0.26)
Respiratory mortality ^a		
Lag 0	0.36 (-0.21, 0.94)	0.36 (-0.21, 0.94)
Average lags 0-1	0.40 (0.11, 0.70)	0.40 (0.11, 0.70)
Sum lags 0-20, unconstrained	3.35 (1.90, 4.83)	3.35 (1.90, 4.83)
Sum lags 0-20, penalized	3.66 (2.25, 5.08)	3.66 (2.25, 5.08)

^aAnalysis for the same day (lag 0), the average of the same and previous day (lag 0-1), the unconstrained distributed lag model for the sum of 0-20 days and the penalized distributed lag model (lag 0-20)

Source: Used with permission of BMJ Group (Samoli et al., 2009).

1	Although the APHENA project (Katsouyanni et al., 2009) did not specifically investigate
2	mortality displacement and therefore did not consider longer lags (e.g., lag >3 days), the
3	study did present O_3 risk estimates for lag 0-1, lag 1, and a distributed lag model of 0-
4	2 days in the Canadian, European, and U.S. datasets. <u>Katsouyanni et al. (2009</u>) found that
5	the results vary somewhat across the regions, but, in general, there was no indication that
6	the distributed lag model with up to a 2-day lag yielded meaningfully larger O ₃ mortality
7	risk estimates than the lag 0-1 and lag 1 results. For example, for all-cause mortality,
8	using the model with natural splines and 8 df/year to adjust for seasonal trends, the
9	reported percent excess risk for mortality for a 40 ppb increase in 1-h max O_3
10	concentrations for lag 0-1, lag 1, and the distributed lag model (lag 0-2) was 2.70%
11	(95% CI: 1.02, 4.40%), 1.42% (95% CI: 0.08, 2.78%), and 3.02% (95% CI: 1.10, 4.89%),
12	respectively. Thus, the observed associations appear to occur over a short time period,
13	(i.e., a few days). Similarly, the Public Health and Air Pollution in Asia (PAPA) study
14	(Wong et al., 2010) also examined multiple lag days (i.e., lag 0, lag 0-1, and lag 0-4), and
15	although it did not specifically examine mortality displacement it does provide additional
16	evidence regarding the timing of mortality effects proceeding O_3 exposure. In a combined

1	and the interview of the former of the second state of the second
1 2	analysis using data from all four cities examined (Bangkok, Hong Kong, Shanghai, and Wuhan), average risk estimates at leg 0.4 were larger than these at leg 0 or leg 0.1 in both
2 3	Wuhan), excess risk estimates at lag 0-4 were larger than those at lag 0 or lag 0-1 in both
	fixed and random effect models (results not presented quantitatively). The larger risk
4	estimates at lag 0-4 can primarily be attributed to the strong associations observed in
5	Bangkok and Shanghai. However, it is worth noting that Bangkok differs from the three
6	Chinese cities included in this analysis in that it has a tropical climate and does not
7	exhibit seasonal patterns of mortality. As a result, <u>Wong et al. (2010)</u> examined the O_3 -
8	mortality associations at lag 0-1 in only the three Chinese cities and found that risk
9	estimates were slightly reduced from 2.26% (95% CI: 1.36, 3.16) in the 4 city analysis to
10	1.84% (0.77, 2.86) in the 3 city analysis for a 30 ppb increase in 8-h max O_3
11	concentrations. Overall, the PAPA study further supports the observation of the
12	APHENA study that associations between O ₃ and mortality occur over a relatively short-
13	time period, but also indicates that it may be difficult to interpret O ₃ -mortality
14	associations across cities with different climates and mortality patterns.
15	When comparing the studies that explicitly examined the potential for mortality
16	displacement in the O_3 -mortality relationship, the results from <u>Samoli et al. (2009</u>), which
17	provide evidence that suggests mortality displacement, are not consistent with those
18	reported by Zanobetti and Schwartz (2008b). However, the shapes of the estimated
19	smooth distributed lag associations are similar (Figure 6-33 and Figure 6-34). A closer
20	examination of these figures shows that in the European data beyond a lag of 5 days the
21	estimates remain negative whereas in the U.S. data the results remain near zero for the
22	corresponding lags. These observed difference could be due to the differences in the
23	model specification between the two studies, specifically the use of: an unconstrained
24	distributed lag model for apparent temperature up to 7 previous days (Zanobetti and
25	Schwartz, 2008b) versus a linear term for humidity and an unconstrained distributed lag
26	model of temperature up to 3 previous days (<u>Samoli et al., 2009</u>); and natural cubic
27	splines with 2 df per season (Zanobetti and Schwartz, 2008b) versus dummy variables per
28	month per year to adjust for season (<u>Samoli et al., 2009</u>). It is important to note that these
29	differences in model specification may have also influenced the city-to-city variation in
30	risk estimates observed in these two studies (i.e., homogenous estimates across cities in
31	Zanobetti and Schwartz (2008b) and heterogeneous estimates across cities in Samoli et
32	al. (2009). Overall, the evidence of mortality displacement remains unclear, but <u>Samoli et</u>
33	al. (2009), Zanobetti and Schwartz (2008b), and Katsouyanni et al. (2009) all suggest that
33 34	
	the positive associations between O_3 and mortality are observed mainly in the first
35	few days after exposure.



Note: The triangles represent the percent increase in all-cause mortality for a 10 μ g/m³ increase in 8-h max O₃ concentrations at each lag; the shaded area represents the 95% CIs. Source: Reprinted with permission of BMJ Group (<u>Samoli et al., 2009</u>).

Figure 6-34 Estimated combined smooth distributed lag in 21 European cities during the summer (June-August) months.

Adaptation

1	Controlled human exposure studies have demonstrated an adaptive response to O_3
2	exposure for respiratory effects, such as lung function decrements, but this issue has not
3	been examined in the epidemiologic investigation of mortality effects of O ₃ . Zanobetti
4	and Schwartz (2008a) examined if there was evidence of an adaptive response in the
5	O ₃ -mortality relationship in 48 U.S. cities from 1989 to 2000 (i.e., the same data analyzed
6	in Zanobetti and Schwartz (2008b). The authors examined all-cause mortality using a
7	case-crossover design to estimate the same-day (i.e., lag 0) effect of O ₃ , matched on
8	referent days from every-3rd-day in the same month and year as the case. Zanobetti and
9	Schwartz (2008a) examined O ₃ -mortality associations by: season, month in the summer
10	season (i.e., May through September), and age categories in the summer season
11	(<u>Table 6-52</u>). The estimated O_3 mortality risk estimate at lag 0 was found to be highest in
12	the summer (1.51% [95% CI: 1.14, 1.87%]; lag 0 for a 30 ppb increase in 8-h max O_3
13	concentrations), and, within the warm months, the association was highest in July (1.96%

1	[95% CI: 1.42, 2.48%]; lag 0). ¹ Upon further examination of the summer months, the
2	authors also observed diminished effects in August (0.84% [95% CI: 0.33, 1.39%]; lag
3	0). Based on these results, the authors concluded that the mortality effects of O_3 appear
4	diminished later in the O_3 season.

Table 6-52Percent excess all-cause mortality per 10 ppb increase in daily
8-h max ozone on the same day, by season, month, and age
groups.

	%	95% CI
By Season		
Winter	-0.13	-0.56, 0.29
Spring	0.35	0.16, 0.54
Summer	0.50	0.38, 0.62
Fall	0.05	-0.14, 0.24
By Month		
Мау	0.48	0.28, 0.68
June	0.46	0.24, 0.68
July	0.65	0.47, 0.82
August	0.28	0.11, 0.46
September	-0.09	-0.35, 0.16
By Age Group		
0-20	0.08	-0.42, 0.57
21-30	0.10	-0.67, 0.87
31-40	0.07	-0.38, 0.52
41-50	0.08	-0.27, 0.43
51-60	0.54	0.19, 0.89
61-70	0.38	0.16, 0.61
71-80	0.50	0.32, 0.67
80	0.29	0.13, 0.44

Source: Zanobetti and Schwartz (2008a).

To further evaluate the potential adaptive response observed in Zanobetti and Schwartz (2008a) the distribution of the O_3 concentrations across the 48 U.S. cities during July and August was examined. Both July and August were found to have comparable means of 48.6 and 47.9 ppb with a reported maximum value of 97.9 and 96.0 ppb, respectively. Thus, the observed reduction in O_3 -related mortality effect estimates in August (0.84%)

¹ These values have been standardized to the increment used throughout the ISA for max 8-h avg increase in O_3 concentrations of 30 ppb. These values differ from those presented in Table 6-52 from <u>Zanobetti and Schwartz (2008a</u>) because the authors presented values for a 10 ppb increase in max 8-h avg O_3 concentrations.

1	compared to July (1.96%) appears to support the existence of an adaptive response.
2	However, unlike an individual's adaptive response to decrements in lung function from
3	short-term O3 exposure, an examination of mortality prevents a direct observation of
4	adaptation. Rather, for mortality the adaptation hypothesis is tested with a tacit
5	assumption that, whatever the mechanism for O_3 -induced mortality, the risk of death
6	from short-term O_3 exposure is reduced over the course of the summer months through
7	repeated exposures. This idea would translate to a smaller population that would die from
8	O_3 exposure towards the end of summer. This may complicate the interpretation of the
9	distributed lag coefficients with long lag periods because the decreased coefficients may
10	reflect diminished effects of the late summer, rather than diminished effects that are
11	constant across the summer. These intertwined issues need to be investigated together in
12	future research.

Exposure Metric

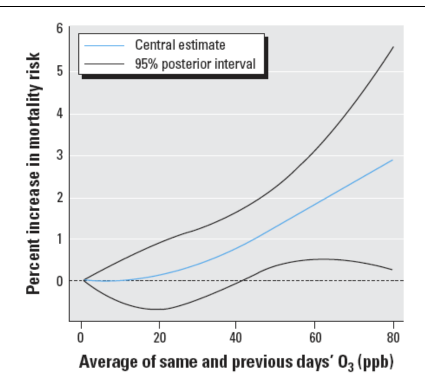
13	When examining the association between short-term O_3 exposure and mortality it is also
14	important to consider the exposure metric used (i.e., 24-h avg, 8-h max, and 1-h max). To
15	date, only a few studies have conducted analyses to examine the impact of different
16	exposure metrics on O_3 mortality risk estimates. In <u>Smith et al. (2009b</u>), the authors
17	examined the effect of different exposure metrics (i.e., 24-h avg , 8-h max, and 1-h max)
18	on O_3 -mortality regression coefficients. When examining whether there are differences in
19	city-specific risk estimates when using different exposure metrics, Smith et al. (2009b)
20	found a rather high correlation ($r = 0.7-0.8$) between risk estimates calculated using
21	24-h avg versus 8-h max and 1-h max versus 8-h max averaging times. These results are
22	consistent with the correlations reported by <u>Darrow et al. (2011a</u>) (Section $6.2.7.3$)
23	between the 8-h max and 24- avg exposure metrics.

24In addition to these recent studies published since the 2006 O3 AQCD, Gryparis et al.25(2004) also supports the high correlation between 1-h max and 8-h max O326concentrations reported in Smith et al. (2009b) and Darrow et al. (2011a) and the27subsequent high degree of similarity between mortality risk estimates calculated using28either metric. Although only a limited number of studies have examined the effect of29different exposure metrics on O3-mortality risk estimates, these studies suggest relatively30comparable results across the exposure metrics used.

Ozone-Mortality C-R Relationship and Threshold Analyses

31	Several of the recent studies evaluated have applied a variety of statistical approaches to
32	examine the shape of the O_3 -mortality C-R relationship and whether a threshold exists.
33	The approach used by <u>Bell et al. (2006</u>) consisted of applying four statistical models to

1	the NMMAPS data, which included 98 U.S. communities for the period 1987-2000.
2	These models included: a linear analysis (i.e., any change in O ₃ concentration can be
3	associated with mortality) (Model 1); a subset analysis (i.e., examining O ₃ -mortality
4	relationship below a specific 24- avg concentration, ranging from 5 to 60 ppb) (Model 2);
5	a threshold analysis (i.e., assuming that an association between O ₃ and mortality is
6	observed above a specific concentration and not below it, using the threshold values set at
7	an increment of 5 ppb between 0 to 60 ppb and evaluating a presence of a local minima in
8	AICs computed at each increment) (Model 3); and nonlinear models using natural cubic
9	splines with boundary knots placed at 0 and 80 ppb, and interior knots placed at 20 and
10	40 ppb (Model 4). A two-stage Bayesian hierarchical model was used to examine these
11	models and O ₃ -mortality risk estimates at the city-level in the first stage analysis and
12	aggregate estimates across cities in the 2nd stage analysis using the average of 0- and
13	1-day lagged 24-h avg O_3 concentrations. The results from all of these models suggest
14	that if a threshold exists it does so well below the current O ₃ NAAQS. When restricting
15	the analysis to all days when the 1997 O_3 NAAQS 8-hour standard (i.e., 84 ppb daily
16	8-h max) is met in each community, Bell et al. (2006) found there was still a 0.60% (95%
17	PI: 0.30, 0.90%) increase in mortality per 20 ppb increase in 24-h avg O ₃ concentrations
18	at lag 0-1. Figure 6-35 shows the combined C-R curve obtained using the nonlinear
19	model (Model 4). Although these results suggest the lack of threshold in the O3-mortality
20	relationship, it is difficult to interpret such a curve because: (1) there is uncertainty
21	around the shape of the C-R curve at 24-h avg O ₃ concentrations generally below 20 ppb,
22	and (2) the C-R curve does not take into consideration the heterogeneity in O ₃ -mortality
23	risk estimates across cities.



Source: Bell et al. (2006)

Figure 6-35 Estimated combined C-R curve for nonaccidental mortality and 24-hour average ozone concentrations at lag 0-1 using the nonlinear (spline) model.

1	Using the same NMMAPS dataset as Bell et al. (2006), Smith et al. (2009b) further
2	examined the O ₃ -mortality C-R relationship. Similar to Bell et al. (2006), Smith et al.
3	(2009b) conduct a subset analysis, but instead of restricting the analysis to days with O_3
4	concentrations below a cutoff the authors only include days above a defined cutoff in the
5	analysis. The results of this "reversed subset" approach are in line with those reported by
6	Bell et al. (2006); consistent positive associations at all cutoff points up to a defined
7	concentration where the total number of days with 24-h avg O_3 concentrations above a
8	value are so limited that the variability around the central estimate is increased. In the
9	Smith et al. (2009b) analysis this observation was initially observed at 45 ppb, with the
10	largest variability at 60 ppb; however, unlike Bell et al. (2006) where 73% of days are
11	excluded when subsetting the data to less than 20 ppb, the authors do not detail the
12	number of days of data included in the subset analyses at higher concentrations. In
13	addition to the subset analysis, Smith et al. (2009b) examined the shape of the C-R curve
14	using a piecewise linear approach with cutpoints at 8-h avg concentrations of 40 ppb,
15	60 ppb, and 80 ppb. Smith et al. (2009b) found that the shape of the C-R curve is similar

1to that reported by Bell et al. (2006) (Figure 6-35), but argue that slopes of the β for each2piece of the curve are highly variable with the largest variation in the 60-80 ppb range.3However, the larger variability around the β between 60-80 would be expected due to the4small number of days with O_3 concentrations within that range in an all-year analysis.5This result is consistent with that observed by Bell et al. (2006), which is presented in6Figure 6-35.

7The APHENA project (Katsouyanni et al., 2009) also analyzed the Canadian and8European datasets (the U.S. data were analyzed for PM10 only) for evidence of a9threshold, using the threshold analysis method (Model 3) applied in Bell et al. (2006)10study described above. There was no evidence of a threshold in the Canadian data11(i.e., the pattern of AIC values for each increment of a potential threshold value varied12across cities, most of which showed no local minima). Likewise, the threshold analysis13conducted using the European data also showed no evidence of a threshold.

14 The PAPA study, did not examine whether a threshold exists in the O₃-mortality C-R 15 relationship, but instead the shape of the C-R curve individually for each city (Bangkok, 16 Hong Kong, Shanghai, and Wuhan) (Wong et al., 2010). Using a natural spline smoother 17 with 3df for the O_3 term, Wong et al. (2010) examined whether non-linearity was present 18 by testing the change in deviance between the smoothed, non-linear, model and an 19 unsmoothed, linear, model with 1 df. For each of the cities, both across the full range of 20 the O_3 distribution and specifically within the range of the 25th to 75th percentile of each 21 city's O_3 24-h avg concentrations (i.e., a range of 9.7 ppb to 60.4 ppb across the cities) 22 there was no evidence of a non-linear relationship in the O₃-mortality C-R curve. It 23 should be noted that the range of the 25th to 75th percentiles in all of the cities, except 24 Wuhan, was lower than that observed in the U.S. using all-year data where the range 25 from the 25th to 75th percentiles is 30 ppb to 50 ppb (Table 3-6).

26 Additional threshold analyses were conducted using NMMAPS data, by Xia and Tong 27 (2006) and Stylianou and Nicolich (2009). Both studies used a new statistical approach developed by Xia and Tong (2006) to examine thresholds in the O₃ mortality C-R 28 29 relationship. The approach consisted of an extended GAM model, which accounted for 30 the cumulative and nonlinear effects of air pollution using a weighted cumulative sum for 31 each pollutant, with the weights (non-increasing further into the past) derived by a 32 restricted minimization method. The authors did not use the term distributed lag model, 33 but their model has the form of distributed lag model, except that it allows for nonlinear 34 functional forms. Using NMMAPS data for 1987-1994 for 3 U.S. cities (Chicago, 35 Pittsburgh, and El Paso), Xia and Tong (2006) found that the extent of cumulative effects 36 of O_3 on mortality were relatively short. While the authors also note that there was 37 evidence of a threshold effect around 24-h avg concentrations of 25 ppb, the threshold

- values estimated in the analysis were sometimes in the range where data density was low. Thus, this threshold analysis needs to be replicated in a larger number of cities to confirm this observation. It should be noted that the model used in this analysis did not include a smooth function of days to adjust for unmeasured temporal confounders, and instead adjusted for season using a temperature term. As a result, these results need to be viewed with caution because some potential temporal confounders (e.g., influenza) do not always follow seasonal patterns of temperature.
- 8 Stylianou and Nicolich (2009) examined the existence of thresholds following an 9 approach similar to Xia and Tong (2006) for all-cause, cardiovascular, and respiratory 10 mortality using data from NMMAPS for nine major U.S. cities (i.e., Baltimore, Chicago, 11 Dallas/Fort Worth, Los Angeles, Miami, New York, Philadelphia, Pittsburgh, and 12 Seattle) for the years 1987-2000. The authors found that PM_{10} and O_3 were the two 13 important predictors of mortality. Stylianou and Nicolich (2009) found that the estimated 14 O₃-mortality risks varied across the nine cities with the models exhibiting apparent 15 thresholds, in the 10-45 ppb range for O_3 (3-day accumulation). However, given the city-16 to-city variation in risk estimates, combining the city-specific estimates into an overall 17 estimate complicates the interpretation of a threshold. Unlike the Xia and Tong (2006) analysis, Stylianou and Nicolich (2009) included a smooth function of time to adjust for 18 19 seasonal/temporal confounding, which could explain the difference in results between the 20 two studies.
- 21 In conclusion, the evaluation of the O₃-mortality C-R relationship did not find any 22 evidence that supports a threshold in the relationship between short-term exposure to O_3 23 and mortality within the range of O₃ concentrations observed in the U.S. Additionally, 24 recent evidence suggests that the shape of the O_3 -mortality C-R curve remains linear 25 across the full range of O_3 concentrations. However, the studies evaluated demonstrated 26 that the heterogeneity in the O_3 -mortality relationship across cities (or regions) 27 complicates the interpretation of a combined C-R curve and threshold analysis. Given the 28 effect modifiers identified in the mortality analyses that are also expected to vary 29 regionally (e.g., temperature, air conditioning prevalence), a national or combined 30 analysis may not be appropriate to identify whether a threshold exists in the O₃-mortality 31 C-R relationship. Overall, the studies evaluated support a linear O₃-mortality C-R 32 relationship and continue to support the conclusions from the 2006 O_3 AOCD, which 33 stated that "if a population threshold level exists in O_3 health effects, it is likely near the 34 lower limit of ambient O_3 concentrations in the United States" (U.S. EPA, 2006b).

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6.6.2.5 Associations of Cause-Specific Mortality and Short-term Ozone Exposure

1 2 3 4 5 6	In the 2006 O_3 AQCD, an evaluation of studies that examined cause-specific mortality found consistent positive associations between short-term O_3 exposure and cardiovascular mortality, with less consistent evidence for associations with respiratory mortality. The majority of the evidence for associations between O_3 exposure and cause- specific mortality were from single-city studies, which had small daily mortality counts and subsequently limited statistical power to detect associations.
7	New multicity studies evaluated in this review build upon and confirm the associations
8	between short-term O_3 exposure and cause-specific mortality identified in the 2006 O_3
9	AQCD (U.S. EPA, 2006b) (Figure 6-36; Table 6-53). In APHENA, a multicontinent
10	study that consisted of the NMMAPS, APHEA2 and Canadian multicity datasets,
11	consistent positive associations were reported for both cardiovascular and respiratory
12	mortality in all-year analyses when focusing on the natural spline model with 8 df/year
13	(Figure 6-36; Table 6-53). The associations between O_3 exposure and cardiovascular and
14	respiratory mortality in all-year analyses were further supported by the multicity PAPA
15	study (<u>Wong et al., 2010</u>). The magnitude of cardiovascular mortality associations were
16	primarily larger in analyses restricted to the summer season compared to those observed
17	in all-year analyses (Figure 6-36; Table 6-53). Additional multicity studies from the U.S.
18	(Zanobetti and Schwartz, 2008b) and Europe (Stafoggia et al., 2010; Samoli et al., 2009)
19	that conducted summer season analyses provide evidence supporting associations
20	between O ₃ exposure and cardiovascular and respiratory mortality that are similar or
21	larger in magnitude compared to those observed in all-year analyses.
22	Of the studies evaluated, only the APHENA study (Katsouyanni et al., 2009) and an
23	Italian multicity study (Stafoggia et al., 2010) conducted an analysis of the potential for
24	copollutant confounding of the O3 cause-specific mortality relationship. When focusing
25	on the natural spline model with 8 df/year and lag 1 results (as discussed in
26	Section <u>6.6.2.1</u>), the APHENA study found that O_3 cause-specific mortality risk estimates
27	were fairly robust to the inclusion of PM_{10} in copollutant models in the European dataset
28	with more variability in the U.S. and Canadian datasets (i.e., copollutant risk estimates
29	increased and decreased for respiratory and cardiovascular mortality). In summer season
30	analyses cardiovascular O_3 mortality risk estimates were robust in the European dataset
31	and attenuated but remained positive in the U.S. datasets; whereas, respiratory O_3
32	mortality risk estimates were attenuated in the European dataset and robust in the U.S.
33	dataset. The authors did not examine copollutant models during the summer season in the
34	Canadian dataset (Figure 6-29; Table 6-45). Interpretation of these results requires
35	caution; however, due to the different PM sampling schedules employed in each of these

1	study locations (i.e., primarily every-6th day in the U.S. and Canadian datasets and
2	every-day in the European dataset). The results of the summer season analyses from the
3	APHENA study (Katsouyanni et al., 2009) are consistent with those from a study of 10
4	Italian cities during the summer months (Stafoggia et al., 2010). Stafoggia et al. (2010)
5	found that cardiovascular (14.3% [95% CI: 6.7, 22.4%]) and cerebrovascular (8.5%
6	[95% CI: 0.06, 16.3%]) mortality O_3 effect estimates were robust to the inclusion of PM_{10}
7	in copollutant models (14.3% [95% CI: 6.7, 23.1%] and 7.3% [95% CI: -1.2, 16.3],
8	respectively), while respiratory mortality O_3 effects estimates (17.6% [95% CI: 1.8,
9	35.5%]) were attenuated, but remained positive (9.2% [95% CI: -6.9, 28.8%]).

udy	Location	Ages	Lag							
ell et al. (2005)a ong et al. (2010)	U.S. and non-U.S. PAPA (4 cities)	All	NR 0-1	Cardiovascular					All-Ye	ar
tsouyanni et al. (2009)	APHENA-U.S.	≥75	DL(0-2)							
	APHENA-Canada		DL(0-2)							
	APHENA-Canada		DL(0-2)b							
	APHENA-Europe		DL(0-2)							
	APHENA-U.S. APHENA-Canada	<75	DL(0-2) DL(0-2)			-				
	APHENA-Canada APHENA-Canada		DL(0-2) DL(0-2)b			,		_		
	APHENA-Europe		DL(0-2)							
	All tella Europe		DE(0 2)		•					
paris et al. (2004)a	21 European cities	All	0-1						Summ	ıer
moli et al. (2009)	21 European cities		0-1							
nobetti and Schwartz (2008) foggia et al. (2010)	48 U.S. cities 10 Italian cities	≥35	0-3 DL(0-5)		-0-		\sim			
souyanni et al. (2010)	APHENA-U.S.	≥35 ≥75	DL(0-3) DL(0-2)							
souyanni et al. (2005)	APHENA-Canada	275	DL(0-2)							
	APHENA-Canada		DL(0-2)b		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~					
	APHENA-Europe		DL(0-2)		Ĩ — — — — — — — — — — — — — — — — — — —					
	APHENA-U.S.	<75	DL(0-2)							
	APHENA-Canada		DL(0-2)							
	APHENA-Canada		DL(0-2)b		Ŷ					
	APHENA-Europe		DL(0-2)							
l et al. (2005)a	U.S. and non-U.S.	All	NR	Respiratory					All-Ye	ar
ong et al. (2010)	PAPA (4 cities)		0-1		+					
souyanni et al. (2009)	APHENA-U.S.		DL(0-2)	-						
	APHENA-Canada APHENA-Canada		DL(0-2)	•						
	APHENA-Canada APHENA-Europe		DL(0-2)b DL(0-2)							
	APHENA-U.S.	≥75	DL(0-2)							
	APHENA-Canada	275	DL(0-2)	← ●						
	APHENA-Canada		DL(0-2)b							
	APHENA-Europe		DL(0-2)	-						
/paris et al. (2004)a	21 European cities	All	0-1						Summ	hor
obetti and Schwartz (2008)	48 U.S. cities		0-3		—————					
Katsouyanni et al. (2009)	APHENA-U.S.		DL(0-2)							
	APHENA-Canada		DL(0-2)						O	
	APHENA-Canada		DL(0-2)b		<u> </u>					
moli et al. (2000)	APHENA-Europe		DL(0-2) 0-1							
noli et al. (2009) foggia et al. (2010)	21 European cities 10 Italian cities	≥35	DL(0-5)					<u> </u>		
souyanni et al. (2010)	APHENA-U.S.	≥35 ≥75	DL(0-3)				_	0		
	APHENA-Canada	2.5	DL(0-2)							
	APHENA-Canada		DL(0-2)b					~		
	APHENA-Europe		DĽ(0-2)	-						
				-10 -5	0 5	10	15	20	25	
								20	25	

Effect estimates are for a 20 ppb increase in 24-h avg; 30 in 8-h max; and 40ppb increase in 1-h max O_3 concentrations. Red = cardiovascular; blue = respiratory; closed circles = all-year analysis; and open circles = summer-only analysis. An "a" represents studies from the 2006 O_3 AQCD. A "b" represents risk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations (Section <u>6.2.7.2</u>).

Figure 6-36 Percent increase in cause-specific mortality.

Study*	Location	Ages	Lag	Avg Time	%Increase (95% CI
Cardiovascular					
All-year - Cardiovascular					
<u>Bell et al. (2005</u>) ^a	U.S. and non-U.S.	All	NR	24-h avg	2.23 (1.36,3.08)
Wong et al. (2010)	PAPA (4 cities)		0-1	8-h max	2.20 (0.06, 4.37)
Katsouyanni et al. (2009)	APHENA-U.S.	≥ 75	DL(0-2)	1-h max	2.30 (-1.33, 6.04)
	APHENA-Canada		DL(0-2)		8.96 (0.75,18.6)
	APHENA-Canada		DL(0-2) ^b		1.1 (0.10,2.20)
	APHENA-europe		DL(0-2)		2.06 (-0.24, 4.31)
	APHENA-U.S.	<75	DL(0-2)		3.83 (-0.16, 7.95)
	APHENA-Canada		DL(0-2)		7.03 (-2.71, 17.7)
	APHENA-Canada		DL(0-2) ^b		0.87 (-0.35, 2.10)
	APHENA-europe		DL(0-2)		1.98 (-1.09, 5.13)
Summer – Cardiovascular					
Gryparis et al. (2004) ^a	21 European cities	All	0-1	8-h max	2.7 (1.29,4.32)
Samoli et al. (2009)	21 European cities		0-1	8-h max	1.48 (0.18, 2.80)
Zanobetti and Schwartz (2008b)	48 U.S. cities		0-3	8-h max	2.42 (1.45, 3.43)
Stafoggia et al. (2010)	10 Italian cities	≥ 35	DL(0-5)	8-h max	14.3 (6.65, 22.4)
Katsouyanni et al. (2009)	APHENA-U.S.	≥ 75	DL(0-2)	1-h max	3.18 (-0.47, 6.95)
	APHENA-Canada		DL(0-2)		1.50 (-2.79, 5.95)
	APHENA-Canada		DL(0-2) ^b		0.19 (-0.36, 0.74)
	APHENA-europe		DL(0-2)		3.67 (0.95, 6.53)
	APHENA-U.S.	<75	DL(0-2)		6.78 (2.70, 11.0)
	APHENA-Canada		DL(0-2)		-1.02 (-4.23, 2.30)
	APHENA-Canada		DL(0-2) ^b		-0.13 (-0.55, 0.29)
	APHENA-europe		DL(0-2)		2.22 (-1.48, 6.04)
Respiratory					
All-years - Respiratory					
Bell et al. (2005) ^a	U.S. and non-U.S.	All	NR	24-h avg	0.94 (-1.02, 2.96)
Wong et al. (2010)	PAPA (4 cities)		0-1	8-h max	2.02 (-0.41, 4.49)
Katsouyanni et al. (2009)	APHENA-U.S.		DL(0-2)	1-h max	2.54 (-3.32, 8.79)
	APHENA-Canada		DL(0-2)		1.02 (-11.9, 15.9)
	APHENA-Canada		DL(0-2) ^b		0.13 (-1.60, 1.90)
	APHENA-europe		DL(0-2)		1.82 (-2.18, 6.04)
	APHENA-U.S.	≥ 75	DL(0-2)		1.10 (-6.48, 9.21)
	APHENA-Canada		DL(0-2)		-4.61 (-19.3, 13.3)
	APHENA-Canada		DL(0-2) ^b		-0.60 (-2.70, 1.60)
	APHENA-europe		DL(0-2)		1.10 (-3.48, 5.95)

Table 6-53Corresponding effect estimates for Figure 6-36.

Study*	Location	Ages	Lag	Avg Time	%Increase (95% CI
Summer - Respiratory					
Gryparis et al. (2004) ^a	21 European cities	All	0-1	8-h max	6.75 (4.38, 9.10)
Zanobetti and Schwartz (2008b)	48 U.S. cities		0-3	8-h max	2.51 (1.14, 3.89)
Katsouyanni et al. (2009)	APHENA-U.S.		DL(0-2)	1-h max	4.40 (-2.10, 11.3)
	APHENA-Canada		DL(0-2)		26.1 (13.3, 41.2)
	APHENA-Canada		DL(0-2) ^b		3.00 (1.60, 4.50)
	APHENA-europe		DL(0-2)		3.83 (-1.33, 9.21)
Samoli et al. (2009)	21 European cities		0-1	8-h max	2.38 (0.65, 4.19)
<u>Stafoggia et al. (2010)</u>	10 Italian cities	≥ 35	DL(0-5)	8-h max	17.6 (1.78, 35.5)
Katsouyanni et al. (2009)	APHENA-U.S.	≥ 75	DL(0-2)	1-h max	4.07 (-4.23, 13.0)
	APHENA-Canada		DL(0-2)		19.5 (2.22, 40.2)
	APHENA-Canada		DL(0-2) ^b		2.30 (0.28, 4.40)
	APHENA-europe		DL(0-2)		2.46 (-3.40, 8.62)

*Studies from Figure 6-36, plus others.

^aStudies from the 2006 O_3 AQCD.

^bRisk estimates from APHENA-Canada standardized to an approximate IQR of 5.1 ppb for a 1-h max increase in O_3 concentrations (Section <u>6.2.7.2</u>).

1	Collectively, the results from the new multicity studies provide evidence of associations
2	between short-term O_3 exposure and cardiovascular and respiratory mortality with
3	additional evidence indicating these associations persist, and in some cases the magnitude
4	of associations are increased, in the summer season. Although copollutant analyses of
5	cause-specific mortality are limited, the APHENA study found that O_3 cause-specific
6	mortality risk estimates were fairly robust to the inclusion of PM_{10} in copollutant models
7	when focusing on the dataset with daily PM_{10} data (i.e., the European dataset), which is
8	supported by the results from Stafoggia et al. (2010). Additionally, APHENA found that
9	O_3 cause-specific mortality risk estimates were moderately to substantially sensitive
10	(e.g., increased or attenuated) to inclusion of PM_{10} in the U.S. and Canadian datasets.
11	However, the mostly every-6th-day sampling schedule for PM_{10} in the U.S. and Canadian
12	datasets greatly reduced their sample size and limits the interpretation of these results.

6.6.3 Summary and Causal Determination

13	The evaluation of new multicity studies that examined the association between short-term
14	O_3 exposure and mortality found evidence which supports the conclusions of the 2006 O_3
15	AQCD. These new studies reported consistent positive associations between short-term
16	O3 exposure and all-cause (nonaccidental) mortality, with associations persisting or
17	increasing in magnitude during the warm season, and provide additional support for
18	associations between O ₃ exposure and cardiovascular and respiratory mortality.

1	Recent studies further examined potential confounders (e.g., copollutants and seasonality)
2	of the O_3 -mortality relationship. Because the PM- O_3 correlation varies across regions,
3	due to the difference in PM chemical constituents, interpretation of the combined effect
4	of PM on the relationship between O_3 and mortality is not straightforward. Unlike
5	previous studies that were limited to primarily examining the confounding effects of
6	PM_{10} , the new studies expanded their analyses to include multiple PM indices (e.g., PM_{10} ,
7	$PM_{2.5}$, and PM components). An examination of copollutant models found evidence that
8	associations between O_3 and all-cause mortality were robust to the inclusion of PM_{10} or
9	$PM_{2.5}$ (Stafoggia et al., 2010; Katsouyanni et al., 2009; Bell et al., 2007), while other
10	studies found evidence for a modest reduction (~20-30%) when examining PM_{10} (Smith
10	et al., 2009b). Additional evidence suggests potential sensitivity (e.g., increases and
12	attenuation) of O_3 mortality risk estimates to copollutants by age group or cause-specific
12	mortality (e.g., respiratory and cardiovascular) (<u>Stafoggia et al., 2010</u> ; <u>Katsouyanni et al.</u> ,
13	2009). An examination of PM components, specifically sulfate, found evidence for
15	reductions in O_3 -mortality risk estimates in copollutant models (<u>Franklin and Schwartz</u> ,
15	2008). Overall, across studies, the potential impact of PM indices on O ₃ -mortality risk
17	
17	estimates tended to be much smaller than the variation in O_3 -mortality risk estimates
	across cities suggesting that O_3 effects are independent of the relationship between PM
19	and mortality. However, interpretation of the potential confounding effects of PM on
20	O_3 -mortality risk estimates requires caution. This is because the PM- O_3 correlation varies
21	across regions, due to the difference in PM components, complicating the interpretation
22	of the combined effect of PM on the relationship between O_3 and mortality. Additionally,
23	the limited PM or PM component datasets used as a result of the every-3rd- and 6th-day
24	PM sampling schedule instituted in most cities limits the overall sample size employed to
25	examine whether PM or one of its components confounds the O ₃ -mortality relationship.
26	An examination of potential seasonal confounding of the O ₃ -mortality relationship found
27	that the extent of smoothing or the methods used for adjustment can influence O_3 risk
28	estimates when not applying enough degrees of freedom to control for temporal/season
29	trends (Katsouyanni et al., 2009). This is because of the opposing seasonal trends
30	between O_3 and mortality.
31	The multicity studies evaluated within this section also examined the regional
32	heterogeneity observed in O ₃ -mortality risk estimates. These studies provide evidence
33	which suggests generally higher O_3 -mortality risk estimates in northeastern U.S. cities
34	with some regions showing no associations between O_3 exposure and mortality
35	(e.g., Southwest, Urban Midwest) (Smith et al., 2009b; Bell and Dominici, 2008).
36	Multicity studies that examined individual- and community-level characteristics
37	identified characteristics that may explain the observed regional heterogeneity in
38	O_3 -mortality risk estimates as well as characteristics of populations potentially at greatest
-	

- 1 risk for O₃-related health effects. An examination of community-level characteristics 2 found an increase in the O_3 -mortality risk estimates in cities with higher unemployment, 3 percentage of the population Black/African-American, percentage of the working 4 population that uses public transportation, lower temperatures, and lower prevalence of 5 central air conditioning (Medina-Ramón and Schwartz, 2008). Additionally, a potential 6 interactive, or synergistic, effect on the O_3 -mortality relationship was observed when 7 examining differences in the O_3 -mortality association across temperature levels (Ren et 8 al., 2008). An examination of individual-level characteristics found evidence that older 9 age, female sex, Black race, having atrial fibrillation, SES indicators (i.e., educational 10 attainment, income level, and employment status), and out-of hospital deaths, specifically 11 in those individuals with diabetes, modify O₃-mortality associations (Cakmak et al., 12 2011; Stafoggia et al., 2010; Medina-Ramón and Schwartz, 2008), and lead to increased 13 risk of O₃-related mortality. Overall, additional research is warranted to further confirm 14 whether these characteristics, individually or in combination, can explain the observed 15 regional heterogeneity.
- 16 Additional studies were evaluated that examined factors that may influence the shape of 17 the O_3 -mortality C-R curve, such as multi-day effects, mortality displacement, adaptation, the use of different exposure metrics (i.e., 24-h avg, 8-h max or 1-h max), and whether a 18 19 threshold exists in the O_3 -mortality relationship. An examination of multiday effects in a 20 U.S. and European multicity study found conflicting evidence for mortality displacement, 21 but both studies suggest that the positive associations between O_3 and mortality are 22 observed mainly in the first few days after exposure (Samoli et al., 2009; Zanobetti and 23 Schwartz, 2008b). A U.S. multicity study found evidence of an adaptive response to O_3 24 exposure, with the highest risk estimates earlier in the O_3 season (i.e., July) and 25 diminished effects later (i.e., August) (Zanobetti and Schwartz, 2008a). However, the 26 evidence of adaptive effects has an implication for the interpretation of multi-day effects, 27 and requires further analysis. The limited number of studies conducted that examined the 28 effect of using different exposure metrics (i.e., 1-h max, 8-h max, and 24-h avg) when 29 examining the O_3 -mortality relationship found relatively comparable O_3 -mortality risk 30 estimates across the exposure metrics used (Smith et al., 2009b; Gryparis et al., 2004). 31 Analyses that specifically focused on the O₃-mortality C-R relationship supported a linear 32 O_3 -mortality relationship and found no evidence of a threshold within the range of O_3 33 concentrations in the U.S., but did observe evidence for potential differences in the C-R 34 relationship across cities (Katsouyanni et al., 2009; Stylianou and Nicolich, 2009; Bell et 35 al., 2006). Collectively, these studies support the conclusions of the 2006 O_3 AQCD that 36 "if a population threshold level exists in O_3 health effects, it is likely near the lower limit 37 of ambient O₃ concentrations in the U.S."

1	Studies that examined the association between short-term O ₃ exposure and cause-specific
2	mortality confirm the associations with both cardiovascular and respiratory mortality
3	reported in the 2006 O ₃ AQCD (<u>Stafoggia et al., 2010</u> ; <u>Wong et al., 2010</u> ; <u>Katsouyanni et</u>
4	al., 2009; Samoli et al., 2009; Zanobetti and Schwartz, 2008b). These associations were
5	primarily larger in magnitude during the summer season compared to all-year analyses.
6	Of the studies that examined the potential confounding effects of PM [i.e., Stafoggia et al.
7	(2010); Katsouyanni et al. (2009)], O_3 mortality associations remained relatively robust
8	in copollutant models, but interpretation of these studies was complicated by the different
9	PM sampling schedules (e.g., every-6th-day) employed in each study. Overall, the strong
10	evidence for respiratory effects due to short-term O_3 exposure (Section <u>6.2</u>) are consistent
11	across disciplines and provides coherence for the respiratory mortality associations
12	observed across studies. However, the strong evidence for O_3 -induced cardiovascular
13	mortality is complicated by toxicological studies that provide initial evidence for a
14	biologically plausible mechanism for O3-induced cardiovascular mortality, but a lack of
15	coherence with controlled human exposure and epidemiologic studies of cardiovascular
16	morbidity that do not demonstrate consistent evidence of O3-induced cardiovascular
17	effects (Section 6.3).
18	In conclusion, the recent epidemiologic studies build upon and confirm the associations
19	between short-term O3 exposure and all-cause and cause-specific mortality reported in the
20	2006 O ₃ AQCD. However, there is a lack of coherence across disciplines and consistency
21	across health outcomes for O_3 -induced cardiovascular morbidity (Section <u>6.3</u>) which do
22	not support the relatively strong epidemiologic evidence for O3-related cardiovascular
23	mortality. Overall, recent studies have provided additional information regarding key
24	uncertainties (previously identified - including the potential confounding effects of
25	copollutants and seasonal trend), individual- and community-level factors that may lead
26	to increased risk of O_3 -induced mortality and the heterogeneity in O_3 -mortality risk
27	estimates, and continued evidence of a linear no-threshold C-R relationship. Although
28	some uncertainties still remain, the collective body of evidence is sufficient to conclude
29	that there is likely to be a causal relationship between short-term O_3 exposure and
30	total mortality.

6.7 Overall Summary

31	The evidence reviewed in this chapter describes the recent findings regarding the health
32	effects of short-term exposure to ambient O_3 concentrations. <u>Table 6-54</u> provides an
33	overview of the causal determinations for each of the health categories evaluated.

Table 6-54Summary of causal determinations for short-term exposures to
ozone.

Health Category	Causal Determination
Respiratory Effects	Causal relationship
Cardiovascular Effects	Suggestive of a causal relationship
Central Nervous System Effects	Suggestive of a causal relationship
Effects on Liver and Xenobiotic Metabolism	Inadequate to infer a causal relationship
Effects on Cutaneous and Ocular Tissues	Inadequate to infer a causal relationship
Total Mortality	Likely to be a causal relationship

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7 INTEGRATED HEALTH EFFECTS OF LONG-TERM OZONE EXPOSURE

7.1 Introduction

1 2 3 4 5	This chapter reviews, summarizes, and integrates the evidence on relationships between health effects and long-term exposures to O_3 . Both epidemiologic and toxicological studies provide a basis for examining long-term O_3 exposure health effects for respiratory effects, cardiovascular effects, reproductive and developmental effects, central nervous system effects, cancer outcomes, and mortality. Long-term exposure has been defined as
6	a duration of approximately 30 days (1 month) or longer ¹ . However, in order to
7	characterize the weight of evidence for the effects of O_3 on reproductive and
8	developmental effects in a consistent, cohesive and integrated manner, results from both
9	short-term and long-term exposure periods are included in that section, and are identified
10	accordingly in the text and tables.
11	Conclusions from the 2006 O ₃ AQCD (U.S. EPA, 2006b) are summarized briefly at the
12	beginning of each section, and the evaluation of evidence from recent studies builds upon
13	what was available during the previous review. For each health outcome (e.g., respiratory
14	disease, lung function), results are summarized for studies from the specific scientific
15	discipline, i.e., epidemiologic and toxicological studies. The major sections
16	(i.e., respiratory, cardiovascular, mortality, reproductive/developmental, cancer) conclude
17	with summaries of the evidence for the various health outcomes within that category and
18	integration of the findings that lead to conclusions regarding causality based upon the
19	framework described in the Preamble to this ISA. Determination of causality is made for
20	the overall health effect category, such as respiratory effects, with coherence and
21	plausibility being based on evidence from across disciplines and also across the suite of
22	related health outcomes, including cause-specific mortality.
23	As mentioned in Chapter $\underline{2}$ (Section $\underline{2.3}$), epidemiologic studies generally present O ₃ -
24	related effect estimates for mortality and morbidity health outcomes based on an
25	incremental change in exposure. Studies traditionally present the relative risk per an
26	incremental change equal to the interquartile range in O ₃ concentrations or some other
27	arbitrary value (e.g., 10 ppb). Additionally, various exposure metrics are used in O_3
28	epidemiologic studies, with the three most common being the maximum 1-h average
29	within a 24-h period (1-h max), the maximum 8-h average within a 24-h period
30	(8-h max), and 24-h average (24-h avg). For the purpose of presenting results from

¹ Unless otherwise specified, the term "chronic" generally refers to an annual exposure duration for epidemiology studies and a duration of greater than 10% of the lifespan of the animal in toxicological studies.

1 studies that use different exposure metrics, EPA consistently applies the same O₃ 2 increments to facilitate comparisons between the results of various studies that may 3 present results for different incremental changes. Differences due to the use of varying 4 exposure metrics (e.g., 1-h max, 24-h avg) become less apparent when averaged across 5 longer exposure periods, because levels are typically lower and less variable. As such, 6 throughout this chapter an increment of 10 ppb was consistently applied across studies, 7 regardless of exposure metric, to facilitate comparisons between the results from these 8 studies.

7.2 Respiratory Effects

9	Studies reviewed in the 2006 O ₃ AQCD examined evidence for relationships between
10	long-term O ₃ exposure (several months to yearly) and effects on respiratory health
11	outcomes including declines in lung function, increases in inflammation, and
12	development of asthma in children and adults. Animal toxicology data provided a clearer
13	picture indicating that long-term O ₃ exposure may have lasting effects. Chronic exposure
14	studies in animals have reported biochemical and morphological changes suggestive of
15	irreversible long-term O ₃ impacts on the lung. In contrast to supportive evidence from
16	chronic animal studies, the epidemiologic studies on longer-term (annual) lung function
17	declines, inflammation, and new asthma development remained inconclusive.
18	Several studies reviewed in the 2006 O ₃ AQCD (Horak et al., 2002; Frischer et al., 1999)
19	collectively indicated that O ₃ exposure averaged over several summer months was
20	associated with smaller increases in lung function growth in children. For longer
21	averaging periods (annual), the definitive analysis in the Children's Health Study (CHS)
22	reported by Gauderman et al. (2004) provided little evidence that such long-term
23	exposure to ambient O_3 was associated with significant deficits in the growth rate of lung
24	function in children in contrast to the effects observed with other pollutants such as acid
25	vapor, NO ₂ , and PM _{2.5} . Limited epidemiologic research examined the relationship
26	between long-term O3 exposures and inflammation. Consistent with evidence of
27	inflammation and allergic responses reported in experimental studies, an association
28	between 30-day average O_3 and increased eosinophil levels was observed in an Austrian
29	study (Frischer et al., 2001). The cross-sectional studies available for the 2006 O_3 AQCD
30	detected no associations between long-term O3 exposures and asthma prevalence, asthma-
31	related symptoms or allergy to common aeroallergens in children after controlling for
32	covariates. However, longitudinal studies provided evidence that long-term O3 exposure
33	influences the risk of asthma development in children (McConnell et al., 2002) and adults
34	(McDonnell et al., 1999a; Greer et al., 1993).

1	New evidence presented below reports interactions between genetic variants and long-
2	term O ₃ exposure in effects on new onset asthma in U.S. cohorts in multi-community
3	studies where protection by specific oxidant gene variants was restricted to children
4	living in low O3 communities. Related studies report coherent relationships between
5	respiratory symptoms among asthmatics and long-term O_3 exposure. Short-term exposure
6	to O_3 is associated with increases in respiratory symptoms and asthma medication use in
7	children with asthma (Section $6.2.4.1$) and asthma hospitalizations in children
8	(Section <u>6.2.7.2</u>). A new line of evidence reports a positive concentration-response
9	relationship between first asthma hospitalization and long-term O_3 exposure. Results
10	from recent studies examining pulmonary function, inflammation, and allergic responses
11	are also presented.

7.2.1 Asthma

7.2.1.1 New Onset Asthma

12 Asthma is a heterogeneous disease with a high degree of temporal variability. Its 13 progression and symptoms can vary within an individual's experience over time. The 14 course of asthma may vary markedly between young children, older children and 15 adolescents, and adults. This variation is probably more dependent on age than on 16 symptoms (NHLBI, 2007). Longitudinal cohort studies have examined associations 17 between long-term O₃ exposures and the onset of asthma in adults and children (McConnell et al., 2002; McDonnell et al., 1999a; Greer et al., 1993), with results 18 19 indicating a direct effect of long-term O_3 exposure on asthma risk in adults and effect 20 modification by O_3 in children. 21 Associations between long-term O_3 exposure and new cases of asthma were reported in a 22 cohort of nonsmoking adults in California (McDonnell et al., 1999a; Greer et al., 1993). 23 The Adventist Health and Smog (AHSMOG) study cohort of 3,914 (age 27 to 87 years, 24 36% male) was drawn from nonsmoking, non-Hispanic white California Seventh Day 25 Adventists, who were surveyed in 1977, 1987, and 1992. To be eligible, subjects had to 26 have lived 10 or more years within 5 miles of their current residence in 1977. Residences 27 from 1977 onward were followed and linked in time and space to interpolate 28 concentrations of O_3 , PM_{10} , SO_4^{2-} , SO_2 , and NO_2 . New asthma cases were defined as self-29 reported doctor-diagnosed asthma at either the 1987 or 1992 follow-up questionnaire 30 among those who had not reported having asthma upon enrollment in 1977. During the 31 10-year follow-up (1977 to 1987), the incidence of new asthma was 2.1% for males and

32 2.2% for females (<u>Greer et al., 1993</u>). Ozone concentration data were not provided. A

- relative risk of 3.12 (95% CI: 1.16, 5.85) per 10-ppb increase in annual mean O₃
 (exposure metric not stated) was observed in males, compared to a relative risk of 0.94
 (95% CI: 0.65, 1.34) in females.
- 4 In the 15-year follow-up study (1977-1992), 3.2% of the eligible males and a slightly 5 greater 4.3% of the eligible females developed adult asthma (McDonnell et al., 1999a). 6 The mean 20-year average (1973-1992) for 8-h avg O₃ (9 a.m. to 5 p.m.) was 46.5 ppb 7 (SD 15.3). For males, the relative risk of developing asthma was 1.31 (95% CI: 1.01, 8 1.71) per 10-ppb increase in 8-h avg O₃. Once again, there was no evidence of a positive 9 association between O_3 and new-onset asthma in females (relative risk of 0.94 [95% CI: 10 (0.87, 1.02)). The lack of an association does not necessarily indicate no effect of O₃ on 11 the development of asthma among females. For example, differences between females 12 and males in time-activity patterns may influence relative exposures to ambient O_3 . 13 During summer 1992, the mean (SD) hours per week spent outdoors for male and female 14 asthma cases were 13.8 (10.6) and 11.4 (10.9), respectively, indicating potential greater misclassification of exposure in females. None of the other pollutants (PM_{10} , SO_4^{2-} , SO_2 , 15 16 and NO₂) were associated with development of asthma in either males or females. 17 Adjusting for copollutants did not diminish the association between O_3 and asthma incidence for males. In no case was the O₃ coefficient reduced by more than 10% in the 18 19 two-pollutant models compared to the model containing O_3 alone. The consistency of the 20 results in the two studies with different follow-up times, as well as the independent and 21 robust association between annual mean O_3 concentrations and asthma incidence, provide 22 supportive evidence that long-term O_3 exposure may be associated with the development 23 of asthma in adult males. However, because the AHSMOG cohort was drawn from a 24 narrow subject definition, the representativeness of this cohort to the general U.S. 25 population may be limited.
- 26 In children, the relationship between long-term O_3 exposure and new onset asthma has 27 been extensively investigated in the CHS. In this cohort, evidence provides stronger 28 support for long-term O_3 exposure modifying the risk of new onset asthma associated 29 with other potential risk factors than having a main effect on new onset asthma. Initiated 30 in the early 1990s, the CHS was originally designed to examine whether long-term 31 exposure to ambient pollutants was related to chronic respiratory outcomes in children in 32 12 communities in southern California (Peters et al., 1999b; Peters et al., 1999a). New-33 onset asthma was classified as having no prior history of asthma at study entry with 34 subsequent report of physician-diagnosed asthma at follow-up with the date of onset 35 assigned to be the midpoint of the interval between the interview date when asthma 36 diagnosis was first reported and the previous interview date. In a cohort recruited during 37 2002-2003 and followed for three years beginning in kindergarten or first grade, 38 McConnell et al. (2010) reported a hazard ratio for new onset asthma of 0.76 (95% CI:

10.38, 1.54) comparing the communities with the highest (59.8 ppb) and lowest (29.5 ppb)2annual average of 8-h avg (10 a.m.-6 p.m.) O3. With adjustment for school and residential3modeled non-freeway traffic-related exposure, the estimated HR for O3 was 1.014(95% CI: 0.49, 2.11).

5 Similarly in a cohort recruited in 1993, asthma risk was not higher for residents of the six 6 high-O₃ communities versus residents of the six low-O₃ communities (McConnell et al., 7 2002). In this study, 3,535 initially nonasthmatic children (ages 9 to 16 years at 8 enrollment) were followed for up to 5 years, during which 265 cases of new-onset asthma 9 were identified. Communities were stratified by 4-year average 1-h max O₃ levels, with 10 six high-O₃ communities (mean 75.4 ppb [SD 6.8]) and six low-O₃ communities (mean 50.1 ppb [SD 11.0]). Within the high-O₃ communities, asthma risk was 3.3 (95% CI: 1.9, 11 12 5.8) times greater for children who played three or more sports as compared with children 13 who played no sports. None of the children who lived in high-O₃ communities and played 14 three or more sports had a family history of asthma. In models with individual sports 15 entered as dummy variables, only tennis was significantly associated with asthma and 16 only in the high O_3 communities. This association was absent in the low- O_3 communities 17 (relative risk of 0.8 [95% CI: 0.4, 1.6]). The overall observed pattern of effects of sports 18 participation on asthma risk was robust to adjustment for SES, history of allergy, family 19 history of asthma, insurance, maternal smoking, and BMI.

20 Analyses aimed at distinguishing the effects of O_3 from effects of other pollutants 21 indicated that in communities with high O_3 and low levels of other pollutants there was a 22 4.2-fold (95% CI: 1.6, 10.7) increased risk of asthma in children playing three or more 23 sports, compared to children who played no sports. The relative risk in children playing 24 three or more sports was slightly lower (3.3 [95% CI: 1.6, 6.9]) in communities with a 25 combination of high levels of O_3 and other pollutants. Ozone concentrations were not 26 strongly correlated with PM₁₀, PM₂₅, NO₂, or inorganic acid vapors, and no associations 27 with asthma were found for these other pollutants. These results provide additional 28 support that the effects of physical activity on asthma are modified by long-term 29 O₃ exposure. Overall, the results from McConnell et al. (2002) suggest that playing sports 30 may indicate greater outdoor activity when O₃ levels are higher and an increased 31 ventilation rate, which may lead to increased O_3 exposure. It should be noted, however, 32 that these findings were based on a small number of new asthma cases (n = 29 among 33 children who played three or more sports) and were not based on a priori hypotheses.

34Recent studies from the CHS provide evidence for gene-environment interactions in35effects on new-onset asthma by indicating that the lower risks associated with specific36genetic variants are found in children who live in lower O3 communities (Islam et al.,372009; Islam et al., 2008; Oryszczyn et al., 2007; Lee et al., 2004b; Gilliland et al., 2002).

1 Risk for new-onset asthma is related in part to genetic susceptibility, behavioral factors 2 and environmental exposure (Gilliland et al., 1999). Gene-environment interactions in 3 asthma have been well discussed in the literature (von Mutius, 2009; Holgate et al., 2007; 4 Martinez, 2007a, b; Rahman et al., 2006; Hoffjan et al., 2005; Kleeberger and Peden, 5 2005; Ober, 2005). Complex chronic diseases, such as asthma, are partially the result of a 6 sequence of biochemical reactions involving exposures to various environmental agents 7 metabolized by a number of different genes (Conti et al., 2003). Oxidative stress has been 8 proposed to underlie these mechanistic hypotheses (Gilliland et al., 1999). Genetic 9 variants may impact disease risk directly or modify disease risk by affecting internal dose 10 of pollutants and other environmental agents and/or their reaction products or by altering cellular and molecular modes of action. Understanding the relation between genetic 11 12 polymorphisms and environmental exposure can help identify high-risk subgroups in the 13 population and provide better insight into pathway mechanisms for these complex 14 diseases.

15 CHS analyses have found that asthma risk is related to interactions between O_3 and 16 variants in genes for enzymes such as heme-oxygenase (HO-1), arginases (ARG1 and 2), 17 and glutathione S transferase P1 (GSTP1) (Himes et al., 2009; Islam et al., 2008; Li et al., 18 2008; Hanene et al., 2007; Ercan et al., 2006; Li et al., 2006a; Tamer et al., 2004; 19 Gilliland et al., 2002). Biological plausibility for these findings is provided by evidence 20 that these enzymes have antioxidant and/or anti-inflammatory activity and participate in 21 well recognized modes of action in asthma pathogenesis. Further, several lines of 22 evidence demonstrate that secondary oxidation products of O_3 initiate the key modes of 23 action that mediate downstream health effects (Section 5.3.2). For example, HO-1 has 24 been found to respond rapidly to oxidants, have anti-inflammatory and anti-oxidant 25 effects (Exner et al., 2004), relax airway smooth muscle, and be induced in airways 26 during asthma (Carter et al., 2004). The GSTP1 Val/Val genotype has been associated 27 with increased risk of having atopic asthma (Tamer et al., 2004). Gene-environment 28 interactions are discussed in greater detail in Section 5.4.2.1.

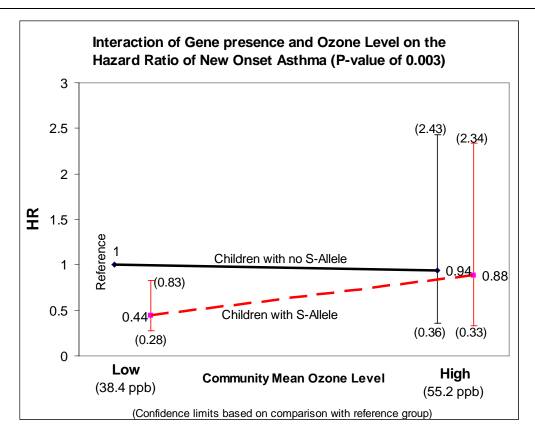
29 Islam et al. (2008) found that functional polymorphisms of the heme oxygenase-1 gene 30 (HMOX-1, [(GT)n repeat]) influenced the risk of new-onset asthma, depending on 31 ethnicity and long-term community O₃ concentrations. Ozone-gene interactions were not 32 found for variants in other antioxidant genes: catalase (CAT [-262C >T -844C >T0]) or 33 and manganese superoxide dismutase (MNSOD, [Ala-9Val]). Analyses were restricted to 34 children of Hispanic (n = 576) or non-Hispanic white ethnicity (n = 1,125) and were 35 conducted with long-term pollutant levels averaged from 1994 to 2003. The effect of 36 ambient air pollution on the relationship between genetic polymorphism and new-onset 37 asthma was assessed using Cox proportional hazard regression models where the community specific average air pollution levels were fitted as a continuous variable 38

together with the appropriate interaction terms for genes and air pollutants and a random effect of community (Berhane et al., 2004).

- 3 Over the follow-up period, 160 new cases of asthma were diagnosed (Islam et al., 2008). 4 For HMOX-1, the interaction (p = 0.003) indicated a greater protective effect of the 5 S-allele (short, <23 (GT)n repeats) compared to the L-allele (long, >23 repeats) among 6 non-Hispanic white children who lived in the low O₃ community (nonparallelism 7 presented in Figure 7-1). Among children residing in low- O_3 communities, the hazard 8 ratio (HR) of new onset asthma associated with the S-allele was 0.44 (95% CI: 0.23, 9 (0.83) compared to non-Hispanic white children who lived in low O₃ communities and had 10 no S-alleles. Biological plausibility for these results is provided by evidence that the S-allele variant of HMOX-1 is more readily induced than those with more numerous 11 12 repeats. The S-allele was found to have a less protective effect in non-Hispanic white 13 children who resided in high O_3 communities (HR = 0.88; [95% CI: 0.33, 2.34] compared 14 to non-Hispanic white children in low O_3 communities with no S-allele). Because 15 HMOX-1 variants were not associated with asthma risk in Hispanic children, effect 16 modification by O_3 was not investigated. No significant interactions were observed 17 between PM_{10} or other pollutants and the HMOX-1 gene; quantitative results were not 18 presented. Average O₃ levels showed low correlation with the other monitored pollutants. 19 The authors did not consider the lack of adjustment for multiple testing to be a concern in 20 this analysis because the selection of the genes was based on a priori hypotheses defined 21 by a well-studied biological pathway, in which oxidative stress serves as the link among 22 O₃ exposure, enzyme activity, and asthma.
- 23 Collectively, results from Islam et al. (2008) indicate that a variant in HMOX-1 that 24 produces a more readily inducible enzyme is associated with lower risk of new-onset 25 asthma in children who live in low O₃ communities. Results were not presented for the 26 main effects relating new-onset asthma to O_3 exposure. However, they do indicate that 27 that in environments of low ambient O_3 , enzymes with greater antioxidative activity may 28 have the capacity to counter any temporary imbalance in an oxidant-antioxidant 29 relationship. However, in the presence of high background O_3 , the protective effect may 30 be attenuated because with higher exposure to oxidants, the antioxidant genes may be at 31 their maximal level of inducibility, and variation in promoters no longer affects levels of 32 expression. Supporting evidence is provided by Schroer et al. (2009), who found that 33 infants with multiple environmental exposures were at increased risk of wheeze 34 regardless of variant in GSTP1, which encodes a gene with antioxidant activity.

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Note: An interaction p-value of 0.003 was obtained from the hierarchical two stage Cox proportional hazard model fitting the community specific O_3 and controlling for random effect of the communities. The interaction indicates there is a greater protective effect of having a heme-oxygenase S-allele compared to having the L-allele among children living in communities with lower long-term ambient ozone concentrations. The HRs are off-set as opposed to overlapping in the figure to allow clearer presentation of the results.

Source: Developed by EPA with data from Islam et al. (2008) (data used with permission of American Thoracic Society).

Figure 7-1 Interaction of heme-oxygenase genetic variants and O₃ level on the Hazard Ratio (HR) of new-onset asthma in the 12 Children's Health Study communities.

1	Expanding on the results of McConnell et al. (2002), Islam et al. (2009) provided
2	evidence that variants in GSTM1 and GSTP1 may influence associations between
3	outdoor exercise and new onset asthma. A primary conclusion that the authors (Islam et
4	al., 2009) reported was that the GSTP1 Ile/Ile and GSTM1 null genotypes increased risk
5	of new onset asthma during adolescence. The highest risk was found for participation in
6	three or more team sports (compared to no sports) among children with GSTP1 Ile/Ile
7	genotype living in high-O ₃ communities (HR: 6.15, [95% CI: 2.2, 7.4]). No three-way
8	interaction was found for GSTM1. These results demonstrate the potential importance of
9	a combination of genetic variability, O ₃ exposure, and outdoor activity on asthma risk. It
10	is important to note that while some studies have found a modifying role of air pollution

1on the association between GSTP1 Ile/Ile and asthma in children (Lee et al., 2004b),2others have found that the GSTP1 Val/Val variant to be associated with greater asthma3prevalence and increase the risk of O3-associated respiratory morbidity (see discussion in4Section 6.2.4.1).

5 The CHS also provided evidence of interactions between O_3 exposure and variants in 6 genes for arginase (Salam et al., 2009). Arginase catalyzes the conversion of L-arginine. 7 Because L-arginine is a precursor of NO, higher arginase activity can limit production of 8 NO and subsequent nitrosative stress. Epidemiologic evidence of associations of arginase 9 variants with asthma are limited (Li et al., 2006a); however, asthmatic subjects have been 10 found to have higher arginase activity than non-asthmatic subjects (Morris et al., 2004). 11 The modifying effect of O₃ and atopy on the association between ARG1 and ARG2 12 haplotypes and asthma were evaluated using likelihood ratio tests with appropriate 13 interaction terms. Having more copies of the ARG1h4 haplotype (compared to having 14 zero copies) was associated with lower odds of asthma, particularly among children with 15 atopic asthma living in high O₃ communities (OR: 0.12; [95% CI: 0.04, 0.43]). Having 16 more copies of the ARG2h3 haplotype (compared to having zero copies) was associated 17 with increased risk of childhood-onset asthma among children in both low and high O_3 18 communities. The implications of findings are somewhat limited because the functional 19 relevance of the ARG1 and ARG2 variants is not clear.

7.2.1.2 Prevalence of Asthma and Asthma Symptoms

- Some cross-sectional studies reviewed in the 2006 O_3 AQCD observed positive relationships between chronic exposure to O_3 and prevalence of asthma and asthmatic symptoms in school children (<u>Ramadour et al., 2000; Wang et al., 1999</u>) while others (<u>Kuo et al., 2002; Charpin et al., 1999</u>) did not. Recent studies provide additional evidence.
- 25 In a cross-sectional nationwide study of 32,672 Taiwanese school children, Hwang et al. 26 (2005) assessed the effects of air pollutants on the risk of asthma. The study population 27 was recruited from elementary and middle schools within 1 km of air monitoring stations. 28 The risk of asthma was related to O_3 in the one-pollutant model. The addition of other 29 pollutants (NO_X, CO₂, SO₂, and PM₁₀), in two-pollutant and three-pollutant models, 30 increased the O₃ risk estimates. The prevalence of childhood asthma was assessed in 31 Portugal by contrasting the risk of asthma between a high O₃ rural area and an area with 32 low O₃ levels (Sousa et al., 2011; Sousa et al., 2009; Sousa et al., 2008). The locations 33 were selected to provide a difference in O_3 levels without the confounding effects of 34 other pollutants. Both evaluation for asthma symptoms and FEV₁ suggested that O₃

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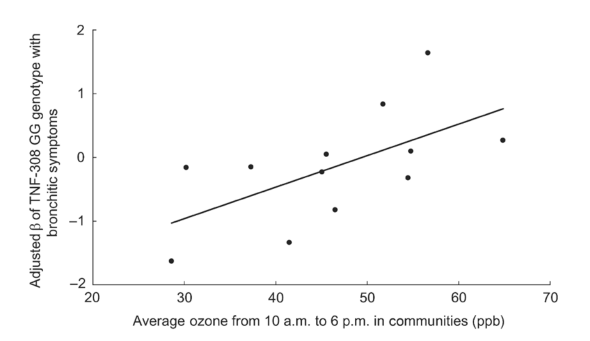
1	
1	increased asthma prevalence. <u>Clark et al. (2010</u>) investigated the effect of exposure to
2	ambient air pollution in utero and during the first year of life on risk of subsequent
3	incidence asthma diagnosis up to 3-4 years of age in a population-based nested case-
4	control study for all children born in southwestern British Columbia in 1999 and 2000
5	(n = 37,401; including 3,482 [9.3%] with asthma). Air pollution exposure for each
6	subject was estimated based on their residential address history using regulatory
7	monitoring data, land use regression modeling, and proximity to stationary pollutant
8	sources. Daily values from the three closest monitors within 50 km were used to calculate
9	exposures. Traffic-related pollutants were associated with the highest risk. Ozone was
10	inversely correlated with the primary traffic-related pollutants ($r = -0.7$ to -0.9). The low
11	reliability of asthma diagnosis in infants makes this study difficult to interpret (Martinez
12	et al., 1995). In a cross-sectional analysis, Akinbami et al. (2010) examined the
13	association between chronic exposure to outdoor pollutants (12-month avg levels by
14	county) and asthma outcomes in a national sample of children ages 3-17 years living in
15	U.S. metropolitan areas (National Health Interview Survey, $N = 34,073$). A 5-ppb
16	increase in estimated 8-h max O_3 concentration (annual average) yielded a positive
17	association for both currently having asthma and for having at least 1 asthma attack in the
18	previous year, while the adjusted odds ratios for other pollutants were not statistically
19	significant. Models in which pollutant value ranges were divided into quartiles produced
20	comparable results. Multi-pollutant models (SO ₂ and PM) produced similar results. The
21	median value for 12-month avg O_3 levels was 39.5 ppb and the IQR was 35.9-43.7 ppb.
22	The adjusted odds for current asthma for the highest quartile (49.9-59.5 ppb) of estimated
23	O ₃ exposure was 1.56 (95% CI: 1.15, 2.10) with a positive concentration-response
24	relationship apparent from the lowest quartile to the highest. Thus, this cross-sectional
25	analysis and <u>Hwang et al. (2005</u>) provides further evidence relating O_3 exposure and the
26	risk of asthma.
27	Relationships between long-term exposure and respiratory symptoms in asthmatic
28	children also were examined in the CHS. McConnell et al. (1999) examined the
29	association between O ₃ levels and the prevalence of chronic lower respiratory tract
30	symptoms in 3,676 cohort children with asthma. In this cross-sectional study, bronchitis,
31	phlegm, and cough were not associated with annual mean 1-h max O ₃ concentrations in
32	children with asthma or wheeze. All other pollutants examined (PM ₁₀ , PM _{2.5} , NO ₂ , and
33	gaseous acid) were associated with an increase in phlegm but not cough. The mean
34	annual average 1-h max O ₃ concentration was 65.6 ppb (range 35.5 to 97.5) across the
35	12 communities. In another CHS analysis, McConnell et al. (2003) evaluated
36	relationships between air pollutants and bronchitic symptoms among 475 children with
37	asthma. The mean 4-year average 8-h avg O_3 (10 a.m6 p.m.) concentration was
38	47.2 ppb (range 28.3 to 65.8) across the 12 communities. For a 10-ppb increase in

8-h avg O_3 averaged over 4 years, the between-community odds ratio was 0.90 (95% CI:

39

1 0.82, 1.00) whereas the within-community (i.e., difference between one- and four-year 2 average) odds ratio was larger, i.e., 1.79 (95% CI: 1.00, 3.21). The authors commented 3 that if the larger within-community effect estimates were correct, then other cross-4 sectional (between-community) studies might have underestimated the true effect of air 5 pollution on bronchitic symptoms in children. These differences might be attributable to 6 confounding by poorly measured or unmeasured risk factors that vary between 7 communities. Within community effects may more accurately represent risk associated 8 with pollutant exposure because the analyses characterize health effects associated with 9 changing pollutant concentrations within a community, thereby minimizing potential 10 confounding by factors that are constant over time within a community. PM_{25} , NO_{2} , and 11 organic carbon also were associated with bronchitic symptoms. In two-pollutant models, 12 the within-community effect estimates for O₃ were markedly reduced and no longer 13 statistically significant in some cases.

14 CHS also examined interactions between TNF- α 308 genotype and long-term O₃ 15 exposure in the occurrence of bronchitic symptoms among children with asthma (Lee et 16 al., 2009b). Increased airway levels of the cytokine TNF- α has been related to 17 inflammation, and the GG genotype has been linked to lower expression of TNF- α . 18 Asthmatic children with the GG genotype had a lower prevalence of bronchitic symptoms 19 compared with children carrying at least one A-allele (e.g., GA or AA genotype). Low-20 versus high- O_3 strata were defined as less than or greater than 50- ppb O_3 avg. Asthmatic 21 children with TNF-308 GG genotype had a significantly reduced risk of bronchitic 22 symptoms with low-O₃ exposure (OR: 0.53 [95% CI: 0.31, 0.91]). The risk was not 23 reduced in children living in high-O₃ communities (OR: 1.42 [95% CI: 0.75, 2.70]). The 24 difference in genotypic effects between low- and high-O₃ environments was statistically 25 significant among asthmatics (P for interaction = 0.01), but not significant among non-26 asthmatic children. Figure 7-2 presents adjusted O₃ community-specific regression 27 coefficients plotted against ambient O₃ concentration, using weights proportional to the 28 inverse variance. Investigators further reported no substantial differences in the effect of 29 the GG genotype on bronchitic symptoms by long-term exposure to PM_{10} , PM_{25} , NO_2 , acid vapor, or second-hand smoke. 30



Note: Using indicator variables for each category of genotype and O_3 exposure, investigators calculated effect estimates for TNF- α GG genotype on the occurrence of bronchitic symptoms among children with asthma. Source: Reprinted with permission of John Wiley & Sons, (Lee et al., 2009b).

Figure 7-2 Ozone modifies the effect of TNF GG genotype on bronchitic symptoms among children with asthma in the CHS.

1 Another CHS analyses reported interrelationships between varia	nts in CAT and
2 myeloperoxidase (MPO) genes, ambient pollutants, and respirate	ory-related school
3 absences for 1,136 Hispanic and non-Hispanic white cohort child	dren (<u>Wenten et al.,</u>
4 <u>2009</u>). A related study (<u>Gilliland et al., 2001</u>), found increased C	₃ exposure to be related
5 greater school absenteeism due to respiratory illness but did not	consider genetic variants.
6 <u>Wenten et al. (2009</u>) hypothesized that variation in the level or f	unction of antioxidant
7 enzymes would modulate respiratory illness risk, especially under	er high levels of
8 oxidative stress expected from high ambient O_3 exposure. The jo	oint effect of variants in
9 these two genes (genetic epistasis) on respiratory illness was exa	mined because the
10 enzyme products operate on the same substrate within the same	biological pathway. Risk
11 of respiratory-related school absences was elevated for children	with CAT GG plus MPO
12 GA or AA genotypes (RR: 1.35 [95% CI: 1.03, 1.77] compared	to GG for both genes)
13 and reduced for children with CAT GA or AA plus MPO GA or	AA (RR: 0.81 [95% CI:
14 0.55, 1.19] compared to GG for both genes). Both CAT GG and	MPO GA or AA
15 genotypes produce a lower activity enzyme. In analyses that stra	tified communities into
16 high and low O ₃ exposure groups by median levels (46.9 ppb), th	ne protective effect of
17 CAT GA or AA plus MPO GA or AA genotype was largely limit	ted to children living in

- 1communities with high ambient O3 levels (RR: 0.42 [95% CI: 0.20, 0.89]). The2association of respiratory-illness absences with functional variants in CAT and MPO that3differ by air pollution levels illustrates the need to consider genetic epistasis in assessing4gene-environment interactions.
- 5 Collective evidence from CHS provides an important demonstration of gene-environment 6 interactions. In the complex gene-environment setting a modifying effect might not be 7 reflected in an exposure main effect. The simultaneous occurrence of main effect and 8 interaction effect can occur. The study of gene-environment interactions helps to dissect 9 disease mechanisms in humans by using information on susceptibility genes to focus on 10 the biological pathways that are most relevant to that disease (Hunter, 2005).
- 11The French Epidemiology study on Genetics and Environment of Asthma (EGEA)12investigated the relationship between ambient air pollution and asthma severity in a
- 13 cohort in five French cities (Paris, Lyon, Marseille, Montpellier, and Grenoble) (Rage et 14 al., 2009b). In this cross-sectional study, asthma severity over the past 12 months was 15 assessed among 328 adult asthmatics using two methods: (1) a four-class severity score 16 that integrated clinical events and type of treatment; and (2) a five-level asthma score 17 based only on symptoms. Two measures of exposure were also assessed: (1) [first 18 method]) closest monitor data from 1991 to 1995 where a total of 93% of the subjects 19 lived within 10 km of a monitor, but where 70% of the O₃ concentrations were 20 back-extrapolated values; and (2) [second method]) a validated spatial model that used 21 geostatistical interpolations and then assigned air pollutants to the geocoded residential 22 addresses of all participants and individually assigned exposure to ambient air pollution
- 23 estimates. Higher asthma severity scores were significantly related to both the 8-h avg O_3 24 during April-September and the number of days with 8-h O_3 averages above 55 ppb. Both 25 exposure assessment methods and severity score methods resulted in very similar 26 findings. Effect estimates of O₃ were similar in three-pollutant models. No PM data were 27 available. Since these estimates were not sensitive to the inclusion of ambient NO_2 in the 28 three-pollutant models, the authors viewed the findings not to be explained by particles 29 which usually have substantial correlations between PM and NO₂. Effect estimates for O₃ 30 in three-pollutant models including O₃, SO₂, and NO₂ yielded OR for O₃-days of 2.74
- (95% CI: 1.68, 4.48) per IQR days of 10-28 (+18) ppb. The effect estimates for SO₂ and
 NO₂ in the three-pollutant model were 1.33 (95% CI: 0.85, 2.11) and 0.94 (95% CI: 0.68,
 1.29) respectively. Taking into account duration of residence did not change the result.
 This study suggests that a higher asthma severity score is related to long-term O₃
 exposure.
- An EGEA follow-up study (Jacquemin et al., In Press), examines the relationship
 between asthma and O₃, NO₂, and PM₁₀. New aspects considered include: (1)

1	examination of three domains of asthma control (symptoms, exacerbations, and lung
2	function); (2) levels of asthma control (controlled, partially controlled, and uncontrolled
3	asthma); and (3) PM_{10} and multi-pollutant analysis. In this cross-sectional analysis,
4	EGEA2 studied 481 adult subjects with current asthma from 2003 to 2007. The IQRs
5	were 11 (41-52) μ g/m ³ for annual O ₃ and 13 (25-38) μ g/m ³ for summer (April-
6	September) O ₃ . The association between asthma control and air pollutants was expressed
7	by ORs (reported for one IQR of the pollutant), derived from multinomial logistic
8	regression. For each factor, the simultaneous assessment of the risk for uncontrolled
9	asthma and for partly controlled asthma was compared with controlled asthma using a
10	composite of the three domains. In crude and adjusted models, O_3 -sum and PM_{10} were
11	positively associated with partly controlled and uncontrolled asthma, with a clear gradient
12	from controlled, partly controlled (OR = 1.53, 95% CI: 1.01, 2.33) and uncontrolled
13	(OR = 2.14, 95% CI: 1.34, 3.43) (from the multinomial logistic regression).

- 14 Separately, they used a composite asthma control classification that used the ordinal 15 logistic regression for risk comparing controlled to partly controlled asthma and 16 comparing partly controlled to uncontrolled asthma. For these two pollutants, the ORs 17 assessed using the ordinal logistic regression were significant (ORs were 1.69 (95% CI: 18 1.22, 2.34) and 1.35 (95% CI: 1.13, 1.64) for O₃-sum and PM₁₀, respectively). For two 19 pollutant models using the ordinal logistic regression, the adjusted ORs for O₃-sum and 20 PM_{10} included simultaneously in a unique model were 1.50 (95% CI: 1.07, 2.11) for O₃-21 sum and 1.28 (95% CI: 1.06, 1.55) for PM_{10} , respectively. This result suggests that the 22 effects of both pollutants are independent.
- 23 The analysis of the associations between air pollution for all asthma subjects and each 24 one of the three asthma control domains showed the following: (1) for lung function 25 defined dichotomously as percent predicted FEV_1 value $\langle or \rangle = 80$ (OR = 1.35, 95% CI: 26 0.80, 2.28 for adjusted O₃-sum); (2) for symptoms defined as asthma attacks or dyspnea 27 or woken by asthma attack or shortness of breath in the past three months (OR = 1.59, 28 95% CI: 1.10, 2.30 for adjusted O_3 -sum); and for exacerbations defined at least one 29 hospitalizations or ER visits in the last year or oral corticosteroids in the past three 30 months (OR = 1.58, 95% CI: 0.97, 2.59 for adjusted O₃-sum). Since the estimates for 31 both pollutants were more stable and significant when using the integrated measure of 32 asthma control, this indicates that the results are not driven by one domain. These results 33 support an effect of long-term exposure to O_3 on asthma control in adulthood in subjects 34 with pre-existing asthma.
- 35Goss et al. (2004) investigated the effect of O_3 on pulmonary exacerbations and lung36function in individuals over the age of 6 years with cystic fibrosis (n = 11,484). The study37included patients enrolled in the Cystic Fibrosis Foundation National Patient Registry,

1	which contains demographic and clinical data collected annually at accredited centers for
2	cystic fibrosis. For 1999 through 2000, the annual mean O_3 concentration, calculated
3	from 1-h averages from 616 monitors in the U.S. EPA Aerometric Information Retrieval
4	System (AIRS), was 51.0 ppb (SD 7.3). Exposure was assessed by linking air pollution
5	values from AIRS with the patient's home ZIP code. No clear association was found
6	between annual mean O_3 and lung function parameters. However, a 10 ppb increase in
7	annual mean O_3 was associated with a 10% (95% CI: 3, 17) increase in the odds of two or
8	more pulmonary exacerbations. Significant excess odds of pulmonary exacerbations also
9	were observed with increased annual mean PM_{10} and $PM_{2.5}$ concentrations. The O_3 effect
10	was robust to adjustment for PM_{10} and $PM_{2.5}$, 8% (95% CI: 1, 15) increase in odds of two
11	or more pulmonary exacerbations per 10 ppb increase in annual mean O ₃ .

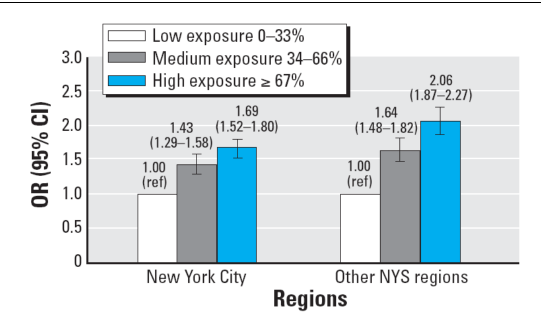
7.2.2 Asthma Hospital Admissions and ED Visits

12	The studies on O ₃ -related hospital discharges and emergency department (ED) visits for
13	as thma and respiratory disease that were available in the 2006 O_3 AQCD mainly looked
14	at the daily time metric. Collectively the short-term O ₃ studies presented earlier in
15	Section 6.2.7.5 indicate that there is evidence for increases in both hospital admissions
16	and ED visits related to both all respiratory outcomes and asthma with stronger
17	associations in the warm months. New studies evaluated long-term O3 exposure metrics,
18	providing a new line of evidence that suggests a positive exposure-response relationship
19	between first asthma hospital admission and long-term O3 exposure.
20	An ecologic study (Moore et al., 2008) evaluated time trends in associations between
21	declining warm-season O_3 concentrations and hospitalization for asthma in children in
22	California's South Coast Air Basin who ranged in age from birth to 19 years. Quarterly
23	average concentrations from 195 spatial grids, 10×10 km, were used. Ozone was the only
24	pollutant associated with increased hospital admissions over the study period. A linear
25	relation was observed for asthma hospital discharges (Moore et al., 2008). A matched
26	case-control study (Karr et al., 2007) was conducted of infant bronchiolitis (ICD 9, code
27	466.1) hospitalization and two measures of long-term pollutant exposure (the month prior
28	to hospitalization and the lifetime average) for O_3 in the South Coast Air Basin of
29	southern California among 18,595 infants born between 1995 and 2000. Ozone was
30	associated with reduced risk in the single-pollutant model, but this relation did not persist
31	in multi-pollutant models (CO, NO ₂ , and PM _{2.5}).
32	In a cross-sectional study, Meng et al. (2010) examined associations between air
33	pollution and asthma morbidity in the San Joaquin Valley in California by using the 2001
34	California Health Interview Survey data from subjects ages 1 to 65+ who reported

1	
1	physician-diagnosed asthma ($n = 1,502$). Subjects were assigned annual average
2	concentrations for O_3 based on residential ZIP code and the closet air monitoring station
3	within 8 km but did not have data on duration of residence. Multi-pollutant models for O_3
4	and PM did not differ substantially from single-pollutant estimates, indicating that
5	pollutant multi-collinearity is not a problem in these analyses. The authors reported
6	increased asthma-related ED visits or hospitalizations for O_3 (OR = 1.49; [95% CI: 1.05,
7	2.11] per 10 ppb) for all ages. Positive associations were obtained for symptoms, but
8	95% confidence intervals included null values. Associations for symptoms for adults
9	(ages 18 +) were observed (OR = 1.40; [95% CI: 1.02, 1.91] per 10 ppb).
10	Associations between air pollution and poorly controlled asthma among adults in
11	Los Angeles and San Diego Counties were investigated using the California Health
12	Interview Survey data collected between November 2000 and September 2001 (Meng et
13	al., 2007). Each respondent was assigned an annual average concentration measured at
14	the nearest station within 5 miles of the residential cross-street intersection. Poorly
15	controlled asthma was defined as having daily or weekly asthma symptoms or at least one
16	ED visit or hospitalization because of asthma during the past 12 months. This cross-
17	sectional study reports an OR of 3.34 (95% CI: 1.01, 11.09) for poorly controlled asthma
18	when comparing those 65 years of age and older above the 90th percentile (28.7 ppb)
19	level to those below that level. Co-pollutant (PM) analysis produced similar results.
20	Evidence associating long-term O3 exposure to first asthma hospital admission in a
20 21	Evidence associating long-term O_3 exposure to first asthma hospital admission in a concentration-response relationship is provided in a retrospective cohort study (<u>Lin et al.</u> ,
21	concentration-response relationship is provided in a retrospective cohort study (Lin et al.,
21 22	concentration-response relationship is provided in a retrospective cohort study (<u>Lin et al.</u> , <u>2008b</u>). This study investigated the association between chronic exposure to O_3 and
21 22 23	concentration-response relationship is provided in a retrospective cohort study (<u>Lin et al.</u> , <u>2008b</u>). This study investigated the association between chronic exposure to O_3 and childhood asthma admissions (defined as a principal diagnosis of ICD9, code 493) by
21 22 23 24	concentration-response relationship is provided in a retrospective cohort study (Lin et al., 2008b). This study investigated the association between chronic exposure to O_3 and childhood asthma admissions (defined as a principal diagnosis of ICD9, code 493) by following a birth cohort of 1,204,396 eligible births born in New York State during
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21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36	concentration-response relationship is provided in a retrospective cohort study (Lin et al., 2008b). This study investigated the association between chronic exposure to O_3 and childhood asthma admissions (defined as a principal diagnosis of ICD9, code 493) by following a birth cohort of 1,204,396 eligible births born in New York State during 1995-1999 to first asthma admission or until 31 December 2000. There were 10,429 (0.87%) children admitted to the hospital for asthma between 1 and 6 years of age. The asthma hospitalization rate in New York State in 1993 was 2.87 per 1,000 (Lin et al., 1999). Three annual indicators (all 8-h max from 10:00 a.m. to 6:00 p.m.) were used to define chronic O_3 exposure: (1) mean concentration during the follow-up period (41.06 ppb); (2) mean concentration during the O_3 season (50.62 ppb); and (3) proportion of follow-up days with O_3 levels >70 ppb. In this study the authors aimed to predict the risk of having asthma admissions in a birth cohort, but the time to the first admission in children that is usually analyzed in survival models was not their primary interest. The effects of co-pollutants were assessed and controlled for using the Air Quality Index (AQI). Interaction terms were used to assess potential effect modifications. A positive association between chronic exposure to O_3 and childhood asthma hospital admissions

1	examined and differences were found for younger children (1-2 years), poor
2	neighborhoods, Medicaid/self-paid births, geographic region and others. As shown in
3	Figure 7-3, positive concentration-response relationships were observed. Asthma
4	admissions were significantly associated with increased O ₃ levels for all chronic exposure
5	indicators (ORs, 1.16-1.68). When estimating the O_3 effect using the exceedance
6	proportion, an increase was observed (OR = 1.68; [95% CI: 1.64, 1.73]) in hospital
7	admissions with an IQR (2.51%) increase in O ₃ . A proportional hazards model for the
8	New York City data was run as a sensitivity analysis and it yielded similar results
9	between asthma admissions and chronic exposure to O_3 (Cox model: HR = 1.14,
10	[95% CI: 1.124, 1.155] is similar to logistic model results: OR = 1.16 [95% CI: 1.15,
11	1.17]) (Lin, 2010). Thus, this study provides evidence associating long-term O_3 exposure
12	to first asthma hospital admission in a concentration-response relationship.
13	In considering relationships between long-term pollutant exposure and chronic disease
14	health endpoints, Künzli (2012) offers two hypotheses relevant to research on air
15	pollution and chronic disease where chronic pathologies are found with acute expressions

- 15 pollution and chronic disease where chronic pathologies are found with acute expressions 16 of the chronic disease: "H1: Exposure provides a basis for the development of the 17 underlying chronic pathology, which increases the pool of people with chronic conditions 18 prone to exacerbations; H2: Exposure triggers an acute event (or a state of frailty that 19 results in an event with a delay of a few days or weeks) among those with the disease." 20 Künzli (2012) states if associations of pollution with events are much larger in the long-21 term studies, it provides some indirect evidence in support of H1. If air pollution 22 increases the pool of subjects with the chronic pathology (H1), more acute events are 23 expected to be seen for higher exposures since events due to various causes are part of the 24 chronic disease pathway.
- 25 Künzli (2012) makes such a comparison noting larger associations with long-term NO_2 26 exposures for adult asthma hospital admissions (Andersen et al., 2012) as compared to 27 short-term NO_2 exposures for asthma hospital admissions (Peel et al., 2005). In a further 28 example, Pope (2007) makes similar conclusions comparing long-term PM mortality 29 study results to short-term PM mortality studies. The results of Lin et al. (2008b) for first 30 asthma hospital admission, presented below, show effect estimates that are larger than 31 those reported in a study of asthma hospital admissions in New York State by Silverman 32 and Ito (2010), discussed in Chapter 6 (both studies are for young children). This 33 provides some support for the hypothesis that O₃ exposure may not only have triggered 34 the events but also increased the pool of asthmatics.



Note: Adjusted for child's sex, age, birth weight, and gestational age; maternal race, ethnicity, age, education, insurance, and smoking status during pregnancy; and regional poverty level and temperature. ORs by low, medium, and high exposure are shown for New York City (NYC: low [37.3 ppb], medium [37.3-38.11 ppb], high [38.11+ ppb] and other New York State regions (Other NYS regions: low [42.58 ppb], medium [42.58-45.06 ppb], high [45.06+ ppb]) for first asthma hospital admission. Source: Lin (2010); Lin et al. (2008b)

Figure 7-3 Ozone-asthma concentration-response relationship using the mean concentration during the entire follow-up period for first asthma hospital admission.

7.2.3 Pulmonary Structure and Function

7.2.3.1 Pulmonary Structure and Function: Evidence from Epidemiology Studies

1	The definitive 8-year follow-up analysis of the first cohort of the CHS, which is
2	discussed in Section 7.2 (Gauderman et al., 2004), provided little evidence that long-term
3	exposure to ambient O_3 was associated with significant deficits in the growth rate of lung
4	function in children. A later CHS study (Islam et al., 2007) examined relationships
5	between air pollution, lung function, and new onset asthma and reported no substantial
6	differences in the effect of O_3 on lung function. Ozone concentrations from the least to
7	most polluted communities (mean annual average of 8-h avg O ₃) ranged from 30 to
8	65 ppb, as compared to the ranges observed for the other pollutants, which had 4-fold- to
9	8-fold differences in concentrations. In a more recent CHS study, Breton et al. (2011)
10	hypothesized that genetic variation in genes on the glutathione metabolic pathway may

1	influence the association between ambient air pollutant exposures and lung function
2	growth in children. They investigated whether genetic variation in glutathione genes
3	GSS, GSR, GCLC, and GCLM was associated with lung function growth in healthy
4	children using data collected on 2,106 children over an 8-year time-period as part of the
5	Children's Health Study. <u>Breton et al. (2011)</u> found that variation in the GSS locus was
6	associated with differences in risk of children for lung function growth deficits associated
8 7	with NO ₂ , PM ₁₀ , PM _{2.5} , elemental carbon, organic carbon, and O ₃ . The negative effects of
8	air pollutants were largely observed within participants who had a particular GSS
9	haplotype. The effects ranged from -124.2 to -149.1 mL for FEV ₁ , -92.9 to -126.7 mL for
10	FVC and -193.9 to -277.9 mL/sec for MMEF for all pollutants except O_3 , for which some
10	positive associations were reported: 25.9 mL for FEV_1 ; 0.1 mL for FVC, and
11	166.5 mL/sec for MMEF. Ozone was associated with larger decreases in lung function in
12	children without this haplotype, when compared to the other pollutants with values of
13	-76.6 mL for FEV ₁ , -17.2 mL for FVC, and -200.3 mL/sec for MMEF, but only the
14	association with MMEF was statistically significant.
15	association with MMEF was statistically significant.
16	As discussed in the 2006 O ₃ AQCD, a study of freshman students at the University of
17	California, Berkeley reported that lifetime exposure to O3 was associated with decreased
18	measures of small airways (<2 mm) function (FEF ₇₅ and FEF ₂₅₋₇₅) (<u>Tager et al., 2005</u>).
19	There was an interaction with the FEF_{25-75}/FVC ratio, a measure of intrinsic airway size.
20	Subjects with a large ratio (indicating an increased airway size relative to their lung
21	volume) were less likely to have decreases in FEF_{75} and FEF_{25-75} for a given estimated
22	lifetime exposure to O ₃ . Kinney and Lippmann (2000) examined 72 nonsmoking adults
23	(mean age 20 years) from the second-year class of students at the U.S. Military Academy
24	in West Point, NY, and reported results that appear to be consistent with a decline in lung
25	function that may in part be due to O_3 exposures over a period of several summer months.
26	Ihorst et al. (2004) examined 2,153 children with a median age of 7.6 years and reported
27	pulmonary function results which indicated that significantly lower FVC and FEV_1
28	increases were associated with higher O3 exposures over the medium-term of several
29	summer months, but not over several months in the winter. Semi-annual mean O_3
30	concentrations ranged from 22 to 54 ppb during the summer months and 4 to 36 ppb
31	during the winter months. Further, over the longer-term 3.5-year period Ihorst et al.
32	(2004) found that higher mean summer months O_3 levels were not associated with growth
33	rates in lung function and for FVC and FEV ₁ , in contrast to the significant medium-term
34	effects. Frischer et al. (1999) found that higher O_3 over one summer season, one winter
35	season, and greater increases from one summer to the next over a three-year period were
36	associated with smaller increases in lung function growth, indicating both medium and
37	longer-term effects.

1	
1	(Mortimer et al., 2008a, b) examined the association of prenatal and lifetime exposures to
2	air pollutants with pulmonary function and allergen sensitization in a subset of asthmatic
3	children (ages 6-11) included in the Fresno Asthmatic Children's Environment Study
4	(FACES). Monthly means of pollutant levels for the years 1989-2000 were created and
5	averaged separately across several important developmental time-periods, including: the
6	entire pregnancy, each trimester, the first 3 years of life, the first 6 years of life, and the
7	entire lifetime. In the first analysis (<u>Mortimer et al., 2008a</u>), negative effects on
8	pulmonary function were found for exposure to PM_{10} , NO_2 , and CO during key neonatal
9	and early life developmental periods. The authors did not find a negative effect of
10	exposure to O_3 within this cohort. In the second analysis (Mortimer et al., 2008b),
11	sensitization to at least one allergen was associated, in general, with higher levels of CO
12	and PM_{10} during the entire pregnancy and second trimester, and higher PM_{10} during the
13	first 2 years of life. Lower exposure to O_3 during the entire pregnancy or second trimester
14	was associated with an increased risk of allergen sensitization. Although the pollutant
15	metrics across time periods were correlated, the strongest associations with the outcomes
16	were observed for prenatal exposures. Though it may be difficult to disentangle the effect
17	of prenatal and postnatal exposures, the models from this group of studies suggest that
18	each time period of exposure may contribute independently to different dimensions of
19	school-aged children's pulmonary function. For 4 of the 8 pulmonary-function measures
20	(FVC, FEV ₁ , PEF, FEF ₂₅₋₇₅), prenatal exposures were more influential on pulmonary
21	function than early-lifetime metrics, while, in contrast, the ratio of measures (FEV_1/FVC
22	and FEF ₂₅₋₇₅ /FVC) were most influenced by postnatal exposures. When lifetime metrics
23	were considered alone, or in combination with the prenatal metrics, the lifetime measures
24	were not associated with any of the outcomes. This suggests that the timing of the O_3
25	exposure may be more important than the overall dose, and prenatal exposures are not
26	just markers for lifetime or current exposures.
27	Latzin et al. (2009) examined whether prenatal exposure to air pollution was associated
28	with lung function changes in the newborn. Tidal breathing, lung volume, ventilation
29	inhomogeneity and eNO were measured in 241 unsedated, sleeping neonates
30	(age = 5 weeks). Consistent with the previous studies, no association was found for
31	prenatal exposure to O ₃ and lung function.
32	In a cross-sectional study of adults, Qian et al. (2005) examined the association of long-
33	term exposure to O_3 and PM_{10} with pulmonary function from data of 10,240 middle-aged
34	subjects who participated in the Atherosclerosis Risk in Communities (ARIC) study in
35	four U.S. communities. A surrogate for long-term O ₃ exposure from daily data was
36	determined at the individual level. Ozone was significantly and negatively associated
37	with measures of pulmonary function.

1	To determine the extent to which long-term exposure to outdoor air pollution accelerates
2	adult decline in lung function, Forbes et al. (2009b) studied the association between
3	chronic exposure to outdoor air pollution and lung function in approximately 42,000
4	adults aged 16 and older who were representatively sampled cross-sectionally from
5	participants in the Health Survey for England (1995, 1996, 1997, and 2001). FEV $_1$ was
6	not associated with O_3 concentrations. In contrast to the results for PM_{10} , NO_2 , and SO_{2} ;
7	combining the results of all the survey years showed that a 5-ppb difference in O_3 was
8	counter-intuitively associated with a higher FEV ₁ by 22 mL.
9	In a prospective cohort study consisting of school-age, non-asthmatic children in
10	Mexico City ($n = 3,170$) who were 8 years of age at the beginning of the study, <u>Rojas-</u>
11	<u>Martinez et al. (2007</u>) evaluated the association between long-term exposure to O_3 , PM_{10}
12	and NO ₂ and lung function growth every 6 months from April 1996 through May 1999.
13	Exposure data were provided by 10 air quality monitor stations located within 2 km of
14	each child's school. Over the study period, 8-h O3 concentrations ranged from 60 ppb
15	(SD, \pm 25) in the northeast area of Mexico City to 90 ppb (SD, \pm 34) in the southwest,
16	with an overall mean of 69.8 ppb. In multi-pollutant models, an IQR increase in mean O_3
17	concentration of 11.3 ppb was associated with an annual deficit in FEV ₁ of 12 mL in girls
18	and 4 mL in boys. Single-pollutant models showed an association between ambient
19	pollutants (O_3 , PM_{10} , and NO_2) and deficits in lung function growth. While the estimates
20	from co-pollutant models were not substantially different than single pollutant models,
21	independent effects for pollutants could not be estimated accurately because the traffic-
22	related pollutants were correlated. To reduce exposure misclassification,
23	microenvironmental and personal exposure assessments were conducted in a randomly
24	selected subsample of 60 children using passive O ₃ samplers. Personal O ₃ concentrations
25	were correlated ($p < 0.05$) with the measurements obtained from the fixed-site air
26	monitoring stations.
27	In the 2006 O_3 AQCD, few studies had investigated the effect of chronic O_3 exposure on
28	pulmonary function. The strongest evidence was for medium-term effects of extended O ₃
29	exposures over several summer months on lung function (FEV ₁) in children, i.e., reduced
30	lung function growth being associated with higher ambient O ₃ levels. Longer-term
31	studies (annual), investigating the association of chronic O_3 exposure on lung function
32	(FEV ₁) such as the definitive 8-year follow-up analysis of the first cohort (Gauderman et
33	<u>al., 2004</u>) provides little evidence that long-term exposure to ambient O_3 at current levels
34	is associated with significant deficits in the growth rate of lung function in children.
35	Analyses indicated that there was no evidence that either 8-h avg O_3 (10 a.m. to 6 p.m.)
36	or 24-h avg O_3 was associated with any measure of lung function growth over a 4-year
37	(age 10 to 14 years; (<u>Gauderman et al., 2000</u>)) or 8-year (age 10 to 18 years; (<u>Gauderman</u>
38	et al., 2004)) period. However, most of the other pollutants examined (including $PM_{2.5}$,

1	NO ₂ , acid vapor, and elemental carbon) were found to be significantly associated with
2	reduced growth in lung function. In addition, there was only about a 2- to 2.5-fold
3	difference in O ₃ concentrations from the least to most polluted communities (mean
4	annual average of 8-h avg O_3 ranged from 30 to 65 ppb), versus the ranges observed for
5	the other pollutants (which had 4- to 8-fold differences in concentrations).
6	Short-term O_3 exposure studies presented in Section <u>6.2.1.2</u> provide a cumulative body of
7	epidemiologic evidence that strongly supports associations between ambient O3 exposure
8	and decrements in lung function among children. For new studies of long-term O_3
9	exposure relationship to pulmonary function, one study, where O_3 and other pollutant
10	levels were higher (90 ppb at high end of the range) than those in the CHS, observes a
11	relationship between O_3 concentration and pulmonary function declines in school-aged
12	children. Two studies of adult cohorts provide mixed results where long- term exposures
13	were at the high end of the range with levels of 49.5 ppb in one study and 27 ppb IQR in
14	the other. Toxicological studies examining monkeys have provided data for airway
15	resistance in an asthma model but this is difficult to compare to FEV_1 results. Thus there
16	is little new evidence to build upon the very limited studies of pulmonary function
17	(FEV_1) from the 2006 O_3 AQCD.

7.2.3.2 Pulmonary Structure and Function: Evidence from Toxicological Studies and Nonhuman Primate Asthma Models

18	Long-term studies in animals allow for greater insight into the potential effects of
19	prolonged exposure to O ₃ , that may not be easily measured in humans, such as structural
20	changes in the respiratory tract. As reviewed in the 1996 and 2006 O_3 AQCDs and
21	Chapter 5 of this ISA, there are both qualitative and quantitative uncertainties in the
22	extrapolation of data generated by rodent toxicology studies to the understanding of
23	health effects in humans. Despite these uncertainties, epidemiologic studies observing
24	functional changes in humans can attain biological plausibility, in conjunction with long-
25	term toxicological studies, particularly O3-inhalation studies performed in non-human
26	primates whose respiratory system most closely resembles that of the human. An
27	important series of studies have used nonhuman primates to examine the effect of O_3
28	alone or in combination with an inhaled allergen, house dust mite antigen, on
29	morphology and lung function. These animals exhibit the hallmarks of allergic asthma
30	defined for humans, including: a positive skin test for HDMA with elevated levels of IgE
31	in serum and IgE-positive cells within the tracheobronchial airway walls; impaired
32	airflow which is reversible by treatment with aerosolized albuterol; increased abundance
33	of immune cells, especially eosinophils, in airway exudates and bronchial lavage; and

1 development of nonspecific airway responsiveness (NHLBI, 2007). Hyde et al. (2006) 2 compared asthma models of rodents (mice) and the nonhuman primate model to 3 responses in humans and concluded that the unique responses to inhaled allergen shown 4 in the rhesus monkeys make it the most appropriate animal model of human asthma. 5 These studies and others have demonstrated changes in pulmonary function and airway 6 morphology in adult and infant nonhuman primates repeatedly exposed to 7 environmentally relevant concentrations of O₃ (Joad et al., 2008; Carey et al., 2007; 8 Plopper et al., 2007; Fanucchi et al., 2006; Joad et al., 2006; Evans et al., 2004; Larson et 9 al., 2004; Tran et al., 2004; Evans et al., 2003; Schelegle et al., 2003; Fanucchi et al., 10 2000; Hyde et al., 1989; Harkema et al., 1987a; Harkema et al., 1987b; Fujinaka et al., 11 1985). Many of the observations found in adult monkeys have also been noted in infant 12 rhesus monkeys, although a direct comparison of the degree of effects between adult and 13 infant monkeys has not been reported. The findings of these nonhuman primate studies 14 have also been observed in rodent studies discussed at the end of this section and 15 included in Table 7-1.

16 The initial observations in adult nonhuman primates have been expanded in a series of 17 experiments using infant rhesus monkeys repeatedly exposed to 0.5 ppm O₃ starting at 1 month of age¹ (Plopper et al., 2007). The purpose of these studies, designed by Plopper 18 19 and colleagues, was to determine if a cyclic regimen of O_3 inhalation would amplify the 20 allergic responses and structural remodeling associated with allergic sensitization and 21 inhalation in the infant rhesus monkey. In terms of pulmonary function changes, after 22 several episodic exposures of infant monkeys to O_3 , they observed a significant increase 23 in the baseline airway resistance, which was accompanied by a small increase in airway 24 responsiveness to inhaled histamine (Schelegle et al., 2003), although neither 25 measurement was statistically different from filtered air control values. Exposure of 26 animals to inhaled house dust mite antigen alone also produced small but not statistically 27 significant changes in baseline airway resistance and airway responsiveness, whereas the 28 combined exposure to both $(O_3 + antigen)$ produced statistically significant and greater 29 than additive changes in both functional measurements. This nonhuman primate evidence 30 of an O₃-induced change in airway resistance and responsiveness supports the biologic 31 plausibility of long-term exposure to O_3 contributing to the effects of asthma in children. 32 To understand which conducting airways and inflammatory mechanisms are involved in 33 O₃-induced airway hyperresponsiveness in the infant rhesus monkey, a follow-up study 34 examined airway responsiveness ex vivo in lung slices (Joad et al., 2006). Using video 35 microscopy to morphometrically evaluate the response of bronchi and respiratory

¹ <u>Schelegle et al. (2003)</u> used a two-by-two block design. Twenty-four infant rhesus monkeys (30 days old) were exposed to 11 episodes (total of 6-months exposure period) of filtered air (FA), house dust mite allergen (HDMA), O₃ (5 days each followed by 9 days of FA). Ozone was delivered for 8h/day at 0.5 ppm. Twelve of the monkeys (HDMA, and HDMA + O₃ groups) were sensitized to house dust mite allergen (HDMA, confirmed by skin testing). To evaluate the potential for recovery, the 5 months of exposure were followed by another 6 months in FA until the monkeys were reevaluated at 12 months of age.

- 1 bronchioles to methacholine, (a bronchoconstricting agent commonly used to evaluate 2 airway responsiveness in asthmatics), the investigators observed differential effects for 3 the two airway sizes. While episodic exposure to O_3 alone (0.5 ppm) had little effect on 4 ex vivo airway responsiveness in bronchi and respiratory bronchioles, exposure to dust 5 mite antigen alone produced airway hyperresponsiveness in the large bronchi, whereas 6 O_3 + antigen produced significant increases in airway hyperresponsiveness only in the 7 respiratory bronchioles. These results suggest that ozone's effect on airway 8 responsiveness occurs predominantly in the smaller bronchioles, where dosimetric 9 models indicate the dose would be higher.
- 10 The functional changes in the conducting airways of infant rhesus monkeys exposed to 11 either O_3 alone or O_3 + antigen were accompanied by a number of cellular and 12 morphological changes, including a significant 4-fold increase in eosinophils, (a cell type 13 important in allergic asthma), in the bronchoalveolar lavage of infant monkeys exposed 14 to O_3 alone. Thus, these studies demonstrate both functional and cellular changes in the 15 lung of infant monkeys after cyclic exposure to 0.5 ppm O₃. This concentration, provides 16 relevant information to understanding the potentially damaging effects of ambient O_3 17 exposure on the respiratory tract of humans. No concentration-response data, however, are available from these nonhuman primate studies. 18
- 19 In addition to these functional and cellular changes, significant structural changes in the 20 respiratory tract have been observed in infant rhesus monkeys exposed to O_3 . During 21 normal respiratory tract development, conducting airways increase in diameter and length 22 in the infant rhesus monkey. Exposure to O_3 alone (5 days of 0.5 ppm O_3 at 8 h/day, 23 followed by 9 days of filtered air exposures for 11 cycles), however, markedly affected 24 the growth pattern of distal conducting airways (Fanucchi et al., 2006). Whereas the first 25 alveolar outpocketing occurred at airway generation 13 or 14 in filtered air-control infant 26 monkeys, the most proximal alveolarized airways occurred at an average of 10 airway 27 generations in O₃-exposed monkeys. Similarly, the diameter and length of the terminal 28 and respiratory bronchioles were significantly decreased in O₃-exposed monkeys. 29 Importantly, the O₃-induced structural pathway changes persisted after recovery in 30 filtered air for 6 months after cessation of the O_3 exposures. These structural effects were 31 accompanied by significant increases in mucus goblet cell mass, alterations in smooth 32 muscle orientation in the respiratory bronchioles, epithelial nerve fiber distribution, and 33 basement membrane zone morphometry. These latter effects are noteworthy because of 34 their potential contribution to airway obstruction and airway hyperresponsiveness which 35 are central features of asthma.
- Because many cellular and biochemical factors are known to contribute to allergic
 asthma, the effect of exposure to O₃ alone or O₃ + antigen on immune system parameters

- 1 was also examined in infant rhesus monkeys. Mast cells, which contribute to asthma via 2 the release of potent proteases, were elevated in animals exposed to antigen alone but O_3 3 alone had little effect on mast cell numbers and the response of animals exposed to O_3 + 4 antigen was not different from that of animals exposed to antigen alone; thus suggesting 5 that mast cells played little role in the interaction between O_3 and antigen in this model of 6 allergic asthma (VanWinkle et al., 2010). Increases in CD4+ and CD8+ lymphocytes 7 were observed at 6 months of age in the blood and bronchoalveolar lavage fluid of infant 8 rhesus monkeys exposed to O_3 + antigen but not in monkeys exposed to either agent 9 alone (Miller et al., 2009). Activated lymphocytes (i.e., CD25+ cells) were 10 morphometrically evaluated in the airway mucosa and significantly increased in infant 11 monkeys exposed to antigen alone or O_3 + antigen. Although O_3 alone had no effect on 12 CD25+ cells, it did alter the anatomic distribution of CD25+ cells within the airways. 13 Ozone had only a small effect on these sets of immune cells and did not produce a strong 14 interaction with an inhaled allergen in this nonhuman primate model.
- 15 In addition to alterations in the immune system, nervous system interactions with 16 epithelial cells are thought to play a contributing role to airway hyperresponsiveness. A 17 critical aspect of postnatal lung development is the laying of nerve axons with specific connections serving to maintain lung homeostasis. Aberrant innervation patterns may 18 19 underlie allergic airways disease pathology and long-term decrements in airway function. 20 As noted in the 2006 O₃ AQCD, exposure of infant rhesus monkeys altered the normal 21 development of neural innervation in the epithelium of the conducting airways (Larson et 22 al., 2004). Significant mean reductions in nerve fiber density were observed in the 23 midlevel airways of animals exposed to O_3 alone (49% reduction), and O_3 + antigen (55% 24 reduction). Moreover, the morphology of nerve bundles was altered. The persistence of 25 these effects was examined after a 6-month recovery period, and although nerve 26 distribution remained atypical, there was a dramatic increase in airway nerve density 27 (hyperinnervation) (Kajekar et al., 2007). Thus, in addition to structural, immune, and 28 inflammatory effects, exposure to O_3 produces alterations in airway innervation which 29 may contribute to O_3 -induced exacerbation of asthma. Evaluation of the pathobiology of 30 airway remodeling in growing lungs of neonates using an animal model where exposure 31 to allergen generates reactive airway disease with all the hallmarks of asthma in humans 32 illustrates that exposure to O₃ and allergen early in life produces a large number of 33 disruptions of fundamental growth and differentiation processes.
- 34A number of studies in both nonhuman primates and rodents demonstrate that O335exposure can increase collagen synthesis and deposition, inducing fibrotic-like changes in36the lung (Last et al., 1994; Chang et al., 1992; Moffatt et al., 1987; Reiser et al., 1987;37Last et al., 1984). Increased collagen content is often associated with elevated abnormal38cross links that appear to be irreversible (Reiser et al., 1987). Generally changes in

- 1 collagen content have been observed in rats exposed to 0.5 ppm O_3 or higher, although 2 extracellular matrix thickening has been observed in the lungs of rats exposed to an urban 3 pattern of O_3 with daily peaks of 0.25 ppm for 38 weeks (Chang et al., 1992; Chang et al., 4 1991). A more recent study using an urban pattern of exposure to $0.5 \text{ ppm } O_3$ 5 demonstrated that O₃-induced collagen deposition in mice is dependent on the activity of 6 TGF- β (Katre et al., 2011). Sex differences have been observed with respect to increased 7 centriacinar collagen deposition and crosslinking, which was observed in female but not 8 male rats exposed to 0.5 and 1.0 ppm O₃ for 20 months (Last et al., 1994). Few other 9 long-term exposure morphological studies have presented sex differences and most only 10 evaluated males.
- 11As described in the 1996 and 2006 O3 AQCDs, perhaps the largest chronic O3 study was12an NIEHS-NTP/HEI funded rodent study conducted by multiple investigators studying a13number of different respiratory tract endpoints (Catalano et al., 1995b). Rats were
- 14 exposed to 0.12, 0.5, or 1.0 ppm O₃ for 6 h/day and 5 d/week for 20 months. The most 15 prominent changes were observed in the nasal cavity where a large fraction of O_3 is 16 absorbed. Alterations in nasal function (increased mucous flow) and structure (goblet cell 17 metaplasia) were observed at 0.5 and 1.0 ppm but not 0.12 ppm O_3 . In the lung, the 18 centriacinar region (CAR) was the anatomical site most affected by O_3 . The epithelial cell 19 lining was changed to resemble that seen in respiratory bronchioles and the interstitial 20 volume was increased. Biochemical analyses demonstrated increased collagen and 21 glycoaminoglycans, an observation that supported the structural changes. As in the nose, 22 these changes were observed only at the two highest exposure concentrations. 23 Importantly, despite these morphologic and biochemical changes after 20 months of 24 exposure, detailed pulmonary function testing revealed little to no measurable change in 25 function. Thus, minor respiratory tract changes were observed after chronic exposure to 26 O_3 up to 1.0 ppm in the F344 rat model.
- 27 It is unclear what the long-term impact of O₃-induced structural changes may be. 28 Simulated seasonal (episodic) exposure studies suggest that such exposures might have 29 cumulative impacts, and a number of studies indicate that structural changes in the 30 respiratory system are persistent or irreversible. For example, O₃-induced hyperplasia 31 was still evident in the nasal epithelia of rats 13 weeks after recovery from 0.5 ppm O₃ 32 exposure (Harkema et al., 1999). In a study of episodic exposure to 0.25 ppm O_3 , Chang 33 et al. (1992) observed no reversal of basement membrane thickening in rat lungs up to 17 34 weeks post-exposure. Thickening of the sub-basement membrane is one of the persistent 35 structural features observed in human asthmatics (NHLBI, 2007). Episodic exposure 36 $(0.25 \text{ ppm O}_3, \text{ every other month})$ of young monkeys induced equivalent morphological 37 changes compared to continuously exposed animals, even though they were exposed for 38 half the time and evaluation occurred a month after exposure ceased as opposed to

- 1 immediately (Tyler et al., 1988). Notably, episodic O_3 exposure increased total lung 2 collagen content, chest wall compliance, and inspiratory capacity, suggesting a delay in 3 lung maturation in episodically-exposed animals. These changes were in contrast to the 4 continuously exposed group, which did not differ from the air exposed group in these 5 particular parameters but did exhibit greater bronchiolitis than the episodically exposed 6 animals. In a study by Harkema and colleagues (Harkema et al., 1993, 1987b), monkeys 7 (both males and females) were acutely exposed for 8 h/day to 0.15 ppm O_3 (6 days) or 8 chronically to 0.15 ppm or 0.3 ppm O_3 (90 days). For most endpoints in the nasal cavity, 9 the observed morphologic changes and inflammation were greater in the monkeys 10 exposed for 6 days compared to 90 days, whereas in the respiratory bronchioles of the 11 same animals, there were no significant time or concentration dependent differences 12 (increased epithelial thickness and proportion of cuboidal cells) between the 6 and 90 day 13 exposure groups.
- 14Stokinger (1962) reported that chronic bronchitis, bronchiolitis, and emphysematous and15fibrotic changes develop in the lung tissues of mice, rats, hamsters, and guinea pigs16exposed 6 h/day, 5 days/week for 14.5 months to a concentration slightly above 1 ppm17O3. Rats continuously exposed for 3 to 5 months to 0.8 ppm O3 develop a disease that18resembles emphysema, and they finally die of respiratory failure (Stephens et al., 1976).19Ozone results in a greater response of fibroblasts in the lesion, thickening of the alveolar20septae, and an increase in number of alveolar macrophages in the proximal alveoli.

Table 7-1Respiratory effects in nonhuman primates and rodents resulting
from long-term ozone exposure

Study	Model	O ₃ (ppm)	Exposure Duration	Effects
Pinkerton et al. (1998); Harkema et al. (1997a); Harkema et al. (1997b); Catalano et al. (1995b); (Chang et al., 1995a); (Chang et al., 1995); Pinkerton et al. (1995); Stockstill et al. (1995); Harkema et al. (1994); Last et al. (1994); Plopper et al. (1994)	Rat, male and female, Fischer F344, 6-8 weeks old	0.12 0.5 1.0	6 h/day, 5 days/week for 20 months	Effects similar to (or a model of) early fibrotic human disease were greater in the periacinar region than in terminal bronchioles. Thickened alveolar septa observed at 0.12 ppm O ₃ . Other effects (e.g., mucous cell metaplasia in the nose, mild fibrotic response in the parenchyma, and increased collagen in CAR of females) observed at 0.5 to 1.0 ppm. Some morphometric changes (epithelial thickening and bronchiolarization) occurred after 2 or 3 months of exposure to 1.0 ppm.
<u>Herbert et al. (1996</u>)	Mice, male and female, B6C3F1, 6-7 weeks old,	0.12 0.50 1.0	6 h/day, 5 days/week for 24 and 30 months	Similar to the response of rats in the same study (see rat above). Effects were seen in the nose and centriacinar region of the lung at 0.5 and 1.0 ppm.
<u>Chang et al. (1991</u>)	Rat, male, F344, 6 weeks old	Continuous: 0.12 or 0.25	Continuous: 12 h/day for 6 weeks	Increased Type 1 and 2 epithelial volume assessed by TEM. Linear relationship observed
		Episodic/urban: baseline 0.06; peak 0.25	Simulated urban pattern; slow rise to peak 9 h/day, 5 days/week, 13 weeks	between increases in Type 1 epithelial cell volume and concentration x time product. Degree of injury not related to pattern of exposure (continuous or episodic).
<u>Chang et al. (1992</u>)	Rat, male, F344, 6 weeks old	baseline 0.06; peak 0.25	Slow rise to peak 9 h/day, 5 days/week, 13 and 78 weeks Recovery in filtered air for 6 or 17 weeks	Progressive epithelial hyperplasia, fibroblast proliferation, and interstitial matrix accumulation observed using TEM. Interstitial matrix thickening due to deposition of basement membrane and collagen fibers. Partial recovery of interstitial matrix during follow-up periods in air; but no resolution of basement membrane thickening.
<u>Barry et al. (1985); (1983)</u>	Rat, male, 1 day old or 6 weeks old	0.12 (adults only) 0.25	12 h/day for 6 weeks	Lung and alveolar development not significantly affected. Increased Type 1 and 2 epithelial cells and AM in CAR alveoli, thickened Type 1 cells with smaller volume and less surface coverage as assessed by TEM (adults and juveniles). In adults, smaller but statistically significant similar changes at 0.12 ppm, suggesting linear concentration- response relationship. No statistically significant age-related effects observed.
<u>Tyler et al. (1988</u>)	Monkey; male, <i>Macaca fascicularis</i> , 7 mo old	0.25	8 h/day, 7 days/week, Daily for 18 mo or episodically every other month for 18 mo Episodic group evaluated 1 mo postexposure	Increased collagen content, chest wall compliance, and inspiratory capacity in episodic group only. Respiratory bronchiolitis in both groups. Episodically exposed group incurred greater alterations in physiology and biochemistry and equivalent changes in morphometry even though exposed for half the time as the daily exposure group.
<u>Harkema et al. (1999</u>)	Rat, male, Fischer F344/N HSD, 10-14 weeks old	0.25 0.5	8 h/day, 7 days/week for 13 weeks	Mucous cell hyperplasia in nasal epithelium after exposure to 0.25 and 0.5 ppm O_3 ; still evident after 13 weeks recovery from 0.5 ppm O_3 exposure.
<u>Van Bree et al. (2002</u>)	Rat, male, Wistar, 7 weeks old, n = 5/group	0.4	23.5 h/day for 1, 3, 7, 28,or 56 days	Acute inflammatory response in BALF reached a maximum at day 1 and resolved within 6 days during exposure. Centriacinar region inflammatory responses throughout O_3 exposure with increased collagen and bronchiolization still present after a recovery period.
Katre et al. (2011)	Mice; male, C57BL/6, 6-8 weeks old	0.5	8 h/day, [5 days/week O_3 , and 2 days filtered air] for 5 or 10 cycles	Sustained elevation in TGF- β and PAI-1 in lung (5 or 10 cycles); elevated α -SMA and increased collagen deposition in airway walls (after 10 cycles). Collagen increase shown to depend on TGF- β .

Study	Model	O₃ (ppm)	Exposure Duration	Effects
<u>Schelegle et al. (2003);</u>	Monkey; Rhesus, 30 days old ^ª	0.5	8 h/day for 5 days, every 5 days for a total of 11 episodes	Goblet cell metaplasia, increased AHR, and increased markers of allergic asthma (e.g., eosinophilia) were observed, suggesting that episodic exposure to O_3 alters postnatal morphogenesis and epithelial differentiation and enhances the allergic effects of house dust mite allergen in the lungs of infant primates.
<u>Harkema et al. (1993, 1987b</u>)	Monkey; <i>Macaca radiata</i> , M, F 2-6 years old	0.15 0.3	8 h/day for 90 days	Significant increase in epithelial thickness in respiratory bronchioles which was accompanied by increase in cuboidal cells; nasal lesions consisted of ciliated cell necrosis and secretory cell hyperplasia; no concentration response effects
<u>Larson et al. (2004</u>)	Monkey; <i>Macaca mulatta</i> , 30 days old ^a	0.5	11 episodes of 5 days each, 8 h/day followed by 9 days of recovery	O_3 or O_3 + house dust mite antigen caused changes in density and number of airway epithelial nerves in small conducting airways. Suggests episodic O_3 alters pattern of neural innervation in epithelial compartment of developing lungs.
<u>Plopper et al. (2007</u>)	Monkey; Rhesus, 30 days old ^a	0.5	5 months of episodic exposure; 5 days O_3 followed by 9 days of filtered air, 8h/day.	Non-significant increases airway resistance and airway responsiveness with O_3 or inhaled allergen alone. Allergen + O_3 produced additive changes in both measures.
Fanucchi et al. (2006)	Monkey; male Rhesus,30 days old	0.5	5 months of episodic exposure; 5 days O ₃ followed by 9 days of filtered air, 8h/day.	Cellular changes and significant structural changes in the distal respiratory tract in infant rhesus monkeys exposed to O ₃ postnatally.
<u>Reiser et al. (1987</u>)	Monkey; male and female Cynomolgus 6-7 mo old	0.61	8 h/day for 1 year	Increased lung collagen content associated with elevated abnormal cross links that were irreversibly deposited.

^asex not reported

1	Collectively, evidence from animal studies strongly suggests that chronic O3 exposure is
2	capable of damaging the distal airways and proximal alveoli, resulting in lung tissue
3	remodeling and leading to apparent irreversible changes. Potentially, persistent
4	inflammation and interstitial remodeling play an important role in the progression and
5	development of chronic lung disease. Further discussion of the modes of action that lead
6	to O_3 -induced morphological changes can be found in Section <u>5.3.7</u> . The findings
7	reported in chronic animal studies offer insight into potential biological mechanisms for
8	the suggested association between seasonal O_3 exposure and reduced lung function
9	development in children as observed in epidemiologic studies (see Section $7.2.3$).
10	Discussion of mechanisms involved in lifestage susceptibility and developmental effects
11	can be found in Section <u>5.4.2.4</u> .

7.2.4 Pulmonary Inflammation, Injury, and Oxidative Stress

12	The 2006 O ₃ AQCD stated that the extensive human clinical and animal toxicological
13	evidence, together with the limited epidemiologic evidence available, suggests a causal
14	role for O ₃ in inflammatory responses in the airways. Short-term exposure epidemiologic

- 1studies discussed earlier in Section 6.2.3.2 show consistent associations of O3 exposure2and increased airway inflammation and oxidative stress. Further discussion of the3mechanisms underlying inflammation and oxidative stress responses can be found in4Section 5.3.3. Though the majority of recent studies focus on short-term exposures,5several epidemiologic and toxicology studies of long-term exposure add to observations6of O3-induced inflammation and injury.
- 7 Inflammatory markers and peak expiratory pulmonary function were examined in 37 8 allergic children with physician-diagnosed mild persistent asthma in a highly polluted 9 urban area in Italy and then again 7 days after relocation to a rural location with 10 significantly lower pollutant levels (Renzetti et al., 2009). The authors observed a 4-fold 11 decrease in nasal eosinophils and a statistically significant decrease in fractional exhaled 12 nitric oxide along with an improvement in lower airway function. Several pollutants were examined, including PM₁₀, NO₂, and O₃, though pollutant-specific results were not 13 14 presented. These results are consistent with studies showing that traffic-related exposures 15 are associated with increased airway inflammation and reduced lung function in children 16 with asthma and contribute to the notion that this negative influence may be rapidly 17 reversible. Exhaled NO (eNO) has been shown to be a useful biomarker for airway 18 inflammation in large population-based studies (Linn et al., 2009). Thus, while the time 19 scale of 7 days between examinations for eNO needs to be evaluated for appropriateness, 20 the results suggest that inflammatory responses are reduced when O_3 levels are decreased.
- 21Chest radiographs (CXR) of 249 children in Mexico City who were chronically exposed22to O_3 and $PM_{2.5}$ were analyzed by Calderón-Garcidueñas et al. (2006). They reported an23association between chronic exposures to O_3 and other pollutants and a significant24increase in abnormal CXR's and lung CTs suggestive of a bronchiolar, peribronchiolar,25and/or alveolar duct inflammatory process, in clinically healthy children with no risk26factors for lung disease. These CXR and CT results should be viewed with caution27because it is difficult to attribute effects to O_3 exposure.
- 28 In a cross-sectional study, Wood et al. (2009) examined the association of outdoor air 29 pollution with respiratory phenotype (PiZZ type) in alpha 1-antitrypsin deficiency 30 (α -ATD) from the U.K. α -ATD registry. This deficiency leads to exacerbated responses 31 to inflammatory stimuli. In total, 304 PiZZ subjects underwent full lung function testing 32 and quantitative high-resolution computed tomography to identify the presence and 33 severity of COPD – emphysema. Mean annual air pollution data for 2006 was matched to 34 the location of patients' houses and used in regression models to identify phenotypic 35 associations with pollution controlling for covariates. Relative trends in O₃ levels were 36 assessed to validate use of a single year's data to indicate long-term exposure and 37 validation; data showed good correlations between modeled and measured data (Stedman

1and Kent, 2008). Regression models showed that estimated higher exposure to O32exposure was associated with worse gas transfer and more severe emphysema, albeit3accounting for only a small proportion of the lung function variability. This suggests that4a gene-specific group demonstrates a long-term O3 exposure effect.

- 5 The similarities of nonhuman primates to humans make them attractive models in which 6 to study the effects of O_3 on the respiratory tract. The nasal mucous membranes, which 7 protect the more distal regions of the respiratory tract, are susceptible to injury from O_3 . 8 Carey et al. (2007) conducted a study of O_3 exposure in infant rhesus macaques, whose 9 nasal airways closely resemble that of humans. Monkeys were exposed either acutely for 10 5 days (8 h/day) to 0.5 ppm O₃, or episodically for several biweekly cycles alternating 11 5 days of 0.5 ppm O_3 with 9 days of filtered air (0 ppm O_3), designed to mimic human 12 exposure (70 days total). All monkeys acutely exposed to O_3 had moderate to marked 13 necrotizing rhinitis, with focal regions of epitheliar exfoliation, numerous infiltrating 14 neutrophils, and some eosinophils. The distribution, character, and severity of lesions in 15 episodically exposed monkeys were similar to that of acutely exposed animals. Neither 16 group exhibited the mucous cell metaplasia proximal to the lesions, observed in adult 17 monkeys exposed continuously to 0.3 ppm O_3 in another study (Harkema et al., 1987a). 18 Adult monkeys also exhibit attenuation of inflammatory responses with continued daily 19 exposure (Harkema et al., 1987a), but inflammation did not resolve over time in young 20 episodically exposed monkeys (Carey et al., 2011). Inflammation in conducting airways 21 has also been observed in rats chronically exposed to O_3 . Using an agar-based technique 22 to fill the alveoli so that only the rat bronchi are lavaged, a 90-day exposure of rats to 23 0.8 ppm O_3 (8 h/day) elicited significantly elevated pro-inflammatory eicosanoids PGE₂ 24 and 12-HETE in the conducting airway compared to filtered air-exposed rats (Schmelzer 25 et al., 2006).
- 26 Persistent inflammation and injury leading to interstitial remodeling may play an 27 important role in the progression and development of chronic lung disease. Chronic 28 airway inflammation is an important component of both asthma and COPD. The 29 epidemiological evidence supporting an association between long-term exposure to O_3 30 and inflammation or injury is limited. However, animal studies clearly demonstrate O₃-31 induced inflammation and injury, which may or may not attenuate with chronic exposure 32 depending on the model. Further discussion of how O_3 initiates inflammation can be 33 found in Section 5.3.3.

7.2.5 Allergic Responses

1 The association of air pollutants with childhood respiratory allergies was examined in the 2 U.S. using the 1999-2005 National Health Interview Survey of approximately 70,000 3 children, and ambient air pollution data from the U.S. EPA, with monitors within 20 4 miles of each child's residential block (Parker et al., 2009). The authors examined the 5 associations between the reporting of respiratory allergy or hay fever and medium-term 6 exposure to O_3 over several summer months, controlling for demographic and geographic 7 factors. Increased respiratory allergy/hay fever was associated with increased O_3 levels 8 (adjusted OR per 10 ppb = 1.20; [95% CI: 1.15, 1.26]). These associations persisted after 9 stratification by urban-rural status, inclusion of multiple pollutants (O₃, SO₂, NO₂, PM), 10 and definition of exposure by differing exposure radii; smaller samples within 5 miles of 11 monitors were remarkably similar to the primary results. No associations between the 12 other pollutants and the reporting of respiratory allergy/hay fever were apparent. 13 Ramadour et al. (2000) reported no relationship between O₃ levels and rhinitis symptoms 14 and hay fever. Hwang et al. (2006) report the prevalence of allergic rhinitis (adjusted OR 15 per 10 ppb = 1.05; [95% CI: 0.98, 1.12]) in a large cross-sectional study in Taiwan. In a 16 large cross-sectional study in France, Penard-Morand et al. (2005) reported a positive 17 relationship between lifetime allergic rhinitis and O₃ exposure in a two-pollutant model 18 with NO₂. These studies related positive outcomes of allergic response and O_3 exposure 19 but with variable strength for the effect estimates. A toxicological study reported that 20 five weeks of continuous exposure to 0.4 ppm O_3 (but not 0.1 or 0.2 ppm O_3) augmented 21 sneezing and nasal secretions in a guinea pig model of nasal allergy (lijima and 22 Kobayashi, 2004). Nasal eosinophils, which participate in allergic disease and 23 inflammation, and allergic antibody levels in serum were also elevated by exposure to 24 concentrations as low as 0.2 ppm (Iijima and Kobayashi, 2004). 25 Nasal eosinophils were observed to decrease by 4-fold in 37 atopic, mildly asthmatic 26 children 7 days after relocation from a highly polluted urban area in Italy to a rural 27 location with significantly lower pollutant levels (Renzetti et al., 2009). Inflammatory

28 and allergic effects of O_3 exposure (30 day mean) such as increased eosinophil levels 29 were observed in children in an Austrian study (Frischer et al., 2001). Episodic exposure 30 of infant rhesus monkeys to 0.5 ppm O_3 for 5 months appears to significantly increase the 31 number and proportion of eosinophils in the blood and airways (lavage) [protocol 32 described above in Section 7.2.3.1 for Fanucchi et al. (2006)] (Maniar-Hew et al., 2011). 33 These changes were not evident at 1 year of age (6 months after O_3 exposure ceased). 34 Increased eosinophils levels have also been observed after acute or prolonged exposures 35 to O_3 in adult bonnet and rhesus monkeys (Hyde et al., 1992; Eustis et al., 1981).

1	Total IgE levels were related to air pollution levels in 369 adult asthmatics in five French
2	centers using generalized estimated equations (GEE) as part of the EGEA study described
3	earlier (Rage et al., 2009a). Geostatistical models were performed on 4×4 km grids to
4	assess individual outdoor air pollution exposure that was assigned to subject's home
5	address. Ozone concentrations were positively related to total IgE levels and an increase
6	of 5 ppb of O_3 resulted in an increase of 20.4% (95% CI: 3.0, 40.7) in total IgE levels.
7	Nearly 75% of the subjects were atopic. In two-pollutant models including O_3 and $NO_{2, the}$
8	O3 effect estimate was decreased by 25% while the NO2 effect estimate was decreased by
9	57%. Associations were not sensitive to adjustment for covariates or the season of IgE
10	measurements. These cross-sectional results suggest that exposure to O ₃ may increase
11	total IgE in adult asthmatics.
12	Although very few toxicological studies of long-term exposure examining allergy are
10	

12Annough very rew toxicological studies of long-term exposure examining anergy are13available, short-term exposure studies in rodents and nonhuman primates demonstrate14allergic skewing of immune responses and enhanced IgE production. Due to the15persistent nature of these responses, the short-term toxicological evidence lends16biological plausibility to the limited epidemiologic findings of an association between17long-term O₃ exposure and allergic outcomes.

7.2.6 Host Defense

18	Short-term exposures to O_3 have been shown to cause decreases in host defenses against
19	infectious lung disease in animal models. Acute O3-induced suppression of alveolar
20	phagocytosis and immune functions observed in animals appears to be transient and
21	attenuated with continuous or repeated exposures, although chronic exposure (weeks,
22	months) has been shown to slow alveolar clearance. In an important study investigating
23	the effects of longer term O ₃ exposure on alveolobronchiolar clearance, rats were exposed
24	to an urban pattern of O_3 (continuous 0.06 ppm, 7 days/week with a slow rise to a peak of
25	0.25 ppm and subsequent decrease to 0.06 ppm over a 9 h period for 5 days/week) for
26	6 weeks and were exposed 3 days later to chrysotile asbestos, which can cause pulmonary
27	fibrosis and neoplasia (Pinkerton et al., 1989). After 30 days, the lungs of the O ₃ -exposed
28	animals had twice the number and mass of asbestos fibers as the air-exposed rats.
29	However, chronic exposures of 0.1 ppm do not cause greater effects on infectivity than
30	short exposures, due to defense parameters becoming reestablished with prolonged
31	exposures. No detrimental effects were seen with a 120-day exposure to 0.5 ppm O_3 on
32	acute lung injury from influenza virus administered immediately before O3 exposure
33	started. However, O ₃ was shown to increase the severity of postinfluenzal alveolitis and
34	lung parenchymal changes (Jakab and Bassett, 1990). A recent study by Maniar-Hew et
35	al. (2011) demonstrated that the immune system of infant rhesus monkeys episodically

1	exposed to 0.5 ppm O_3 for 5 months ¹ appeared to be altered in ways that could diminish
2	host defenses. Reduced numbers of circulating leukocytes were observed, particularly
3	polymorphonuclear leukocytes (PMNs) and lymphocytes, which were decreased in the
4	blood and airways (bronchoalveolar lavage). These changes did not persist at 1 year of
5	age (6 months postexposure); rather, increased numbers of monocytes were observed at
6	that time point. Challenge with LPS, a bacterial ligand that activates monocytes and other
7	innate immune cells, elicited lower responses in O3-exposed animals even though the
8	relevant reactive cell population was increased. This was observed in both an in vivo
9	inhalation challenge and an ex vivo challenge of peripheral blood mononuclear cells.
10	Thus a decreased ability to respond to pathogenic signals was observed six months after
11	O_3 exposure ceased, in both the lungs and periphery.

7.2.7 Respiratory Mortality

12	A limited number of epidemiologic studies have assessed the relationship between long-
13	term exposure to O_3 and mortality. The 2006 O_3 AQCD concluded that an insufficient
14	amount of evidence existed "to suggest a causal relationship between chronic O_3
15	exposure and increased risk for mortality in humans" (U.S. EPA, 2006b). Though total
16	and cardio-pulmonary mortality were considered in these studies, respiratory mortality
17	was not specifically considered. In the most recent follow-up analysis of the ACS cohort
18	(Jerrett et al., 2009), cardiopulmonary deaths were subdivided into respiratory and
19	cardiovascular, separately, as opposed to combined in the Pope et al. (2002) work. A
20	10-ppb increment in exposure to O_3 elevated the risk of death from respiratory causes and
21	this effect was robust to the inclusion of $PM_{2.5}$. The association between increased O_3
22	concentrations and increased risk of death from respiratory causes was insensitive to the
23	use of a random-effects survival model allowing for spatial clustering within the
24	metropolitan area and state of residence, and to adjustment for several ecologic variables
25	considered individually. Additionally, a recent study (Zanobetti and Schwartz, 2011)
26	observed an association between long-term exposure to O ₃ and elevated risk of mortality
27	among Medicare enrollees that had previously experienced an emergency hospital
28	admission due to COPD.

7.2.8 Summary and Causal Determination

29 30 The epidemiologic studies reviewed in the 2006 O_3 AQCD detected no associations between long-term (annual) O_3 exposures and asthma-related symptoms, asthma

¹ Exposure protocol is described above in Section <u>7.2.3.2</u> for <u>Fanucchi et al. (2006</u>).

- 1 prevalence, or allergy to common aeroallergens among children after controlling for 2 covariates. Little evidence was available to relate long-term exposure to ambient O_3 3 concentrations with deficits in the growth rate of lung function in children. Additionally, 4 limited evidence was available evaluating the relationship between long-term O_3 5 concentrations and pulmonary inflammation and other endpoints. From toxicological 6 studies, it appeared that O₃-induced inflammation tapered off during long-term 7 exposures, but that hyperplastic and fibrotic changes remained elevated and in some 8 cases even worsened after a postexposure period in clean air. Episodic exposures were 9 also known to cause more severe pulmonary morphologic changes than continuous 10 exposure (U.S. EPA, 2006b).
- 11 The recent epidemiologic evidence base consists of studies using a variety of designs and 12 analysis methods evaluating the relationship between long-term exposure to ambient O_3 13 concentrations and measures of respiratory health effects and mortality conducted by 14 different research groups in different locations. See Table 7-2 for O₃ concentrations 15 associated with selected studies. Table 7-2 is organized by longitudinal and cross-16 sectional studies both presented alphabetically. The positive results from various designs 17 and locations support a relationship between long-term exposure to ambient O₃ concentrations and respiratory health effects and mortality. 18
- 19 Earlier studies reported associations of new-onset asthma and O₃ in an adult cohort in 20 California (McDonnell et al., 1999a; Greer et al., 1993) but only in males. In the CHS 21 cohort of children in 12 Southern California communities, long-term exposure to O_3 22 concentrations was not associated with increased risk of developing asthma (McConnell 23 et al., 2010); however, greater outdoor exercise was associated with development of 24 asthma in children living in communities with higher ambient O_3 concentrations 25 (McConnell et al., 2002). Recent CHS studies examined interactions among genetic 26 variants, long-term O_3 exposure, and new onset asthma in children. These prospective 27 cohort studies are methodologically rigorous epidemiology studies, and evidence 28 indicates gene- O_3 interactions. These studies have provided data supporting decreased 29 risk of certain different genetic variants on new onset asthma (e.g., HMOX-1, ARG) that 30 is limited to children either in low (Islam et al., 2008) or high (Salam et al., 2009) O₃ 31 communities. Gene-environment interaction also was demonstrated with findings that 32 greater outdoor exercise increased risk of asthma in GSTP1 Ile/Ile children living in high 33 O_3 communities (Islam et al., 2009). Biological plausibility for these these gene- O_3 34 environment interactions is provided by evidence that these enzymes have antioxidant 35 and/or anti-inflammatory activity and participate in well recognized modes of action in 36 asthma pathogenesis. As O_3 is a source of oxidants in the airways, oxidative stress serves 37 as the link among O_3 exposure, enzyme activity, and asthma.

Table 7-2Summary of selected key new studies examining annual ozone
exposure and respiratory health effects

Study; Health Effect; Location	Annual Mean O ₃ Concentration (ppb)	O₃ Range (ppb) Percentiles
Longitudinal		
<u>Islam et al. (2008);</u> New-onset asthma; CHS	55.2 high vs. 38.4 low communities 10:00 a.m. to 6:00 p.m. average	See left
<u>Islam et al. (2009);</u> New-onset asthma; CHS	55.2 high vs. 38.4 low communities 10:00 a.m. to 6:00 p.m.	See left
Lin et al. (2008b); First asthma hospital admission;	Range of mean O_3 concentrations over the 10 New York Regions 37.51 to 47.78	See left
New York State - 10 regions	8-h max 10:00 a.m. to 6:00 p.m.	
<u>Salam et al. (2009);</u> Childhood onset asthma; CHS	$O_{\rm 3}$ greater than or less than 50 ppb	See left
Cross-sectional		
<u>Akinbami et al. (2010</u>); Current asthma United States	12 month median 39.8 8hr max	IQR 35.9 to 43.7
<u>Hwang et al. (2005</u>); Prevalence of asthma; Taiwan	Mean 23.14	Range 18.65 to 31.17
Jacquemin et al. (In Press);	Median 46.9 ppb;	25th-75th
Asthma control in adults; Five French cities	8-h average	41-52
. <u>Lee et al. (2009b</u>); Bronchitic symptoms in asthmatic children; CHS	Above and below 50 ppb	See left
Meng et al. (2010);	Median 30.3 ppb	25-75% range
Asthma ED visits or hospitalizations; San Joaquin Valley, CA	Yearly based on hourly	27.1 to 34.0
<u>Moore et al. (2008);</u>	Median 87.8 ppb	Range
Asthma hospital admissions; South Coast Basin	Quarterly 1hr daily max	28.6 to 199.9
Rage et al. (2009a);	Mean 30 ppb	25th-75th
Asthma severity; Five French cities	8-h average	21-36
<u>Wenten et al. (2009);</u>	Median 46.9 ppb;	Min-Max
Respiratory school absence, U.S.	10a.m. – 6 p.m. average	27.6-65.3

1Studies using a cross-sectional design provide support for a relationship between long-2term O3 exposure and health effects in asthmatics. A long-term O3 exposure study relates3bronchitic symptoms to TNF-308 genotype asthmatic children with ambient O3 exposure4in the CHS (Lee et al., 2009b). A study in five French cities reports effects on asthma5severity related to long-term O3 exposure (Rage et al., 2009b). A follow-up study of this

1	cohort (<u>Jacquemin et al., In Press</u>) supports an effect of cumulative long-term O ₃
2	exposure on asthma control in adulthood in subjects with pre-existing asthma. Akinbami
3	et al. (2010) and Hwang et al. (2005) provide further evidence relating O_3 exposures and
4	the risk of asthma. For the respiratory health of a cohort based on the general U.S.
5	population, risk of respiratory-related school absences was elevated for children with the
6	CAT and MPO variant genes related to communities with high ambient O_3 levels
7	(<u>Wenten et al., 2009</u>).
8	Long-term O3 exposure was related to first childhood asthma hospital admissions in a
9	positive concentration-response relationship in a New York State birth cohort (Lin et al.,
10	2008b). A separate hospitalization cross-sectional study in San Joaquin Valley, California
11	reports similar findings (Meng et al., 2010). Another study relates asthma hospital
12	admissions to quarterly average O3 in the South Coast Air Basin of California (Moore et
13	<u>al., 2008</u>).
14	Information from toxicological studies indicates that long term exposure to O_3 during
15	gestation or development can result in irreversible morphological changes in the lung,
16	which in turn can influence the function of the respiratory tract. Studies by Plopper and
17	colleagues using an allergic asthma model have demonstrated changes in pulmonary
18	function and airway morphology in adult and infant nonhuman primates repeatedly
19	exposed to environmentally relevant concentrations of O ₃ (Fanucchi et al., 2006; Joad et
20	al., 2006; Schelegle et al., 2003; Harkema et al., 1987b). This nonhuman primate
21	evidence of an O ₃ -induced change in airway responsiveness supports the biologic
22	plausibility of long term exposure to O ₃ contributing to effects of asthma in children.
23	Results from epidemiologic studies examining long-term O ₃ exposure and pulmonary
24	function effects are inconclusive with some new studies relating effects at higher
25	exposure levels. The definitive 8-year follow-up analysis of the first cohort of the CHS,
26	which is discussed in Section 7.2 (Gauderman et al., 2004), provided little evidence that
27	long-term exposure to ambient O ₃ was associated with significant deficits in the growth
28	rate of lung function in children. Other cross-sectional studies provide mixed results.
29	Several studies (see <u>Table 7-3</u>) provide results adjusted for potential confounders,
30	presenting results for both O_3 and PM (single and multipollutant models) as well as other
31	pollutants where PM effects were not provided. As shown in the table, O3 associations
32	are generally robust to adjustment for potential confounding by PM.

Table 7-3Studies providing evidence concerning potential confounding by
PM for available endpoints.

Study Endpoint	Exposure	Single Pollutant O ₃	Single Pollutant PM	O₃ with PM	PM with O ₃
Asthma Related Health I	Effect Endpoint				
Akinbami et al. (2010) Asthma prevalence in children	IQR 35.9-43.7 ppb	1.56 (1.15, 2.10)	PM _{2.5} 1.43 (0.98, 2.10)	Adjusted for SO ₂ ,PM _{2.5} ,PM ₁₀ 1.86 (1.02-3.40)	PM _{2.5} 1.24 (0.70-2.21)
				Adjusted for PM _{2.5} , PM ₁₀ 1.36 (0.91-2.02)	PM _{2.5} 1.26 0.80-1.98)
<u>Hwang et al. (2005</u>) Asthma risk in children	10 ppb O ₃	1.138 (1.001, 1.293	0.934 (0.909, 0.960)	PM ₁₀ 1.253 (1.089, 1.442)	0.925 (0.899, 0.952)
<u>Jacquemin et al. (In</u> <u>Press</u>) Asthma control in adults	IQR 25-38 ppb O ₃ summer	1.69 (1.22, 2.34)	1.33 (1.06, 1.67)	PM ₁₀ 1.50 (1.07, 2.11)	1.28 (1.06, 1.55)
Lee et al. (2009b) Bronchitic symptoms asthmatics	High O ₃ >50 ppb	1.42 (0.75, 2.70)	NA	No substantial differences PM ₁₀ , PM _{2.5}	NA
Lin et al. (2008b) Asthma admissions in children	IQR 2.5%	1.16 (1.15, 1.17)	NA	Air Quality Index 1.24 (1.23, 1.25)	NA
Meng et al. (2007) Asthma control	1 ppm	1.70 (0.91, 3.18)	PM ₁₀ 2.06 (1.17, 3.61) women	Did not differ	NA
<u>Meng et al. (2010)</u> Asthma ED visits, Hospitalization	10 ppb	1.49 (1.05, 2.11)	PM ₁₀ 1.29 (0.99, 1.69)	Did not differ	NA
Rage et al. (2009b) Asthma severity in adults	IQR 28.5-33.9 ppb	2.53 (1.69, 3.79)	NA	No PM data Three pollutant (O_3, NO_2, SO_2) 2.74 (1.68, 4.48)	NA
Other Respiratory Health	n Effect Endpoints				
<u>Karr et al. (2007</u>) Bronchiolitis Hospitalization	10 ppb	0.92 (0.88, 0.96)	1.09 (1.04, 1.14)	PM _{2.5} 1.02 (0.94, 1.10)	1.09 (1.03, 1.15)
Parker et al. (2009) Respiratory allergy	10 ррb	1.24 (1.15, 1.34)	1.23 (1.04, 1.46)	Multi-pollutant 1.18 (1.09, 1.27)	1.29 (1.07, 1.56)
Rojas-Martinez et al. (2007) FEV ₁ (mL) Deficit Girls	11.3 ppb IQR	-24 (-30, -19)	PM ₁₀ IQR 36.4 ug/m ³ -29(-36, -21)	-17 (-23, -12)	-24 (-31,-16)

The highest quartile is shown for all results

NA = not available

1	There is limited evidence for an association between long-term exposure to ambient O_3
2	concentrations and respiratory mortality (Jerrett et al., 2009) and this effect was robust to
3	the inclusion of $PM_{2.5}$. The association between increased O_3 concentrations and
4	increased risk of death from respiratory causes was insensitive to a number of different
5	model specifications. Additionally, there is evidence that long-term exposure to O_3 is
6	associated with mortality among individuals that had previously experienced an
7	emergency hospital admission due to COPD (Zanobetti and Schwartz, 2011).
8	Taken together, the recent epidemiologic studies of respiratory health effects (including
9	respiratory symptoms, new-onset asthma and respiratory mortality) combined with
10	toxicological studies in rodents and nonhuman primates, provide biologically plausible
11	evidence that there is likely to be a causal relationship between long-term exposure
12	to O_3 and respiratory effects. The strongest epidemiologic evidence for a relationship
13	between long-term O3 exposure and respiratory effects is provided by studies that
14	demonstrate interactions between exercise or different genetic variants and long-term
15	measures of O_3 exposure on new-onset asthma in children; and increased respiratory
16	symptom effects in asthmatics. Additional studies of respiratory health effects and a
17	study of respiratory mortality provide a collective body of evidence supporting these
18	relationships. Studies considering other pollutants provide data suggesting that the effects
19	related to O_3 are independent from potential effects of the other pollutants. Some studies
20	provide evidence for a positive concentration-response relationship. Short-term studies
21	provide supportive evidence with increases in respiratory symptoms and asthma
22	medication use, hospital admissions and ED visits for all respiratory outcomes and
23	asthma, and decrements in lung function in children. The recent epidemiologic and
24	toxicological data base provides a compelling case to support the hypothesis that a
25	relationship exists between long-term exposure to ambient O_3 and measures of
26	respiratory health effects.

7.3 Cardiovascular Effects

7.3.1 Cardiovascular Disease

7.3.1.1 Cardiovascular Epidemiology

27	Long-term exposure to O ₃ and its effects on cardiovascular morbidity were not
28	considered in the 2006 O ₃ AQCD. However, recent studies have assessed the chronic
29	effects of O ₃ concentration on cardiovascular morbidity (<u>Chuang et al., 2011; Forbes et</u>

1	al., 2009a; Chen et al., 2007a). The association between O_3 concentration and markers of
2	lipid peroxidation and antioxidant capacity was examined among 120 nonsmoking
3	healthy college students, aged 18-22 years, from the University of California, Berkeley
4	(February—June 2002) (Chen et al., 2007a). By design, students were chosen from
5	geographic areas so they had experienced different concentrations of O ₃ over their
6	lifetimes and during recent summer vacation in either greater Los Angeles (LA) or the
7	San Francisco Bay Area (SF). A marker of lipid peroxidation, 8-isoprostane (8-iso-PGF)
8	in plasma, was assessed. This marker is formed continuously under normal physiological
9	conditions but has been found at elevated concentrations in response to environmental
10	exposures. A marker of overall antioxidant capacity, ferric reducing ability of plasma
11	(FRAP), was also measured. The lifetime average O_3 concentration estimates (from
12	estimated monthly averages) did not show much overlap between the two geographic
13	areas [median (range): LA, 42.9 ppb (28.5-65.3); SF, 26.9 ppb (17.6-33.5)]. Estimated
14	lifetime average O ₃ concentration was related to 8-iso-PGF [$\beta = 0.025$ (pg/mL)/8-h ppb
15	O_3 , p = 0.0007]. For the 17-ppb lifetime O_3 concentration difference between LA and SF
16	participants, there was a 17.41-pg/mL (95% CI: 15.43, 19.39) increase in 8-iso-PGF. No
17	evidence of association was observed between lifetime O ₃ concentration and FRAP
18	$[\beta = -2.21 \text{ (pg/mL)/8-h ppb O}_3, p = 0.45]$. The authors note that O ₃ was highly correlated
19	with $PM_{10-2.5}$ and NO_2 in this study population; however, their inclusion in the O_3 models
20	did not substantially modify the magnitude of the associations with O ₃ . Because the
21	average lifetime concentration results were supported by shorter-term exposure period
22	results from analyses considering O_3 concentrations up to 30 days prior to sampling, the
23	authors conclude that persistent exposure to O ₃ can lead to sustained oxidative stress and
24	increased lipid peroxidation. However, because there was not much overlap in average
25	lifetime O ₃ concentration estimates between LA and SF, it is possible that the risk
26	estimates involving the lifetime O ₃ exposures could be confounded by unmeasured
27	factors related to other differences between the two cities.
28	Forbes et al. (2009a) used the annual average exposures to assess the relationship
29	between chronic ambient air pollution and levels of fibrinogen and C-reactive protein
30	(CRP) in a cross-sectional study conducted in England. Data were collected from the
31	Health Survey of England for 1994, 1998, and 2003. The sampling strategy was designed
32	to obtain a representative sample of the English population; however, due to small group
33	sizes, only data from white ethnic groups were analyzed. For analyses, the annual
34	concentrations of O_3 were averaged for the year of data collection and the previous year
35	with the exception of 1994 (because pollutant data were not available for 1993). Median
36	O_3 concentrations were 26.7 ppb, 25.4 ppb, and 28 ppb for 1994, 1998, and 2003,
37	respectively. Year specific adjusted effect estimates were created and combined in a
38	meta-analysis. No evidence of association was observed for O_3 and levels of fibrinogen
39	or CRP (e.g., the combined estimates for the percent change in fibrinogen and CRP for a

1	10 ppb increase in O_3 were -0.28 [95% CI: -2.43, 1.92] and -3.05 [95% CI: -16.10,
2	12.02], respectively).
3	A study was performed in Taiwan to examine the association between long-term O_3
4	concentrations and blood pressure and blood markers using the Social Environment and
5	Biomarkers of Aging Study (SEBAS) (Chuang et al., 2011). Individuals included in the
6	study were 54 years of age and older. The mean annual O_3 concentration during the study
7	period was 22.95 ppb (SD 6.76 ppb). Positive associations were observed between O_3
8	concentrations and both systolic and diastolic blood pressure [changes in systolic and
9	diastolic blood pressure were 21.51mmHg (95% CI: 16.90, 26.13) and 20.56 mmHg
10	(95% CI: 18.14, 22.97) per 8.95 ppb increase in O_3 , respectively). Increased O_3
11	concentrations were also associated with increased levels of total cholesterol, fasting
12	glucose, hemoglobin A1c, and neutrophils. No associations were observed between O_3
13	concentrations and triglyceride and IL-6 levels. The observed associations were reduced
14	when other pollutants were added to the models. Further research will be important for
15	understanding the effects, if any, of chronic O ₃ exposure on cardiovascular morbidity
16	risk.

7.3.1.2 Cardiovascular Toxicology

17	
17	Three new studies have investigated the cardiovascular effects of long-term exposure to
18	O_3 in animal models (See <u>Table 7-3</u> for study details). In addition to the short-term
19	exposure effects described in Section $6.3.3$, a recent study found that O_3 exposure in
20	genetically hyperlipidemic mice enhanced aortic atherosclerotic lesion area compared to
21	air exposed controls (Chuang et al., 2009). Chuang et al. (2009) not only provided
22	evidence for increased atherogenesis in susceptible mice, but also reported an elevated
23	vascular inflammatory and redox state in wild-type mice and infant primates
24	(Section $6.3.3$). This study is compelling in that it identifies biochemical and cellular
25	events responsible for transducing the airway epithelial reactions of O_3 into
26	proinflammatory responses that are apparent in the extrapulmonary vasculature (Cole and
27	<u>Freeman, 2009</u>).
•	
28	Another recent study provides further evidence for increased vascular inflammation and
29	oxidation and long term effects in the extrapulmonary space. Rats episodically exposed to
30	O ₃ for 16 weeks presented marked increases in gene expression of biomarkers of
31	oxidative stress, thrombosis, vasoconstriction, and proteolysis (Kodavanti et al., 2011).
32	Ozone exposure upregulated aortic mRNA expression of heme oxygenase-1 (HO-1),
33	tissue plasminogen activator (tPA), plasminogen activator inhibitor-1 (PAI-1), von
34	Willebrand factor (vWf), thrombomodulin, endothelial nitric oxide synthase (eNOS),

1	endothelin-1 (ET-1), matrix metalloprotease-2 (MMP-2), matrix metalloprotease-3
2	(MMP-3), and tissue inhibitor of matrix metalloprotease-2 (TIMP-2). In addition, O_3
3	exposure depleted some cardiac mitochondrial phospholipid fatty acids (C16:0 and
4	C18:1), which may be the result of oxidative modifications. The authors speculate that
5	oxidatively modified lipids and proteins produced in the lung and heart promote vascular
6	pathology through activation of lectin-like oxidized-low density lipoprotein receptor-1
7	(LOX-1). Activated LOX-1 induces expression of a number of the biomarkers induced by
8	O ₃ exposure and is considered pro-atherogenic. Both LOX-1 mRNA and protein were
9	increased in mouse aorta after O ₃ exposure. This study provides a possible pathway and
10	further support to the observed O_3 induced atherosclerosis.
11	Vascular occlusion resulting from atherosclerosis can block blood flow through vessels
12	causing ischemia. The restoration of blood flow or reperfusion can cause injury to the
13	tissue from subsequent inflammation and oxidative damage. Ozone exposure enhanced
14	the sensitivity to myocardial ischemia-reperfusion (I/R) injury in rats while increasing
15	oxidative stress levels and pro-inflammatory mediators and decreasing production of
16	anti-inflammatory proteins (Perepu et al., 2010). Both long- and short-term O_3 exposure
17	decreased the left ventricular developed pressure, rate of change of pressure
18	development, and rate of change of pressure decay and increased left ventricular end
19	diastolic pressure in isolated perfused hearts (Section $6.3.3$ for short-term exposure
20	discussion). In this ex vivo heart model, O_3 induced oxidative stress by decreasing SOD
21	enzyme activity and increasing malondialdehyde levels. Ozone also elicited a
22	proinflammatory state evident by an increase in TNF- α and a decrease in the
23	anti-inflammatory cytokine IL-10. The authors conclude that O ₃ exposure will result in a
24	greater I/R injury.
25	Overall, the few animal studies that have been conducted suggest that long-term O_3
26	exposure may result in cardiovascular effects. These studies demonstrate O3-induced
27	atherosclerosis and injury. In addition, evidence is presented for a potential mechanism
28	for the development of vascular pathology that involves increased oxidative stress and
29	proinflammatory mediators, activation of LOX-1 by O_3 oxidized lipids and proteins, and

Further discussion of the mechanisms that may lead to cardiovascular effects from O₃
exposure can be found in Section <u>5.3.8</u>.

30

upregulation of genes responsible for proteolysis, thrombosis, and vasoconstriction.

Study	Model	O₃ (ppm)	Exposure Duration	Effects
Chuang et al. (2009)	Mice; ApoE ^{-/-} ; M; 6 weeks	0.5	8 wks, 5 days/week, 8 h/day	Enhanced aortic atherosclerotic lesion area compared to air controls.
<u>Kodavanti et al.</u> (2011)	Rat; Wistar; M; 10-12 weeks	0.4	16 wks, 1 day/week, 5 h/day	Increased vascular inflammation and oxidative stress, possibly through activation of LOX-1 signaling.
<u>Perepu et al. (2010</u>)	Rat; Sprague-Dawley; Weight: 50-75 g	0.8	56 days, 8 h/day	Enhanced the sensitivity to myocardial I/R injury while increasing oxidative stress and pro-inflammatory mediators and decreasing production of anti-inflammatory proteins.

Table 7-3Characterization of Study Details for Section 7.3.1.2.

No previous studies investigated cardiovascular effects from long-term exposure to O_3 . For details, see Section <u>7.3.1.2</u>

7.3.2 Cardiovascular Mortality

1	A limited number of epidemiologic studies have assessed the relationship between long-
2	term exposure to O_3 and mortality. The 2006 O_3 AQCD concluded that an insufficient
3	amount of evidence existed "to suggest a causal relationship between chronic O_3
4	exposure and increased risk for mortality in humans" (U.S. EPA, 2006b). Though total
5	and cardio-pulmonary mortality were considered in these studies, cardiovascular
6	mortality was not specifically considered. In the most recent follow-up analysis of the
7	ACS cohort (Jerrett et al., 2009), cardiopulmonary deaths were subdivided into
8	respiratory and cardiovascular, separately, as opposed to combined in the Pope et al.
9	(2002) work. A 10-ppb increment in exposure to O_3 elevated the risk of death from the
10	cardiopulmonary, cardiovascular, and ischemic heart disease. Inclusion of $PM_{2.5}$ as a
11	copollutant attenuated the association with exposure to O_3 for all of the cardiovascular
12	endpoints to become null. Additionally, a recent study (Zanobetti and Schwartz, 2011)
13	observed an association between long-term exposure to O ₃ and elevated risk of mortality
14	among Medicare enrollees that had previously experienced an emergency hospital
15	admission due to congestive heart failure (CHF) or myocardial infarction (MI).

7.3.3 Summary and Causal Determination

16	Previous AQCDs did not address the cardiovascular effects of long-term O ₃ exposure due
17	to limited data availability. The evidence remains limited; however the emerging data is
18	supportive of a role for O_3 in chronic cardiovascular diseases. Few epidemiologic studies
19	have investigated cardiovascular morbidity after long-term O ₃ exposure, and the majority
20	only assessed cardiovascular disease related biomarkers. The studies used annual or

1	multi-year averages of air monitoring data for exposure assessment. As described in
2	Section 4.6, this exposure assignment method is typical of long-term epidemiologic
3	studies, and analyses suggest that annual average concentrations are representative of
4	exposure metrics accounting for residential mobility. A study on O ₃ and cardiovascular
5	mortality reported no association after adjustment for PM _{2.5} levels. Further epidemiologic
6	studies on cardiovascular morbidity and mortality after long-term exposure have not been
7	published.
8	Toxicological evidence on long-term O ₃ exposure is also limited but three strong
9	toxicological studies have been published since the previous AQCD. These studies
10	provide evidence for O_3 enhanced atherosclerosis and I/R injury, corresponding with
11	development of a systemic oxidative, proinflammatory environment. Further discussion
12	of the mechanisms that may lead to cardiovascular effects can be found in Section $5.3.8$.
13	Although questions exist for how O_3 inhalation causes systemic effects, a recent study
14	proposes a mechanism for development of vascular pathology that involves activation of
15	LOX-1 by O_3 oxidized lipids and proteins. This activation may also be responsible for O_3
16	induced changes in genes involved in proteolysis, thrombosis, and vasoconstriction.
17	Taking into consideration the findings of toxicological studies, and the emerging
18	evidence from epidemiologic studies, the generally limited body of evidence is
19	suggestive of a causal relationship between long-term exposures to O_3 and
20	cardiovascular effects.

7.4 Reproductive and Developmental Effects

21 Although the body of literature characterizing the health effects associated with exposure 22 to O_3 is large and continues to grow, the research focusing on adverse birth outcomes is 23 relatively small. Among these studies, various measures of birth weight and fetal growth, 24 such as low birth weight (LBW), small for gestational age (SGA), and intrauterine 25 growth restriction (IUGR), and preterm birth (<37-week gestation; [PTB]) have received 26 more attention in air pollution research, while congenital malformations are less studied. 27 There are also recent studies on reproductive and developmental effects and infant 28 mortality. 29 A major issue in studying environmental exposures and reproductive and developmental 30 effects (including infant mortality) is selecting the relevant exposure period, since the 31 biological mechanisms leading to these outcomes and the critical periods of exposure are 32 poorly understood. To account for this, many epidemiologic studies evaluate multiple 33 exposure periods, including long-term (months to years) exposure periods, such as entire 34 pregnancy, individual trimesters or months of pregnancy, and short-term (days to weeks)

- 1 exposure periods such as the days and weeks immediately preceding birth. Due to the 2 length of gestation in rodents (18-24 days, on average), animal toxicological studies 3 investigating the effects of O_3 generally utilize short-term exposure periods. Thus, an 4 epidemiologic study that uses the entire pregnancy as the exposure period is considered 5 to have a long-term exposure period (about 40 weeks, on average), while a toxicological 6 study conducted with rats that also uses the entire pregnancy as the exposure period is 7 considered to have a short-term exposure period (about 18-24 days, on average). In order 8 to characterize the weight of evidence for the effects of O₃ on reproductive and 9 developmental effects in a consistent, cohesive and integrated manner, results from both 10 short-term and long-term exposure periods are included in this section and are identified 11 accordingly in the text and tables throughout this section.
- 12 Due to the poorly understood biological mechanisms and uncertainty regarding relevant 13 exposure studies, all of the studies of reproductive and developmental outcomes, 14 including infant mortality, are evaluated in this section. Infant development processes, 15 much like fetal development processes, may be particularly sensitive to O_3 -induced 16 health effects. Exposures proximate to the death may be most relevant if exposure causes 17 an acute effect. However, exposure occurring in early life might affect critical growth and 18 development, with results observable later in the first year of life, or cumulative exposure 19 during the first year of life may be the most important determinant. In dealing with the 20 uncertainties surrounding these issues, studies have considered several exposure metrics 21 based on different periods of exposure, including both short- and long-term exposure 22 periods. In the toxicological literature, a challenge in interpreting data from studies that 23 use very young murine pups, is that pups can have differential exposure to O_3 doses, 24 versus their respective dams, because of the physiology and behavior associated with the 25 early postnatal period. Namely, young pups tend to nuzzle close to their mothers and are 26 often housed in cages with litter used in nest formation. Both the dam's fur and the 27 bedding can absorb and react with O_3 , decreasing the dose that a young animal might 28 receive. The reproductive and developmental studies are characterized in this chapter, as 29 they contribute to the weight of evidence for an effect of O₃ on reproductive and 30 developmental effects.
- 31 Infants and fetal development processes may be particularly at-risk for O_3 -induced health 32 effects, and although the physical mechanisms are not fully understood, several 33 hypotheses have been proposed; these include: oxidative stress, systemic inflammation, 34 vascular dysfunction and impaired immune function (Section 5.3). Study of these 35 outcomes can be difficult given the need for detailed exposure data and potential 36 residential movement of mothers during pregnancy. Air pollution epidemiologic studies 37 reviewed in the 2006 O₃ AQCD examined impacts on birth-related endpoints, including 38 intrauterine, perinatal, postneonatal, and infant deaths; premature births; intrauterine

1growth retardation; very low birth weight (weight <1,500 grams) and low birth weight</th>2(weight <2,500 grams); and birth defects. However, in the limited number of studies that</td>3investigated O₃, no associations were found between O₃ and birth outcomes, with the4possible exception of birth defects.

- 5 Several recent articles have reviewed methodological issues relating to the study of 6 outdoor air pollution and adverse birth outcomes (Chen et al., 2010a; Woodruff et al., 7 2009; Ritz and Wilhelm, 2008; Slama et al., 2008). Some of the key challenges to 8 interpretation of these study results include the difficulty in assessing exposure as most 9 studies use existing monitoring networks to estimate individual exposure to ambient air 10 pollution; the inability to control for potential confounders such as other risk factors that affect birth outcomes (e.g., smoking); evaluating the exposure window (e.g., trimester) of 11 12 importance; and limited evidence on the physiological mechanism of these effects (Ritz 13 and Wilhelm, 2008; Slama et al., 2008).
- 14 Overall, the evidence for an association between exposure to ambient O₃ and 15 reproductive and developmental outcomes is growing, yet remains relatively small. 16 Recently, an international collaboration was formed to better understand the relationships 17 between air pollution and adverse birth outcomes and to examine some of these 18 methodological issues through standardized parallel analyses in datasets from different 19 countries (Woodruff et al., 2010). Initial results from this collaboration have examined 20 PM and birth weight (Parker et al., 2011); work on O₃ has not yet been performed. 21 Although early animal studies (Kaylock et al., 1980) found that exposure to O_3 in the late 22 gestation of pregnancy in rats led to some abnormal reproductive performances for 23 neonates, to date human studies have reported inconsistent results for the association of 24 ambient O₃ concentrations and birth outcomes.

7.4.1 Effects on Sperm

25	A limited amount of research has been conducted to examine the association between air
26	pollution and male reproductive outcomes, specifically semen quality. To date, the
27	epidemiologic studies have considered various exposure durations before semen
28	collection that encompass either the entire period of spermatogenesis (i.e., 90 days) or
29	key periods of sperm development that correspond to epididymal storage, development of
30	sperm motility, and spermatogenesis. In an analysis conducted as part of the Teplice
31	Program, 18-year-old men residing in the heavily polluted district of Teplice in the Czech
32	Republic were found to be at greater risk of having abnormalities in sperm morphology
33	and chromatin integrity than men of similar age residing in Prachatice, a less polluted
34	district (Selevan et al., 2000; Sram et al., 1999). A follow-up longitudinal study

conducted on a subset of the same men from Teplice revealed associations between total episodic air pollution and abnormalities in sperm chromatin (<u>Rubes et al., 2005</u>). A limitation of these studies is that they did not identify specific pollutants or their concentrations.

- 5 More recent epidemiologic studies conducted in the U.S. have also reported associations 6 between ambient air pollution and sperm quality for individual air pollutants, including 7 O₃ and PM_{2.5}. In a repeated measures study in Los Angeles, CA, <u>Sokol et al. (2006)</u> 8 reported a reduction in average sperm concentration during three exposure windows 9 (short-term exposures of 0-9, 10-14, and 70-90 days before semen collection, as well as 10 long-term exposures of 0-90 days before semen collection) associated with high ambient 11 levels of O_3 in healthy sperm donors. This effect persisted under a joint additive model 12 for O_3 , CO, NO_2 and PM_{10} . The authors did not detect a reduction in sperm count. Hansen 13 et al. (2010) investigated the effect of exposure to O_3 and $PM_{2.5}$ (using the same exposure 14 windows used by Sokol et al. (2006) on sperm quality in three southeastern counties 15 (Wake County, NC; Shelby County, TN; Galveston County, TX). Outcomes included 16 sperm concentration and count, morphology, DNA integrity and chromatin maturity. 17 Overall, the authors found both protective and adverse effects, although some results 18 suggested adverse effects on sperm concentration, count and morphology.
- 19 The biological mechanisms linking ambient air pollution to decreased sperm quality have 20 yet to be determined, though O_3 -induced oxidative stress, inflammatory reactions, and the 21 induction of the formation of circulating toxic species have been suggested as possible 22 mechanisms (see Section 5.3.8). Decremental effects on testicular morphology have been 23 demonstrated in a toxicological study with histological evidence of O₃-induced depletion 24 of germ cells in testicular tissue and decreased seminiferous tubule epithelial layer. 25 Jedlinska-Krakowska et al. (2006) demonstrated histopathological evidence of impaired 26 spermatogenesis (round spermatids/ spermatocytes, giant spermatid cells, and focal 27 epithelial desquamation with denudation to the basement membrane). The exposure 28 protocol used five-month-old adult rats exposed to O_3 as adults (long-term exposure, 29 0.5 ppm, 5 h/day for 50 days). This degeneration could be rescued by vitamin E 30 administration, indicating an antioxidant effect. Vitamin C administration had no effect at 31 low doses of ascorbic acid and exacerbated the O₃-dependent damage at high doses, as 32 would be expected as vitamin C can be a radical generator instead of an antioxidant at 33 higher doses. In summary, this study provided toxicological evidence of impaired 34 spermatogenesis with O₃ exposure that was rescued with certain antioxidant 35 supplementation.
- 36Overall, there is limited epidemiologic evidence for an association with O3 concentration37and decreased sperm concentration. A recent toxicological study provides limited

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2

3

4

1 2 evidence for a possible biological mechanism (histopathology showing impaired spermatogenesis) for such an association.

7.4.2 Effects on Reproduction

3 Evidence suggests that exposure to air pollutants during pregnancy may be associated 4 with adverse birth outcomes, which has been attributed to the increased sensitivity of the 5 fetus due to physiologic immaturity. Gametes (i.e., ova and sperm) may be even more at-6 risk, especially outside of the human body, as occurs with assisted reproduction. Smokers 7 require twice the number of in vitro fertilization (IVF) attempts to conceive as non-8 smokers (Feichtinger et al., 1997), suggesting that a preconception exposure can be 9 harmful to pregnancy. A recent study used an established national-scale, log-normal 10 kriging method to spatially estimate daily mean concentrations of criteria pollutants at 11 addresses of women undergoing their first IVF cycle and at their IVF labs from 2000 to 12 2007 in the northeastern U.S. (Legro et al., 2010). Increasing O₃ concentration at the 13 patient's address during ovulation induction (short-term exposure, ~12 days) was 14 significantly associated with an increased chance of live birth (OR = 1.13, [95% CI: 1.05, 15 1.22] per 10 ppb increase), but with decreased odds of live birth when exposed from 16 embryo transfer to live birth (long-term exposure, ~ 200 days) (OR = 0.79, [95% CI: 0.69, 17 (0.90) per 10 ppb increase). After controlling for NO₂ in a copollutant model, however, O₃ 18 was no longer significantly associated with IVF failure. The results of this study suggest 19 that short-term exposure to O_3 during ovulation was beneficial (perhaps due to early 20 conditioning to O_3), whereas long-term exposure to O_3 (e.g., during gestation) was 21 detrimental, and reduced the likelihood of a live birth. 22 In most toxicological studies, reproductive success appears to be unaffected by O₃ 23 exposure. Nonetheless, one study has reported that 25% of the BALB/c mouse dams in 24 the highest O₃ exposure group (1.2 ppm, short-term exposure GD9-18) did not complete 25 a successful pregnancy, a significant reduction (Sharkhuu et al., 2011). Ozone 26 administration (continuous 0.4, 0.8 or 1.2 ppm O_3) to CD-1 mouse dams during the

27 majority of pregnancy (short-term exposure, PD7-17, which excludes the 28 pre-implantation period), led to no adverse effects on reproductive success (proportion of 29 successful pregnancies, litter size, sex ratio, frequency of still birth, or neonatal mortality) 30 (Bignami et al., 1994). There was a nearly statistically significant increase in pregnancy 31 duration (0.8 and 1.2 ppm O_3). Initially, dam body weight (0.8 and 1.2 ppm O_3), water 32 consumption (0.4, 0.8 and 1.2 ppm O₃) and food consumption (0.4, 0.8 and 1.2 ppm O₃) 33 during pregnancy were decreased with O_3 exposure but these deficits dissipated a week or 34 two after the initial exposure (Bignami et al., 1994). The anorexigenic effect of O₃ 35 exposure on the pregnant dam appears to dissipate with time; the dams seem to adapt to

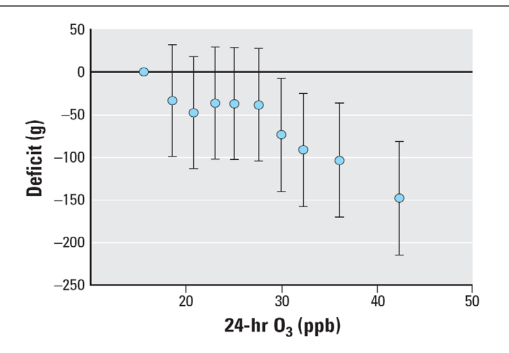
1 the O_3 exposure. In males, data exist showing morphological evidence of alte	red
2 spermatogenesis in O ₃ exposed animals (<u>Jedlinska-Krakowska et al., 2006</u>). S	ome
3 evidence suggests that O ₃ may affect reproductive success when combined w	ith other
4 chemicals. <u>Kavlock et al. (1979</u>) showed that O_3 acted synergistically with so	dium
5 salicylate to increase the rate of pup resorptions after midgestational exposure	e (1.0 ppm
6 O ₃ , short-term exposure, GD9-GD12). At low concentrations of O ₃ exposure,	
7 toxicological studies show reproductive effects to include a transient anorexi	genic effect
8 of O_3 on gestational weight gain, and a synergistic effect of O_3 on salicylate-in	nduced pup
9 resorptions; other fecundity, pregnancy- and gestation-related outcomes appe	ar
10 unaffected by O_3 exposure.	

11Collectively, there is very little epidemiologic evidence for the effect of short- or long-12term exposure to O3 on reproductive success, and the reproductive success in rats appears13to be unaffected in toxicological studies of short-term exposure to O3.

7.4.3 Birth Weight

- 14With birth weight routinely collected in vital statistics and being a powerful predictor of15infant mortality, it is the most studied outcome within air pollution-birth outcome16research. Air pollution researchers have analyzed birth weight as a continuous variable17and/or as a dichotomized variable in the form of LBW (<2,500 g [5 lbs, 8 oz]).</td>
- 18Birth weight is primarily determined by gestational age and intrauterine growth, but also19depends on maternal, placental and fetal factors as well as on environmental influences.20In both developed and developing countries, LBW is the most important predictor for21neonatal mortality and is a significant determinant of postneonatal mortality and22morbidity. Studies report that infants who are smallest at birth have a higher incidence of23diseases and disabilities, which continue into adulthood (Hack and Fanaroff, 1999).
- 24 The strongest evidence for an effect of O_3 on birth weight comes from the Children's 25 Health Study conducted in southern California. In this study, Salam et al. (2005) report 26 that maternal exposure to 24-h avg O_3 concentrations averaged over the entire pregnancy 27 was associated with reduced birth weight (39.3 g decrease [95% CI: -55.8, -22.8] in birth 28 weight per 10 ppb and 8-h avg (19.2-g decrease [95% CI: -27.7, -10.7] in birth weight per 29 10 ppb). This effect was stronger for concentrations averaged over the second and third 30 trimesters. PM_{10} , NO_2 and CO concentrations averaged over the entire pregnancy were 31 not statistically significantly associated with birth weight, although CO concentrations in 32 the first trimester and PM₁₀ concentrations in the third trimester were associated with a 33 decrease in birth weight. Additionally, the authors observed a concentration-response 34 relationship of birth weight with 24-h avg O₃ concentrations averaged over the entire

1pregnancy that was clearest above the 30-ppb level (see Figure 7-4). Relative to the2lowest decile of 24-h avg O3, estimates for the next 5 lowest deciles were approximately3-40 g to -50 g, with no clear trend and with 95% confidence bounds that included zero.4The highest four deciles of O3 exposure showed an approximately linear decrease in birth5weight, and all four 95% CIs excluded zero, and ranged from mean decreases of674 grams to decreases of 148 grams.



Note: Deficits are plotted against the decile-group-specific median O_3 exposure. Error bars represent 95% CIs. Indicator variables for each decile of O_3 exposure (except the least-exposed group) were included in a mixed model. Source: <u>Salam et al. (2005</u>).'

Figure 7-4 Birthweight deficit by decile of 24-h avg ozone concentration averaged over the entire pregnancy compared with the decile group with the lowest ozone exposure.

7	Several additional studies conducted in the U.S. and Canada also investigated the
8	association between ambient O_3 concentrations and birth weight and report some weak
9	evidence for an association. Morello-Frosch et al. (2010) estimated ambient O_3
10	concentrations throughout pregnancy and for each trimester in the neighborhoods of
11	women who delivered term singleton births between 1996 and 2006 in California. A
12	10-ppb increase in the O_3 concentration averaged across the entire pregnancy was
13	associated with a 5.7-g decrease (95% CI: -6.6, -4.9) in birth weight when exposures
14	were calculated using monitors within 10 km of the maternal address at date of birth.

1	When the distance from the monitor was restricted to 3 km, the decrease in birth weight
2	associated with a 10-ppb increase in O_3 concentration was 8.9 g (95% CI: -10.6, -7.1).
3	These results persisted in copollutant models and in models that stratified by trimester of
4	exposure, SES, and race. <u>Darrow et al. (2011b</u>) did not observe an association with birth
5	weight and O_3 concentrations during two exposure periods of interest (i.e., the first month
6	and last trimester), but did find an association with reduced birth weight when examining
7	the cumulative air pollution concentration during the entire pregnancy period.
8	Additionally, they observed effect modification by race and ethnicity, such that
9	associations between birth weight and third-trimester O_3 concentrations were
10	significantly stronger in Hispanics and non-Hispanic African Americans than in non-
11	Hispanic whites. Chen et al. (2002) used 8-h avg O_3 concentrations to create exposure
12	variables based on average maternal exposure for each trimester. Ozone was not found to
13	be related to birth weight in single-pollutant models, though the O_3 effect during the third
14	trimester was borderline statistically significant in a copollutant model with PM_{10} .
15	Several studies found no association between ambient O3 concentrations and birth
16	weight. Wilhelm and Ritz (2005) extended previous analyses of term LBW (Ritz et al.,
17	2000; <u>Ritz and Yu, 1999</u>) to include the period 1994-2000. The authors examined varying
18	residential distances from monitoring stations to see if the distance affected risk
19	estimation, exploring the possibility that effect attenuation may result from local pollutant
20	heterogeneity inadequately captured by ambient monitors. As in their previous studies,
21	the authors observed associations between elevated concentrations of CO and PM_{10} both
22	early and late in pregnancy and risk of term LBW. After adjusting for CO and/or PM_{10}
23	the authors did not observe associations between O_3 and term LBW in any of their
24	models. Brauer et al. (2008) evaluated the impacts of air pollution (CO, NO ₂ , NO, O ₃ ,
25	SO ₂ , PM _{2.5} , PM ₁₀) on birth weight for the period 1999-2002 using spatiotemporal
26	residential exposure metrics by month of pregnancy in Vancouver, BC. Quantitative
27	results were not presented for the association between O_3 and LBW, though the authors
28	observed associations that were largely protective. Dugandzic et al. (2006) examined the
29	association between LBW and ambient levels of air pollutants by trimester of exposure
30	among a cohort of term singleton births from 1988-2000. Though there was some
31	indication of an association with SO_2 and PM_{10} , there were no effects for O_3 .
32	Similarly, studies conducted in Australia, Latin America, and Asia report limited
33	evidence for an association between ambient O ₃ and measures of birth weight. In Sydney,
34	Australia, Mannes et al. (2005) found that O_3 concentrations in the second trimester of
35	pregnancy had small adverse effects on birth weight (7.5-g decrease; [95% CI: -13.8, 1.2]
36	per 10 ppb), although this effect disappeared when the analysis was limited to births with
37	a maternal address within 5 km of a monitoring station (87.7-g increase; [95% CI: 10.5,
38	164.9] per 10 ppb). Hansen et al. (2007) reported that trimester and monthly specific

1	exposures to all pollutants were not statistically significantly associated with a reduction
2	in birth weight in Brisbane, Australia. In Sao Paulo, Brazil, Gouveia et al. (2004) found
3	that O_3 exhibited a small inverse relation with birth weight over the third trimester (6.0-g
4	decrease; [95% CI: -30.8, 18.8] per 10 ppb). Lin et al. (2004b) reported a positive, though
5	not statistically significant, exposure-response relationship for O_3 during the entire
6	pregnancy in a Taiwanese study. In a study performed in Korea, Ha et al. (2001) reported
7	no O ₃ effect during the first trimester of pregnancy, but they found that during the third
8	trimester of pregnancy O_3 was associated with LBW (RR = 1.05 [95% CI: 1.02, 1.08] per
9	10 ppb).

Study	Location Sample Size	Mean O₃ (ppb)	Exposure assessment	Effect Estimate ^a (95% CI)
<u>Salam et al. (2005</u>)	California, U.S. (n = 3,901)	24-h avg: 27.3 8 h: 50.6	ZIP code level	Entire pregnancy: -39.3 g (-55.8, -22.8) T1: -6.1 g (-16.8, 4.8) T2: -20.0 g (-31.7, -8.4) T3: -20.7 g (-32.1, -9.3)
<u>Morello-Frosch et al.</u> (2010)	California, U.S. (n = 3,545,177)	24-h avg: 23.5	Nearest Monitor (within 10, 5, 3 km)	Entire pregnancy: -5.7 g (-6.6, -4.9) T1: -2.1 g (-2.9, -1.4) T2: -2.3 g (-3.1, -1.5) T3: -1.3 g (-2.1, -0.6)
Darrow et al. (2011b)	Atlanta, GA (N=406,627)	8-h max: 44.8	Population-weighted spatial average	Entire pregnancy: -12.3 g (-17.8, -6.8) First 28 days -0.5 g (-3.0, 2.1) T3: -0.9g (-4.5, 2.8)
<u>Chen et al. (2002</u>)	Northern Nevada, U.S. (n = 36,305)	8-h: 27.2	County level	Entire pregnancy: 20.9 g (6.3, 35.5) T1: 23.4 g (-35.6 , 82.4) T2: -19.4 g (-77.0, 38.2) T3: 7.7 g (-50.9, 66.3)
Wilhelm and Ritz (2005)	Los Angeles County, CA (n = 136,134)	1-h: 21.1-22.2	Varying distances from monitor	T1: NR T3: NR 6 weeks before birth: NR
<u>Brauer et al. (2008</u>)	Vancouver, BC, Canada (n = 70,249)	24-h avg: 14	Nearest Monitor (within 10 km) Inverse Distance Weighting (IDW)	Entire pregnancy: NR First 30 days of pregnancy: NR Last 30 days of pregnancy: NR T1: NR T3: NR
Dugandzic et al. (2006)	Nova Scotia, Canada (n = 74,284)	24-h avg: 21	Nearest Monitor (within 25 km)	T1: 0.97 (0.81, 1.18) ^d T2: 1.06 (0.87, 1.27) ^d T3: 1.01 (0.83-1.24) ^d

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Study	Location Sample Size	Mean O₃ (ppb)	Exposure assessment	Effect Estimate ^a (95% CI)
<u>Mannes et al. (2005)</u>	Sydney, Australia	1-h max:	Citywide avg and	T1: -0.9 g (-6.6, 4.8)
	(n = 138,056)	31.6	<5 km from monitor	T2: -7.5 g (-13.8, 1.2)
				T3: -4.5 g (-10.8, 1.8)
				Last 30 days:
				-1.1 g (-5.6, 3.4)
<u>Hansen et al. (2007)</u>	Brisbane, Australia	8 h max:	Citywide avg	T1: 2.8 g (-10.5, 16.0)
	(n = 26,617)	26.7		T2: 4.4 g (-11.4, 20.1)
				T3: 11.3 g (-4.4, 27.1)
<u>Gouveia et al. (2004)</u>	Sao Paulo, Brazil	1-h max:	Citywide avg	T1: -3.2 g (-25.6, 19)
	(n = 179,460)	31.5		T2: -0.2 g (-23.8, 23.4)
				T3: -6.0 g (-30.8, -18.8)
Lin et al. (2004b)	Kaohsiung and Taipei, Taiwan	24-h avg: 15.86- 47.78	Nearest monitor (within 3 km)	Entire pregnancy: 1.13 (0.92, 1.38) [°]
	(n = 92,288)			T1: 1.02 (0.85, 1.22) ^c
				T2: 0.93 (0.78, 1.12) ^c
				T3: 1.05 (0.87, 1.26) ^c
<u>Ha et al. (2001</u>)	Seoul, Korea	8-h avg:	Citywide avg	T1: 0.87 (0.81, 0.94) ^c
	(n = 276,763)	22.4-23.3 ^b		T3: 1.05 (1.02, 1.08) ^c

^aChange in birthweight per 10 ppb change in O_3

^bMedian

^cOdds ratios of LBW; Highest quartile of exposure compared to lowest quartile of exposure

^dRelative risk of LBW per 10 ppb change in O_3

T1 = First Trimester, T2 = Second Trimester, T3 = Third Trimester

NR: No quantitative results reported

1	Table 7-4 provides a brief overview of the epidemiologic studies of birth weight. In
2	summary, only the Children's Health Study conducted in southern California (Salam et
3	<u>al., 2005</u>) provides strong evidence for an effect of ambient O_3 on birth weight. The study
4	by Morello-Frosch et al. (2010), also conducted in California, provides support for the
5	results of the Children's Health Study. Additional studies, conducted in the U.S., Canada,
6	Australia, Latin America, and Asia, provide limited and inconsistent evidence to support
7	the effect reported in the Children's Health Study. The toxicological literature on the
8	effect of O ₃ on birth weight is sparse. In some studies, the reporting of birth weight may
9	be avoided because birth weight can be confounded by decreased litter size resulting
10	from an increased rate of pup resorption (aborted pups) in O3 exposed dams. In one
11	toxicological study by Haro and Paz (1993), no differences in litter size were observed
12	and decreased birth weight in pups from dams who were exposed to 1ppm O_3 during
13	pregnancy (short-term exposure, ~22 days) was reported. A second animal toxicology
14	study recapitulated these finding with pregnant BALB/c mice that exposed to O_3
15	(1.2 ppm, short-term exposure, GD9-18) producing pups with significantly decreased
16	birth weights (Sharkhuu et al., 2011).

7.4.4 Preterm Birth

1	Preterm birth (PTB) is a syndrome (<u>Romero et al., 2006</u>) that is characterized by multiple
2	etiologies. It is therefore unusual to be able to identify an exact cause for each PTB. In
3	addition, PTB is not an adverse outcome in itself, but an important determinant of health
4	status (i.e., neonatal morbidity and mortality). Although some overlap exists for common
5	risk factors, different etiologic entities related to distinct risk factor profiles and leading
6	to different neonatal and postneonatal complications are attributed to PTB and measures
7	of fetal growth. Although both restricted fetal growth and PTB can result in LBW,
8	prematurity does not have to result in LBW or growth restricted babies.
9	A major issue in studying environmental exposures and PTB is selecting the relevant
10	exposure period, since the biological mechanisms leading to PTB and the critical periods
11	of vulnerability are poorly understood (Bobak, 2000). Short-term exposures proximate to
12	the birth may be most relevant if exposure causes an acute effect. However, exposure
13	occurring in early gestation might affect placentation, with results observable later in
14	pregnancy, or cumulative exposure during pregnancy may be the most important
15	determinant. The studies reviewed have dealt with this issue in different ways. Many
16	have considered several exposure metrics based on different periods of exposure. Often
17	the time periods used are the first month (or first trimester) of pregnancy and the
18	last month (or 6 weeks) prior to delivery. Using a time interval prior to delivery
19	introduces an additional problem since cases and controls are not in the same stage of
20	development when they are compared. For example, a preterm infant delivered at
21	36 weeks is a 32-week fetus 4 weeks prior to birth, while an infant born at term
22	(40 weeks) is a 36-week fetus 4 weeks prior to birth.
23	Recently, investigators have examined the association of PTB with both short-term
24	(i.e., hours, days, or weeks) and long-term (i.e., months or years) exposure periods. Time-
25	series studies have been used to examine the association between air pollution
26	concentrations during the days immediately preceding birth. An advantage of these time-
27	series studies is that this approach can remove the influence of covariates that vary across
28	individuals over a short period of time. Retrospective cohort and case-control studies
29	have been used to examine long-term exposure periods, often averaging air pollution
30	concentrations over months or trimesters of pregnancy.
31	Studies of PTB fail to show consistency in pollutants and periods during pregnancy when
32	an effect occurs. For example, while some studies find the strongest effects associated
33	with exposures early in pregnancy, others report effects when the exposure is limited to
34	the second or third trimester. However, the effect of air pollutant exposure during
35	pregnancy on PTB has a biological basis. There is an expanding list of possible

1 2	mechanisms that may explain the association between O_3 exposure and PTB (see Section <u>5.4.2.4</u>).
3	Many studies of PTB compare exposure in quartiles, using the lowest quartile as the
4	reference (or control) group. No studies use a truly unexposed control group. If exposure
5	in the lowest quartile confers risk, than it may be difficult to demonstrate additional risk
6	associated with a higher quartile. Thus negative studies must be interpreted with caution.
7	Preterm birth occurs both naturally (idiopathic PTB), and as a result of medical
8	intervention (iatrogenic PTB). <u>Ritz et al. (2007</u>); (2000) excluded all births by Cesarean
9	section to limit their studies to idiopathic PTB. No other studies attempted to distinguish
10	the type of PTB, although air pollution exposure maybe associated with only one type.
11	This is a source of potential effect misclassification.
12	Generally, studies of air pollution and birth outcomes conducted in North America and
13	the United Kingdom have not identified an association between PTB and maternal
14	exposure to O_3 . Most recently, <u>Darrow et al. (2009</u>) used vital record data to construct a
15	retrospective cohort of 476,489 births occurring between 1994 and 2004 in 5 central
16	counties of metropolitan Atlanta. Using a time-series approach, the authors examined
17	aggregated daily counts of PTB in relation to ambient levels of CO, NO ₂ , SO ₂ , O ₃ , PM ₁₀ ,
18	$PM_{2.5}$ and speciated PM measurements. This study investigated 3 gestational windows of
19	short- and long-term exposure: the final week of gestation (short-term exposure), and the
20	first month of gestation and the final 6 weeks of gestation (long-term exposure). The
21	authors did not observe associations of PTB with O3 concentrations for any of the
22	exposure periods.
23	A number of U.S. studies were conducted in southern California, and report somewhat
24	inconsistent results. <u>Ritz et al. (2000</u>) evaluated the effect of air pollution (CO, NO ₂ , O ₃ ,
25	PM_{10}) exposure during pregnancy on the occurrence of PTB in a cohort of 97,518
26	neonates born in southern California between 1989 and 1993. The authors use both short-
27	and long-term exposure windows, averaging pollutant measures taken at the closest air-
28	monitoring station over distinct periods, such as 1, 2, 4, 6, 8, 12, and 26 weeks before
29	birth and the whole pregnancy period. Additionally, they calculated average exposures
30	for the first and second months of pregnancy. The authors found no consistent effects
31	associated with O_3 concentration over any of the pregnancy periods in single or
32	multipollutant models. Wilhelm and Ritz (2005) extended previous analyses of PTB (Ritz
33	et al., 2000; Ritz and Yu, 1999) in California to include 1994-2000. The authors
34	examined varying residential distances from monitoring stations to see if the distance
35	affected risk estimation, because effect attenuation may result from local pollutant
36	heterogeneity inadequately captured by ambient monitors. The authors analyzed the
37	association between long-term O_3 exposure during varying periods of pregnancy and

- 1 PTB, finding a positive association between O_3 levels in both the first trimester of 2 pregnancy (RR = 1.23 [95% CI: 1.06, 1.42] per 10 ppb increase in 24-h avg O₃) and the 3 first month of pregnancy (results for first trimester exposure were similar, but slightly 4 smaller, quantitative results not presented) in models containing all pollutants. No 5 association was observed between O_3 in the 6 weeks before birth and preterm delivery. 6 Finally, Ritz et al. (2007) conducted a case-control survey nested within a birth cohort 7 and assessed the extent to which residual confounding and exposure misclassification 8 impacted air pollution effect estimates. The authors calculated mean long-term exposure 9 levels for three gestational periods: the entire pregnancy, the first trimester, and the last 10 6 weeks before delivery. Though positive associations were observed for CO and PM_{25} , 11 no consistent patterns of increase in the odds of PTB for O₃ or NO₂ were observed.
- 12 A study conducted in Canada evaluated the impacts of air pollution (including CO, NO₂, 13 NO, O₃, SO₂, PM_{2.5}, and PM₁₀) on PTBs (1999-2002) using spatiotemporal residential exposure metrics by month of pregnancy (long-term exposure) in Vancouver, BC (Brauer 14 15 et al., 2008). The authors did not observe consistent associations with any of the 16 pregnancy average exposure metrics except for $PM_{2.5}$ for PTB. The O₃ associations were 17 largely protective, and no quantitative results were presented for O_3 . Additionally, Lee et 18 al. (2008c) used time-series techniques to investigate the associations of short-term 19 exposure to O_3 and PTB in London, England. In addition to exposure on the day of birth, 20 cumulative exposure up to 1 week before birth was investigated. The risk of PTB did not 21 increase with exposure to the levels of ambient air pollution experienced by this 22 population.
- 23 Conversely, studies conducted in Australia and China provide evidence for an association 24 between ambient O_3 and PTB. Hansen et al. (2006) reported that long-term exposure to 25 O_3 during the first trimester was associated with an increased risk of PTB (OR = 1.38, 26 [95% CI: 1.14, 1.69] per 10 ppb increase). Although the test for trend was significant due 27 to the strong effect in the highest quartile, there was not an obvious exposure-response 28 pattern across the quartiles of O₃ during the first trimester. The effect estimate was 29 diminished and lost statistical significance when PM₁₀ was included in the model 30 (OR = 1.23, [95% CI: 0.97, 1.59] per 10 ppb increase). Maternal exposure to O₃ during 31 the 90 days prior to birth showed a weak, positive association with PTB (OR = 1.09, 32 [95% CI: 0.85, 1.39] per 10 ppb increase). Jalaludin et al. (2007) found that O₃ levels in 33 the month and three months preceding birth had a statistically significant association with 34 PTB. Ozone levels in the first trimester of pregnancy were associated with increased risks 35 for PTBs (OR = 1.15 [95% CI: 1.05, 1.24] per 10 ppb increase in 1-h max O_3 36 concentration), and remained a significant predictor of PTB in copollutant models (ORs 37 between 1.07 and 1.10). Jiang et al. (2007) examined the effect of short- and long-term 38 exposure to air pollution on PTB, including risk in relation to levels of pollutants for a

1	single day exposure window with lags from 0 to 6 days before birth. An increase of
2	10 ppb of the 8-week avg of O ₃ corresponded to 9.47% (95% CI: 0.70, 18.7%) increase in
3	PTBs. Increases in PTB were also observed for PM_{10} , SO ₂ , and NO ₂ . The authors did not
4	observe any significant effect of short-term exposure to outdoor air pollution on PTB
5	among the 1-day time windows examined in the week before birth.
6	Little data is available from toxicological studies; a study reported a nearly statistically
7	significant increase in pregnancy duration (short-term exposure) in mice when exposed to
8	0.8 or 1.2 ppm O ₃ . This phenomenon was most likely due to the anorexigenic effect of
9	relatively high O ₃ concentrations (<u>Bignami et al., 1994</u>).
10	Table 7-5 provides a brief overview of the epidemiologic studies of PTB. In summary,
11	the evidence is consistent when examining short-term exposure to O ₃ during late
12	pregnancy and reports no association with PTB. However when long-term exposure to O_3
13	early in pregnancy is examined the results are inconsistent. Generally, studies conducted
14	in the U.S., Canada, and England find no association with O ₃ and PTB, while studies
15	conducted in Australia and China report an O ₃ effect on PTB.

Study	Location Sample Size	Mean O₃ (ppb)	Exposure assessment	Effect Estimate ^a (95% CI)
Darrow et al. (2009)	Atlanta, GA (n = 476,489)	8-h max: 44.1	Population-weighted spatial averages Nearest Monitor (within 4 miles)	First month: 0.98 (0.97, 1.00) Last week: 0.99 (0.98, 1.00) Last 6 weeks: 1.00 (0.98, 1.02)
<u>Ritz et al. (2000</u>)	California, U.S. (n = 97,158)	8 h: 36.9	<2 mi of monitor	First month: NR Last 6 weeks: NR
Wilhelm and Ritz (2005)	Los Angeles, CA (n = 106,483)	1 h: 21.1- 22.2	Varying distances to monitor	First month: 1.23 (1.06, 1.42) T1: NR T2: 1.38 (1.14, 1.66) Last 6 weeks: NR
<u>Ritz et al. (2007</u>)	Los Angeles, CA (n = 58,316)	24-h avg: 22.5	Nearest monitor to ZIP code	Entire pregnancy: NR T1: 0.93 (0.82, 1.06) Last 6 weeks: NR
Brauer et al. (2008)	Vancouver, BC, Canada (n = 70,249)	24-h avg: 14	Nearest Monitor (within 10 km) Inverse Distance Weighting (IDW)	Entire pregnancy: NR First 30 days of pregnancy: NR Last 30 days of pregnancy: NR T1: NR T3: NR
Lee et al. (2008c)	London, UK	24-h avg: NR	1 monitor	Lag 0: 1.00 (1.00, 1.01)
Hansen et al. (2006)	Brisbane, Australia (n = 28,200)	8-h max: 26.7	Citywide avg	T1: 1.39 (1.15, 1.70) T3: 1.09 (0.88, 1.39)
<u>Jalaludin et al.</u> (2007)	Sydney, Australia (n = 123,840)	1-h max: 30.9	Citywide avg and <5 km from monitor	First month: 1.04 (0.95, 1.13) T1: 1.15 (1.05, 1.24) T3: 0.98 (0.89, 1.07) Last month: 0.98 (0.88, 1.06)
<u>Jiang et al. (2007)</u>	Shanghai, China (n = 3,346 preterm births)	8-h avg: 32.7	Citywide avg	4 wks before birth: 1.06 (1.00, 1.12) 6 wks before birth: 1.06 (0.99, 1.13) 8 wks before birth: 1.09 (1.01, 1.19) L0: NR (results presented in figure) L1: NR (results presented in figure) L2: NR (results presented in figure) L3: NR (results presented in figure) L4: NR (results presented in figure) L5: NR (results presented in figure) L6: NR (results presented in figure)

Table 7-5 Brief summary of epidemiologic studies of PTB

^aRelative risk of PTB per 10 ppb change in O_3 .

T1 = First Trimester, T2 = Second Trimester, T3 = Third Trimester.

L0 = Lag 0, L1= Lag 1, L2 = Lag 2, L3 = Lag 3, L4 = Lag 4, L5 = Lag 5, L6 = Lag 6.

NR: No quantitative results reported.

7.4.5 Fetal Growth

1 2 3 4 5 6 7 8 9	Low birth weight has often been used as an outcome measure because it is easily available and accurately recorded on birth certificates. However, LBW may result from either short gestation, or inadequate growth in utero. Most of the studies investigating air pollution exposure and LBW limited their analyses to term infants to focus on inadequate growth. A number of studies were identified that specifically addressed growth restriction in utero by identifying infants who failed to meet specific growth standards. Usually these infants had birth weight less than the 10th percentile for gestational age, using an external standard. Many of these studies have been previously discussed, since they also examined other reproductive outcomes (i.e., LBW or PTB).
10 11 12 13 14 15	Fetal growth is influenced by maternal, placental, and fetal factors. The biological mechanisms by which air pollutants may influence the developing fetus remain largely unknown. Several mechanisms have been proposed, and are the same as those hypothesized for birth weight (see Section $5.4.2.4$). Additionally, in animal toxicology studies, O ₃ causes transient anorexia in exposed pregnant dams. This may be one of many possible contributors to O ₃ -dependent decreased fetal growth.
16 17 18 19 20 21 22 23 24	A limitation of environmental studies that use birth weight as a proxy measure of fetal growth is that patterns of fetal growth during pregnancy cannot be assessed. This is particularly important when investigating pollutant exposures during early pregnancy as birth weight is recorded many months after the exposure period. The insult of air pollution may have a transient effect on fetal growth, where growth is hindered at one point in time but catches up at a later point. For example, maternal smoking during pregnancy can alter the growth rate of individual body segments of the fetus at variable developmental stages, as the fetus experiences selective growth restriction and augmentation (Lampl and Jeanty, 2003).
25 26 27 28 29 30 31 32 33 34 35 36	The terms small-for-gestational-age (SGA), which is defined as a birth weight <10th percentile for gestational age (and often sex and/or race), and intrauterine growth retardation (IUGR) are often used interchangeably. However, this definition of SGA does have limitations. For example, using it for IUGR may overestimate the percentage of "growth-restricted" neonates as it is unlikely that 10% of neonates have growth restriction (Wollmann, 1998). On the other hand, when the 10th percentile is based on the distribution of live births at a population level, the percentage of SGA among PTB is most likely underestimated (Hutcheon and Platt, 2008). Nevertheless, SGA represents a statistical description of a small neonate, whereas the term IUGR is reserved for those with clinical evidence of abnormal growth. Thus all IUGR neonates will be SGA, but not all SGA neonates with be IUGR (Wollmann, 1998). In the following section the terms SGA and IUGR are referred to as each cited study used the terms.

- 1 Over the past decade a number of studies examined various metrics of fetal growth 2 restriction. Salam et al. (2005) assessed the effect of increasing O_3 concentrations on 3 IUGR in a population of infants born in California from 1975-1987 as part of the 4 Children's Health Study. The authors reported that maternal O₃ exposures averaged over 5 the entire pregnancy and during the third trimester were associated with increased risk of 6 IUGR. A 10-ppb difference in 24-h maternal O₃ exposure during the third trimester 7 increased the risk of IUGR by 11% (95% CI: 0, 20%). Brauer et al. (2008) evaluated the 8 impacts of air pollution (CO, NO₂, NO, O₃, SO₂, PM₂₅, PM₁₀) on SGA (1999-2002) using 9 spatiotemporal residential exposure metrics by month of pregnancy in Vancouver, BC. 10 The O₃ associations were largely protective (OR = 0.87, [95% CI: 0.81, 0.93] for a 11 10 ppb increase in inverse distance weighted SGA), and no additional quantitative results 12 were presented for O_3 . Liu et al. (2007b) examined the association between IUGR among 13 singleton term live births and SO₂, NO₂, CO, O₃, and PM_{2.5} in 3 Canadian cities for the 14 period 1985-2000. No increase in the risk of IUGR in relation to exposure to O_3 averaged 15 over each month and trimester of pregnancy was noted.
- 16 Three studies conducted in Australia provide evidence for an association between 17 ambient O₃ and fetal growth restriction. Hansen et al. (2007) examined SGA among 18 singleton, full-term births in Brisbane, Australia in relation to ambient air pollution (bsp, 19 PM_{10} , NO₂, O₃) during pregnancy. They also examined head circumference and crown-20 heel length in a subsample of term neonates. Trimester specific exposures to all pollutants 21 were not statistically significantly associated with a reduction in head circumference or 22 an increased risk of SGA. When monthly-specific exposures were examined, the authors 23 observed an increased risk of SGA associated with exposure to O₃ during month 4 24 (OR = 1.11 [95% CI: 1.00, 1.24] per 10 ppb increase). In a subsequent study, Hansen et 25 al. (2008) examined the possible associations between fetal ultrasonic measurements and 26 ambient air pollution (PM₁₀, O₃, NO₂, SO₂) during early pregnancy. This study had two 27 strengths: (1) fetal growth was assessed during pregnancy as opposed to at birth; and (2) 28 there was little delay between exposures and fetal growth measurements, which reduces 29 potential confounding and uses exposures that are concurrent with the observed growth 30 pattern of the fetus. Fetal ultrasound biometric measurements were recorded for biparietal diameter (BPD), femur length, abdominal circumference, and head circumference. To 31 32 further improve exposure assessment, the authors restricted the samples to include only 33 scans from women for whom the centroid of their postcode was within 14 km of an air 34 pollution monitoring site. Ozone during days 31-60 was associated with decreases in all 35 of the fetal growth measurements, and a 1.78 mm reduction in abdomen circumference 36 per 10 ppb increase in O_3 concentration, though this effect did not persist in copollutant 37 models. The change in ultrasound measurements associated with O₃ during days 31-60 of 38 gestation indicated that increasing O_3 concentration decreased the magnitude of 39 ultrasound measurements for women living within 2 km of the monitoring site. The

1	relationship decreased toward the null as the distance from the monitoring sites increased.
2	When assessing effect modification due to SES, there was some evidence of effect
3	modification for most of the associations, with the effects of air pollution stronger in the
4	highest SES quartile. In the third study, Mannes et al. (2005) estimated the effects of
5	pollutant (PM ₁₀ , PM _{2.5} , NO ₂ , CO and O ₃) exposure in the first, second and third trimesters
6	of pregnancy and risk of SGA in Sydney, Australia. Citywide average air pollutant
7	concentrations in the last month, third trimester, and first trimester of pregnancy had no
8	effect on SGA. Concentrations of O ₃ in the second trimester of pregnancy had small but
9	adverse effects on SGA (OR = 1.10 [95% CI: 1.00, 1.14] per 10 ppb increment). This
10	effect disappeared when the analysis was limited to births with a maternal address within
11	5 km of a monitoring station (OR = 1.00 [95% CI: 0.60, 1.79] per 10 ppb increment).
12	Very little information from toxicological studies is available to address effects on fetal
13	growth. However, there is evidence to suggest that prenatal (short-term) exposure to O_3
14	can affect postnatal growth. A few studies reported that mice or rats exposed
15	developmentally (gestationally \pm lactationally) to O ₃ had deficits in body weight gain in
16	the postpartum period (Bignami et al., 1994; Haro and Paz, 1993; Kavlock et al., 1980).
17	Table 7-6 provides a brief overview of the epidemiologic studies of fetal growth
18	restriction. In summary, the evidence is inconsistent when examining exposure to O_3 and
19	fetal growth restriction. Similar to PTB, studies conducted in Australia have reported an
20	effect of O ₃ on fetal growth, whereas studies conducted in other areas generally have not
21	found such an effect. This may be due to the restriction of births to those within 2-14 km
22	of a monitoring station, as was done in the Australian studies.

Study	Location (Sample Size)	Mean O ₃ (ppb)	Exposure assessment	Effect Estimate ^a (95% CI)
<u>Salam et al.</u> (2005)	California, U.S. (n = 3901)	24-h avg: 27.3 8 h: 50.6	ZIP code level	Entire pregnancy: 1.16 (1.00, 1.32) T1:1.00 (0.94, 1.11) T2: 1.06 (1.00, 1.12) T3: 1.11 (1.00, 1.17)
<u>Brauer et al.</u> (2008)	Vancouver, BC, Canada (n = 70,249)	24-h avg: 14	Nearest Monitor (within 10 km) Inverse Distance Weighting (IDW)	Entire pregnancy: NR First 30 days of pregnancy: NR Last 30 days of pregnancy: NR T1: NR T3: NR
<u>Liu et al. (2007b</u>)	Calgary, Edmonton, and Montreal, Canada (n = 16,430)	24-h avg: 16.5 1-h max: 31.2	Census Subdivision avg	Entire pregnancy: NR (results presented in figure) T1: NR (results presented in figure) T2: NR (results presented in figure) T3: NR (results presented in figure)
<u>Hansen et al.</u> (2007)	Brisbane, Australia (n = 26,617)	8-h max: 26.7	Citywide avg	T1: 1.01 (0.89, 1.15) T2: 1.00 (0.86, 1.17) T3: 0.83 (0.71, 0.97)
<u>Hansen et al.</u> (2008)	Brisbane, Australia (n = 15,623)	8-h avg: 24.8	Within 2 km of monitor	M1: -0.32 (-1.56, 0.91) ^b M2: -0.58 (-1.97, 0.80) ^b M3: 0.26 (-1.07, 1.59) ^b M4: 0.11 (-0.98, 1.21) ^b
<u>Mannes et al.</u> (2005)	Sydney, Australia (n = 138,056)	1-h max: 31.6	Citywide avg and <5 km from monitor	T1: 0.90 (0.48, 1.34) T2: 1.00 (0.60, 1.79) T3: 1.10 (0.66, 1.97) Last 30 days of pregnancy: 1.10 (0.74 1.79)

Table 7-6Brief summary of epidemiologic studies of fetal growth.

^aRelative risk of fetal growth restriction per 10 ppb change in O₃, unless otherwise noted.

^bMean change in fetal ultrasonic measure of head circumference recorded between 13 and 26 weeks gestation for a 10-ppb increase in maternal exposure to O_3 during early pregnancy

T1 = First Trimester, T2 = Second Trimester, T3 = Third Trimester

M1 = Month 1, M2 = Month 2, M3 = Month 3, M4 = Month 4

NR: No quantitative results reported

7.4.6 Postnatal Growth

1	Postnatal weight and height are routinely measured in children as indicators of growth
2	and somatic changes. Toxicological studies often follow these endpoints to ascertain if a
3	known exposure has an effect in the postnatal window, an effect which can be permanent.
4	Time-pregnant BALB/c mice were exposed to O_3 (0, 0.4, 0.8, or 1.2 ppm) GD9-18
5	(short-term exposure) with parturition at GD20-21 (Sharkhuu et al., 2011). As the
6	offspring aged, postnatal litter body weight continued to be significantly decreased in the
7	highest concentration (1.2 ppm) O_3 group at PND3 and PND7. When the pups were

1	weighed separately by sex at PND42, the males with the highest concentration of O_3
2	exposure (1.2 ppm, GD9-18) had significant decrements in body weight (Sharkhuu et al.,
3	<u>2011</u>).
4	Significant decrements in body weight at 4 weeks of age were reported in C57Bl/6 mice
5	that were exposed to postnatal O_3 (short-term exposure, PND2-28 exposure, 1 ppm O_3 ,
6	3 hours/day, 3 days/week) (Auten et al., 2012). Animals with co-exposure to in utero DE
7	(short-term exposure, dam GD9-GD17; inhalation 0.5 or 2.0 mg/m ³ O ₃ ; 4 h/day via
8	inhalation; or oropharyngeal aspiration DEPs, 2×/week) + postnatal O ₃ (aforementioned
9	short-term exposure) also had significantly reduced body weight.

7.4.7 Birth Defects

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Despite the growing body of literature evaluating the association between ambient air pollution and various adverse birth outcomes, relatively few studies have investigated the effect of temporal variations in ambient air pollution on birth defects. Heart defects and oral clefts have been the focus of the majority of these recent studies, given the higher prevalence than other birth defects and associated mortality. Mechanistically, air pollutants could be involved in the etiology of birth defects via a number of key events (see Section 5.4.2.4).

17 Several studies have been conducted examining the relationship between O_3 exposure 18 during pregnancy and birth defects and reported a positive association with cardiac 19 defects. The earliest of these studies was conducted in southern California (Ritz et al., 20 2002). This study evaluated the effect of air pollution on the occurrence of cardiac birth 21 defects in neonates and fetuses delivered in southern California in 1987-1993. Maternal 22 exposure estimates were based on data from the fixed site closest to the mother's ZIP 23 code area. When using a case-control design where cases were matched to 10 randomly 24 selected controls, results showed increased risks for aortic artery and valve defects 25 (OR = 1.56 [95% CI: 1.16, 2.09] per 10 ppb O₃), pulmonary artery and valve anomalies 26 (OR = 1.34 [95% CI: 0.96, 1.87] per 10 ppb O₃), and conotruncal defects (OR = 1.36) 27 [95% CI: 0.91, 2.03] per 10 ppb O₃) in a dose-response manner with second-month O₃ 28 exposure. A study conducted in Texas (Gilboa et al., 2005) looked at a similar period of 29 exposure but reported no association with most of the birth defects studied (O_3) 30 concentration was studied using quartiles with the lowest representing <18 ppb and the 31 highest representing ≥ 31 ppb). The authors found slightly elevated odds ratios for 32 pulmonary artery and valve defects. They also detected an inverse association between O_3 33 exposure and isolated ventricular septal defects. Overall, this study provided some weak 34 evidence that air pollution increases the risk of cardiac defects. Hansen et al. (2009)

1	investigated the possible association between ambient air pollution concentrations
2	averaged over weeks 3-8 of pregnancy and the risk of cardiac defects. When analyzing all
3	births with exposure estimates for O_3 from the nearest monitor there was no indication for
4	an association with cardiac defects. There was also no adverse association when
5 6	restricting the analyses to only include births where the mother resided within 12 km of a
	monitoring station. However, among births within 6 km of a monitor, a 10 ppb increase
7	in O_3 was associated with an increased risk of pulmonary artery and valve defects
8	(OR = 8.76 [95% CI: 1.80, 56.55]). As indicated by the very wide credible intervals,
9	there were very few cases in the sensitivity analyses for births within 6 km of a monitor,
10	and this effect could be a result of type I errors. <u>Dadvand et al. (2011</u>) investigated the
11	association between maternal exposure to ambient air pollution concentrations averaged
12	over weeks 3-8 of pregnancy and the occurrence of cardiac birth defects in England.
13	Similar to <u>Hansen et al. (2009</u>), they found no associations with maternal exposure to O_3
14	except for when the analysis was limited to those subjects residing within a 16 km
15	distance of a monitoring station (OR for malformations of pulmonary and tricuspid
16	valves=1.64 [95% CI: 1.04, 2.60] per 10 ppb increase in O ₃).
17	Despite the association between O_3 and cardiac defects observed in the above studies, a
18	recent study did not observe an increased risk of cardiac birth defects associated with
19	ambient O_3 concentrations. The study, conducted in Atlanta, GA, examined O_3 exposure
20	during weeks 3-7 of of pregnancy and reported no association with risk of cardiovascular
21	malformations (Strickland et al., 2009).
22	Several of these studies have also examined the relationship between O_3 exposure during
23	pregnancy and oral cleft defects. The study by <u>Ritz et al. (2002</u>) evaluated the effect of air
24	pollution on the occurrence of orofacial birth defects and did not observe strong
25	associations between ambient O_3 concentration and orofacial defects. They did report an
26	OR of 1.13 (95% CI: 0.90, 1.40) per 10 ppb during the second trimester for cleft lip with
27	or without cleft palate. Similarly, <u>Gilboa et al. (2005</u>) reported an OR of 1.09 (95% CI:
28	0.70, 1.69) for oral cleft defects when the fourth quartile was contrasted with the first
29	quartile of exposure during 3-8 weeks of pregnancy. <u>Hansen et al. (2009</u>) reported no
30	indication for an association with cleft defects and air pollution concentrations averaged
31	over weeks 3-8 of pregnancy. <u>Hwang and Jaakkola (2008</u>) conducted a population-based
32	case-control study to investigate exposure to ambient air pollution and the risk of cleft lip
33	with or without cleft palate in Taiwan. The risk of cleft lip with or without cleft palate
34	was increased in relation to O_3 levels in the first gestational month (OR = 1.17 [95% CI:
35	1.01, 1.36] per 10 ppb) and second gestational month (OR = 1.22 [95% CI: 1.03, 1.46]
36	per 10 ppb), but was not related to any of the other pollutants. In three-pollutant models,
37	the effect estimates for O_3 exposure were stable for the four different combinations of
38	-
50	pollutants and were all statistically significant. Marshall et al. (2010) compared estimated

1	exposure to ambient pollutants during early pregnancy (6 week period from 5 to 10
2	weeks into the gestational period) among mothers of children with oral cleft defects to
3	that among mothers of controls. The authors observed no consistent elevated associations
4	between any of the air pollutants examined and cleft malformations, though there was a
5	weak association between cases of cleft palate only and increasing O ₃ concentrations.
6	This association increased when cases and controls were limited to those with residences
7	within 10 km of the closest O_3 monitor (OR = 2.2 [95% CI: 1.0, 4.9], comparing highest
8	quartile [>33 ppb] to lowest quartile [<15 ppb]).
9	A limited number of toxicological studies have examined birth defects in animals
10	exposed gestationally to O ₃ . Kavlock et al. (1979) exposed pregnant rats to O ₃ for precise
11	periods during organogenesis. No significant teratogenic effects were found in rats
12	exposed 8 h/day to concentrations of O_3 varying from 0.44 to 1.97 ppm during early
13	(days 6-9), mid (days 9-12), or late (days 17 to 20) gestation, or the entire period of
14	organogenesis (days 6-15) (short-term exposures). Earlier research found eyelid
15	malformation following gestational and postnatal exposure to 0.2 ppm O ₃ (Veninga,
16	<u>1967</u>).
16 17	<u>1967</u>). <u>Table 7-7</u> provides a brief overview of the epidemiologic studies of birth defects. These
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17 18 19 20 21 22 23 24 25 26	Table 7-7 provides a brief overview of the epidemiologic studies of birth defects. These studies have focused on cardiac and oral cleft defects, and the results from these studies are not entirely consistent. This inconsistency could be due to the absence of true associations between O_3 and risks of cardiovascular malformations and oral cleft defects; it could also be due to differences in populations, pollution levels, outcome definitions, or analytical approaches. The lack of consistency of associations between O_3 and cardiovascular malformations or oral cleft defects might be due to issues relating to statistical power or measurement error. A recent meta-analysis of air pollution and congenital anomalies concluded that there was no statistically significant increase in risk
17 18 19 20 21 22 23 24 25 26 27	Table 7-7 provides a brief overview of the epidemiologic studies of birth defects. These studies have focused on cardiac and oral cleft defects, and the results from these studies are not entirely consistent. This inconsistency could be due to the absence of true associations between O_3 and risks of cardiovascular malformations and oral cleft defects; it could also be due to differences in populations, pollution levels, outcome definitions, or analytical approaches. The lack of consistency of associations between O_3 and cardiovascular malformations or oral cleft defects might be due to issues relating to statistical power or measurement error. A recent meta-analysis of air pollution and congenital anomalies concluded that there was no statistically significant increase in risk of congenital anomalies and O_3 (Vrijheid et al., 2011). These authors note that heterogeneity in the results of these studies may be due to inherent differences in study location, study design, and/or analytic methods, and comment that these studies have not
17 18 19 20 21 22 23 24 25 26 27 28	Table 7-7 provides a brief overview of the epidemiologic studies of birth defects. These studies have focused on cardiac and oral cleft defects, and the results from these studies are not entirely consistent. This inconsistency could be due to the absence of true associations between O_3 and risks of cardiovascular malformations and oral cleft defects; it could also be due to differences in populations, pollution levels, outcome definitions, or analytical approaches. The lack of consistency of associations between O_3 and cardiovascular malformations or oral cleft defects might be due to issues relating to statistical power or measurement error. A recent meta-analysis of air pollution and congenital anomalies concluded that there was no statistically significant increase in risk of congenital anomalies and O_3 (Vrijheid et al., 2011). These authors note that heterogeneity in the results of these studies may be due to inherent differences in study

Study	Outcomes Examined	Location (Sample Size)	Mean O₃ (ppb)	Exposure Assessment	Exposure Window
<u>Ritz et al. (2002</u>)	Cardiac and Cleft Defects	Southern California (n = 3,549 cases; 10,649 controls)	24-h avg: NR	Nearest Monitor (within 10 mi)	Month 1,2,3 Trimester 2,3 3-mo period prior to conception
<u>Gilboa et al. (2005</u>)	Cardiac and Cleft Defects	7 Counties in TX (n = 5,338 cases; 4,580 controls)	24-h avg: NR	Nearest Monitor	Weeks 3-8 of gestation
<u>Hwang and Jaakkola</u> (2008)	Oral Cleft Defects	Taiwan (n = 653 cases; 6,530 controls)	24-h avg: 27.31	Inverse Distance Weighting (IDW)	Months 1,2,3
Strickland et al. (2009)	Cardiac Defects	Atlanta, GA (n = 3,338 cases)	8-h max: 39.8-43.3	Weighted citywide avg	Weeks 3-7 of gestation
<u>Hansen et al. (2009</u>)	Cardiac and Cleft Defects	Brisbane, Australia (n = 150,308 births)	8-h max: 25.8	Nearest Monitor	Weeks 3-8 of gestation
Marshall et al. (2010)	Oral Cleft Defects	New Jersey (n = 717 cases; 12,925 controls)	24-h avg: 25	Nearest Monitor (within 40 km)	Weeks 5-10 of gestation
<u>Dadvand et al.</u> (2011)	Cardiac Defects	Northeast England (n = 2,140 cases; 14,256 controls)	24-h avg: 18.8	Nearest Monitor	Weeks 3-8 of gestation`1

Table 7-7 Brief summary of epidemiologic studies of birth defects

7.4.8 Developmental Respiratory Effects

1	The issue of prenatal exposure has assumed increasing importance since ambient air
1	
2	pollution exposures of pregnant women have been shown to lead to adverse pregnancy
3	outcomes, as well as to respiratory morbidity and mortality in the first year of life.
4	Growth and development of the respiratory system take place mainly during the prenatal
5	and early postnatal periods. This early developmental phase is thought to be very
6	important in determining long-term lung growth. Studies have recently examined this
7	emerging issue. Several studies were included in Section <u>7.2.1</u> and Section <u>7.2.3</u> , and are
8	included here because they reported both prenatal and post-natal exposure periods.
9	Mortimer et al. (2008a); (2008b) examined the association of prenatal and lifetime
10	exposures to air pollutants with pulmonary function and allergen sensitization in a subset
11	of asthmatic children (ages 6-11) included in the Fresno Asthmatic Children's
12	Environment Study (FACES). Monthly means of pollutant levels for the years 1989-2000
13	were created and averaged separately across several important developmental time-
14	periods, including the entire pregnancy, each trimester, the first 3 years of life, the first
15	6 years of life, and the entire lifetime. The 8-h avg O_3 concentrations were approximately

1	50 ppb for each of the exposure metrics (estimated from figure). In the first analysis
2	(Mortimer et al., 2008a), negative effects on pulmonary function were found for exposure
3	to PM ₁₀ , NO ₂ , and CO during key neonatal and early life developmental periods. The
4	authors did not find a negative effect of exposure to O3 among this cohort. In the second
5	analysis (Mortimer et al., 2008b), sensitization to at least one allergen was associated, in
6	general, with higher levels of CO and PM_{10} during the entire pregnancy and second
7	trimester and higher PM_{10} during the first 2 years of life. Lower exposure to O_3 during the
8	entire pregnancy or second trimester was associated with an increased risk of allergen
9	sensitization. Although the pollutant metrics across time periods are correlated, the
10	strongest associations with the outcomes were observed for prenatal exposures. Though it
11	may be difficult to disentangle the effect of prenatal and postnatal exposures, the models
12	from this group of studies suggest that each time period of exposure may contribute
13	independently to different dimensions of school-aged children's pulmonary function. For
14	4 of the 8 pulmonary-function measures (FVC, FEV ₁ , PEF, FEF ₂₅₋₇₅), prenatal exposures
15	were more influential on pulmonary function than early-lifetime metrics, while, in
16	contrast, the ratio of measures (FEV $_1$ /FVC and FEF $_{25-75}$ /FVC) were most influenced by
17	postnatal exposures. When lifetime metrics were considered alone, or in combination
18	with the prenatal metrics, the lifetime measures were not associated with any of the
19	outcomes, suggesting the timing of the exposure may be more important than the overall
20	dose and prenatal exposures are not just markers for lifetime or current exposures.

- 21 <u>Clark et al. (2010)</u> investigated the effect of exposure to ambient air pollution in utero 22 and during the first year of life on risk of subsequent asthma diagnosis (incident asthma 23 diagnosis up to age 3-4) in a population-based nested case-control study. Air pollution 24 exposure for each subject based on their residential address history was estimated using 25 regulatory monitoring data, land use regression modeling, and proximity to stationary 26 pollution sources. An average exposure was calculated for the duration of pregnancy 27 (~15 ppb) and the first year of life (~14 ppb). In contrast to the Mortimer et al. (2008a); 28 (2008b) studies, the effect estimates for first-year exposure were generally larger than for 29 in utero exposures. However, similar to the Mortimer et al. studies, the observed 30 associations with O_3 were largely protective. Because of the relatively high correlation 31 between in utero and first-year exposures for many pollutants, it was difficult to discern 32 the relative importance of the individual exposure periods.
- 33Latzin et al. (2009) examined whether prenatal exposure to air pollution was associated34with lung function changes in the newborn. Tidal breathing, lung volume, ventilation35inhomogeneity and eNO were measured in 241 unsedated, sleeping neonates (age=365 weeks). The median of the 24-h avg O3 concentrations averaged across the post-natal37period was ~44 ppb. Consistent with the previous studies, no association was found for38prenatal exposure to O3 and lung function.

- 1The new toxicological literature since the 2006 O3 AQCD, covering respiratory changes2related to developmental O3 exposure, reports ultrastructural changes in bronchiole3development, alterations in placental and pup cytokines, and increased pup airway hyper-4reactivity. These studies are detailed below.
- 5 Fetal rat lung bronchiole development is triphasic, comprised of the glandular phase 6 (measured at GD18), the canalicular phase (GD20), and the saccular phase (GD21). The 7 ultrastructural lung development in fetuses of pregnant rats exposed to 1-ppm O₃ (12 8 h/day, out to either GD18, GD20 or GD21) was examined by electron microscopy during 9 these three phases. In the glandular phase, bronchiolar columnar epithelial cells in fetuses 10 of dams exposed to O₃ had cytoplasmic damage and swollen mitochondria. Bronchial 11 epithelium at the canalicular phase in O_3 exposed pups had delayed maturation in 12 differentiation, i.e., glycogen abundance in secretory cells had not diminished as it should 13 with this phase of development. Congruent with this finding, delayed maturation of 14 tracheal epithelium following early neonatal O_3 exposure (1 ppm, 4-5 h/day for first week 15 of life) in lambs has been previously reported (Mariassy et al., 1990; Mariassy et al., 16 1989). Also at the canalicular phase, atypical cells were seen in the bronchiolar lumen of 17 O₃-exposed rat fetuses. Finally, in the saccular phase, mitochondrial degradation was 18 present in the non-ciliated bronchiolar cells of rats exposed in utero to O₃. In conclusion, 19 O₃ exposure of pregnant rats produced ultra-structural damage to near-term fetal 20 bronchiolar epithelium (López et al., 2008).
- 21 Exposure of laboratory animals to multiple airborne pollutants can differentially affect 22 pup physiology. One study showed that exposure of C57BL/6 mouse dams to 0.48 mg 23 PM intratracheally twice weekly for 3 weeks during pregnancy augmented O₃-induced 24 airway hyper-reactivity in juvenile offspring. Maternal PM exposure also significantly 25 increased placental cytokines above vehicle-instilled controls. Pup postnatal O₃ exposure 26 (1 ppm 3 h/day, every other day, thrice weekly for 4 weeks) induced significantly 27 increased cytokine levels (IL-1 β , TNF- α , KC, and IL-6) in whole lung versus postnatal 28 air exposed groups; this was further exacerbated with gestational PM exposure (Auten et 29 al., 2009). In further studies by the same laboratory, O₃-induced AHR was studied in 30 rodent offspring after dam gestational exposure to inhaled diesel exhaust (Auten et al., 31 2012). Pregnant C57Bl/6 mice were exposed to diesel exhaust GD9-17 (0.5 or 2.0 mg/m³) 32 O_3 , 4h/day) via inhalation or in a separate set of animals via oropharyngeal aspiration of 33 freshly generated DEPs ($2\times$ /week). Postnatally, the offspring were exposed to O₃ starting 34 at PND2 (1 ppm O₃, 3 hours/day, 3 days/week for 4 weeks). Juvenile mice were then 35 subjected to measurements of pulmonary mechanisms (at 4 weeks of age and then at 8 36 weeks of age). Increased inflammation of the placenta and lungs of DE exposed fetuses 37 was reported at GD18. In animals with postnatal O₃ exposure alone, elevated 38 inflammation was seen with significant increased levels of BAL cytokines; these O₃-

- 1 related elevated levels were significantly exacerbated with prenatal DE exposure 2 $(DE+O_3)$. At PND28, DE+O₃ exposed offspring had significant impairment of alveolar 3 development as measured with secondary alveolar crest development, a finding that was 4 absent in all other exposure groups (O₃ alone, DE alone). Postnatal O₃ exposure induced 5 AHR in methacholine challenged animals at 4 weeks of age and was exacerbated with the 6 higher dose of DE exposure (DE+ O_3). At 8 weeks of age, O_3 exposed pups had persistent 7 AHR (+/-DE) that was significantly augmented in DE+O₃ pups. In summary, gestational 8 DE exposure induced an inflammatory response which, when combined with postnatal O_3 9 exposure impaired alveolar development, and caused an exacerbated and longer-lasting 10 O₃-induced AHR in offspring.
- 11 A series of experiments using infant rhesus monkeys repeatedly exposed to 0.5 ppm O_3 12 starting at one-month of age have examined the effect of O_3 alone or in combination with 13 an inhaled allergen on morphology and lung function (Plopper et al., 2007). Exposure to 14 O₃ alone or allergen alone produced small but not statistically significant changes in 15 baseline airway resistance and airway responsiveness, but the combined exposure to both 16 O_3 + antigen produced statistically significant and greater than additive changes in both 17 functional measurements. Additionally, cellular changes and significant structural 18 changes in the respiratory tract have been observed in infant rhesus monkeys exposed to 19 O₃ (Fanucchi et al., 2006). A more detailed description of these studies can be found in 20 Section 7.2.3 (Pulmonary Structure and Function), with mechanistic information found in 21 Section 5.4.2.4.
- 22Lung immunological response in O3 exposed pups was followed by analyzing BAL and23lung tissue. Sprague Dawley (SD) pups were exposed to a single 3h exposure of air or O324(0.6 ppm) on PND 13 (Han et al., 2011). Bronchoalveolar lavage (BAL) was performed2510 hours after the end of O3 exposure. BALF polymorphonuclear leukocytes (PMNs) and26total BALF protein were significantly elevated in O3 exposed pups. Lung tissue from O327exposed pups had significant elevations of manganese superoxide dismutase (SOD)28protein and significant decrements of extra-cellular SOD protein.
- 29 Various immunological outcomes were followed in offspring after their pregnant dams 30 (BALB/c mice) were exposed gestationally to O₃ (0, 0.4, 0.8, or 1.2 ppm, GD9-18) 31 (Sharkhuu et al., 2011). Delayed type hypersensitivity (DTH) was initiated with initial 32 BSA injection at 6 weeks of age and then challenge 7 days later. The normal edematous 33 response of the exposed footpad (thickness after BSA injection) was recorded as an 34 indicator of DTH. In female offspring, normal footpad swelling with BSA injection that 35 was seen in air exposed animals was significantly attenuated with O_3 exposure (0.8 and 36 1.2 ppm O_3), implying immune suppression of O_3 exposure specifically in DTH. Humoral 37 immunity was measured with the sheep red blood cell (SRBC) response. Animals

- 1received primary immunization with SRBC and then blood was drawn for SRBC IgM2measurement. A SRBC booster was given 2 weeks later with blood collected 5 days after3booster for IgG measurement. Maternal O3 exposure had no effect on humoral immunity4in the offspring as measured by IgG and IgM titers after SRBC primary and booster5immunizations (Sharkhuu et al., 2011).
- 6 Toxicity assessment and allergen sensitization was also assessed in these O_3 exposed 7 offspring. At PND42, animals were euthanized for analysis of immune and inflammatory 8 markers (immune proteins, inflammatory cells, T-cell populations in the spleen). A subset 9 of the animals was intra-nasally instilled or sensitized with ovalbumin on either PND2 10 and 3 or PND42 and 43. All animals were challenged with OVA on PND54, 55, and 56. 11 One day after final OVA challenge, lung function, lung inflammation and immune 12 response were determined. Offspring of O_3 exposed dams that were initially sensitized at 13 PDN3 (early) or PND42 (late) were tested to determine the level of allergic sensitization 14 or asthma-like inflammation after OVA challenge. Female offspring sensitized early in 15 life developed significant eosinophilia (1.2 ppm O_3) and elevated serum OVA-specific 16 IgE (1.2 ppm O_3), which is a marker of airway allergic inflammation. The females that 17 were sensitized early also had significant decrements in BALF total cells, macrophages, 18 and lymphocytes (1.2 ppm O_3). Offspring that were sensitized later (PND42) in life did 19 not develop the aforementioned changes in BALF, but these animals did develop modest, 20 albeit significant neutropenia (0.8 and 1.2 ppm O₃) (Sharkhuu et al., 2011).
- 21BALF cytology in non-sensitized animals was followed. BALF of offspring born to dams22exposed to O3 was relatively unaffected (cytokines, inflammatory cell numbers/types) as23were splenic T-cell subpopulations. LDH was significantly elevated in BALF of females24whose mothers were exposed to 1.2 ppm during pregnancy (Sharkhuu et al., 2011). In25summary, the females born to mothers exposed to O3 developed modest26immunocompromise. Males were unaffected (Sharkhuu et al., 2011).
- 27Overall, animal toxicological studies have reported ultrastructural changes in bronchiole28development, alterations in placental and pup cytokines, and increased pup airway hyper-29reactivity related to exposure to O_3 during the developmental period. Epidemiologic30studies have found no association between prenatal exposure to O_3 and growth and31development of the respiratory system. Fetal origins of disease have received a lot of32attention recently, thus additional research to further explore the inconsistencies between33these two lines of evidence is warranted.

7.4.9 Developmental Central Nervous System Effects

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The following sections describe the results of toxicological studies of O_3 and developmental central nervous system effects. No epidemiologic studies of this association have been published.

7.4.9.1 Laterality

4	Two reports of laterality changes in mice developmentally exposed to O_3 have been
5	reported in the literature. Mice developmentally exposed to 0.6 ppm O_3 (6 days before
6	breeding to weaning at PND21) showed a turning preference (left turns) distinct from air
7	exposed controls (clockwise turns) (Dell'Omo et al., 1995); in previous studies this
8	behavior in mice has been found to correlate with specific structural asymmetries of the
9	hippocampal mossy fiber projections (Schöpke et al., 1991). The 2006 O ₃ AQCD
10	evidence for the effect of O_3 on laterality or handedness demonstrated that rats exposed to
11	O3 during fetal and neonatal life showed limited, gender-specific changes in handedness
12	after exposure to the intermediate concentration of O_3 (only seen in female mice exposed
13	to 0.6 ppm O_3 , and not in males at 0.6 ppm or in either sex of 0.3 or 0.9 ppm O_3 with
14	exposure from 6 days before breeding to PND26) (Petruzzi et al., 1999).

7.4.9.2 Brain Morphology and Neurochemical Changes

15	The nucleus tractus solitarius (NTS), a medullary area of respiratory control, of adult
16	animals exposed prenatally to 0.5 ppm O_3 (12h/day, ED5-eD20) had significantly less
17	tyrosine hydroxylase staining versus control (Boussouar et al., 2009). Tyrosine
18	hydroxylase is the rate-limiting enzyme for dopamine synthesis and serves as a precursor
19	for catecholamine synthesis; thus, decreased staining is used as a marker of dopaminergic
20	or catecholaminergic cell or activity loss in these regions and thus functions in neuronal
21	plasticity. After physical restraint stress, control animals respond at the histological level
22	with Fos activation, a marker of neuronal activity, and tyrosine hydroxylase activation in
23	the NTS, a response which is absent or attenuated in adult animals exposed prenatally to
24	0.5 ppm O_3 (Boussouar et al., 2009) when compared to control air exposed animals who
25	also were restrained. The O3-exposed offspring in this study were cross-fostered to
26	control air exposed dams to avoid O3-dependent dam related neonatal effects on offspring
27	outcomes (i.e., dam behavioral or lactational contributions to pup outcomes) (Boussouar
28	<u>et al., 2009</u>).

1	Developmental exposure to 0.3 or 0.6 ppm O_3 prior to mating pair formation through
2	GD17 induced significant increased levels of BDNF in the striatum of adult (PND140)
3	O ₃ exposed offspring as compared to control air exposed animals; these O ₃ -exposed
4	animals also had significantly decreased level of NGF in the hippocampus versus control
5	(<u>Santucci et al., 2006</u>).
6	
6	Changes in the pup cerebellum with prenatal 1 ppm O ₃ exposure include altered
7	morphology (Romero-Velazquez et al., 2002; Rivas-Manzano and Paz, 1999), decreased
8	total area (Romero-Velazquez et al., 2002), decreased number of Purkinje cells (Romero-
9	Velazquez et al., 2002), and altered monoamine neurotransmitter content with the
10	catecholamine system affected and the indoleamine system unaffected by O_3 (Gonzalez-
11	<u>Pina et al., 2008</u>).

7.4.9.3 Neurobehavioral Outcomes

- O₃ administration to dams during pregnancy with or without early neonatal exposure has
 been shown to contribute to multiple neurobehavioral outcomes in offspring that are
 described in further detail below.
- 15O3 administration (0.4, 0.8 or 1.2 ppm O3) during the majority of pregnancy (PD7-17) of16CD-1 mice did not affect pup behavioral outcomes including early behavioral ultrasonic17vocalizations and more permanent later measurements (PND60 or 61) including pup18activity, habituation and exploration and d-amphetamine-induced hyperactivity (Bignami
- 19et al., 1994); these pups were all cross-fostered or reared on non- O3 exposed dams.20Testing for aggressive behavior in mice continuously exposed to O3 (0.3 or 0.6 ppm from2130 days prior to mating to GD17) revealed that mice had significantly increased
- 22defensive/ submissive behavior (increased freezing posturing on the first day only of a23multiple-day exam) versus air exposed controls (Santucci et al., 2006). Similarly,24continuous exposure of adult animals to O3 induced significant increases in fear behavior25and decreased aggression as measured by significantly decreased freezing behavior26(Petruzzi et al., 1995).
- 27Developmentally exposed animals also had significantly decreased amount of time spent28nose sniffing other mice (Santucci et al., 2006); this social behavior deficit, decreased29sniffing time, was not found in an earlier study with similar exposures (Petruzzi et al.,301995), but sniffing of specific body areas was measured in Santucci et al. (2006) and total31number of sniffs of the entire body was measured in Petruzzi et al. (1995). The two32toxicology studies exploring social behavior (sniffing) employ different study designs33and find opposite effects in animals exposed to O3.

7.4.9.4 Sleep Aberrations after Developmental Ozone Exposure

1	The effect of gestational O ₃ exposure (1 ppm O ₃ daily for 12h/day, during dark period for
2	the entire pregnancy) on sleep patterns in rat offspring was followed using 24 h
3	polysomnographic recordings at 30, 60 and 90 days of age (Haro and Paz, 1993).
4	Ozone-exposed pups manifested with inverted sleep-wake patterns or circadian rhythm
5	phase-shift. Rat vigilance was characterized in wakefulness, slow wave sleep (SWS), and
6	paradoxical sleep (PS) using previously characterized criteria. The O3 exposed offspring
7	spent longer time in the wakefulness state during the light period, more time in SWS
8	during the period of darkness, and showed significant decrements in PS. Chronic O_3
9	inhalation significantly decreased the duration of PS during both the light and dark
10	periods (Haro and Paz, 1993). These effects were consistent at all time periods measured
11	(30, 60 and 90 days of age). These sleep effects reported after developmental exposures
12	expand upon the existing literature on sleep aberrations in adult animals exposed to O_3
13	[rodents: (Paz and Huitron-Resendiz, 1996; Arito et al., 1992); and cats: (Paz and Bazan-
14	Perkins, 1992)]. A role for inhibition of cyclooxygenase-2 and the interleukins and
15	prostaglandins in the O_3 -dependent sleep changes potentially exists with evidence from a
16	publication on indomethacin pretreatment attenuating O3-induced sleep aberrations in
17	adult male animals (<u>Rubio and Paz, 2003</u>).

7.4.10 Early Life Mortality

18 Infants may be particularly at risk for the effects of air pollution. Within the first year of 19 life, infants develop rapidly; therefore their sensitivity may change within weeks or 20 months. During the neonatal and post-neonatal periods, the developing lung is highly 21 sensitive to environmental toxicants. The lung is not well developed at birth, with 80% of 22 alveoli being formed postnatally. An important question regarding the association 23 between O_3 and infant mortality is the critical window of exposure during development 24 for which infants are at risk. Several age intervals have been explored: neonatal 25 (<1 month); postneonatal (1 month to 1 year); and an overall interval for infants that 26 includes both the neonatal and postneonatal periods (<1 year). Within these various age 27 categories, multiple causes of deaths have been investigated, particularly total deaths and 28 respiratory-related deaths. The studies reflect a variety of study designs, exposure 29 periods, regions, and adjustment for confounders. As discussed below, a handful of 30 studies have examined the effect of ambient air pollution on neonatal and postneonatal 31 mortality, with the former the least studied. These studies varied somewhat with regard to 32 the outcomes and exposure periods examined and study designs employed.

7.4.10.1 Stillbirth

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Pereira et al. (1998) investigated the association among daily counts of intrauterine mortality (over 28 weeks of gestation) and air pollutant concentrations in Sao Paulo, Brazil from 1991 through 1992. The association was strong for NO₂, but lesser for SO₂ and CO. These associations exhibited a short lag time, less than 5 days. No significant association was detected between short-term O_3 exposure and intrauterine mortality.

7.4.10.2 Infant Mortality, Less than 1 Year

6	Ritz et al. (2006) linked birth and death certificates for infants who died between 1989
7	and 2000 to evaluate the influence of outdoor air pollution on infant death in the South
8	Coast Air Basin of California. The authors examined short- and long-term exposure
9	periods 2 weeks, 1 month, 2 months, and 6 months before each case subject's death and
10	reported no association between ambient levels of O ₃ and infant mortality. Similarly,
11	Diaz et al. (2004) analyzed the effects of extreme temperatures and short-term exposure
12	to air pollutants on daily mortality in children less than 1 year of age in Madrid, Spain,
13	from 1986 to 1997 and observed no statistically significant association between mortality
14	and O_3 concentrations. <u>Hajat et al. (2007</u>) analyzed time-series data of daily infant
15	mortality counts in 10 major cities in the UK to quantify any associations with short-term
16	changes in air pollution. When the results from the 10 cities were combined there was no
17	relationship between O ₃ and infant mortality, even after restricting the analysis to just the
18	summer months.
19	Conversely, a time-series study of infant mortality conducted in the southwestern part of

19Conversely, a time-series study of infant mortality conducted in the southwestern part of20Mexico City in the years 1993-1995 found that infant mortality was associated with21short-term exposure to NO2 and O3 3-5 days before death, but not as consistently as with22PM. A 10-ppb increase in 24-h avg O3 was associated with a 2.78% increase (95% CI:230.29, 5.26%) in infant mortality (lag 3) (Loomis et al., 1999). This increase was24attenuated, although still positive when evaluated in a two-pollutant model with PM2.5.25One-hour max concentrations of O3 exceeded prevailing Mexican and international26standards nearly every day.

7.4.10.3 Neonatal Mortality, Less than 1 Month

27	Several studies have evaluated ambient O ₃ concentrations and neonatal mortality and
28	observed no association. <u>Ritz et al. (2006</u>) linked birth and death certificates for infants
29	who died between 1989 and 2000 to evaluate the influence of outdoor air pollution on

1	infant death in the South Coast Air Basin of California. The authors examined short- and
2	long-term exposure periods 2 weeks, 1 month, 2 months, and 6 months before each case
3	subject's death and reported no association between ambient levels of O_3 and neonatal
4	mortality. Hajat et al. (2007) analyzed time-series data of daily infant mortality counts in
5	10 major cities in the UK to quantify any associations with short-term changes in air
6	pollution. When the results from the 10 cities were combined there was no relationship
7	between O_3 and neonatal mortality, even after restricting the analysis to just the summer
8	months. Lin et al. (2004a) assessed the impact of short-term changes in air pollutants on
9	the number of daily neonatal deaths in Sao Paulo, Brazil. The authors observed no
10	association between ambient levels of O_3 and neonatal mortality.

7.4.10.4 Postneonatal Mortality, 1 Month to 1 Year

11	A number of studies focused on the postneonatal period when examining the effects of O_3
12	on infant mortality. <u>Ritz et al. (2006</u>) linked birth and death certificates for infants who
13	died between 1989 and 2000 to evaluate the influence of outdoor air pollution on infant
14	death in the South Coast Air Basin of California. The authors examined short- and long-
15	term exposure periods 2 weeks, 1 month, 2 months, and 6 months before each case
16	subject's death and reported no association between ambient levels of O ₃ and
17	postneonatal mortality. Woodruff et al. (2008) evaluated the county-level relationship
18	between cause-specific postneonatal infant mortality and long-term early-life exposure
19	(first 2 months of life) to air pollutants across the U.S. Similarly, they found no
20	association between O3 exposure and deaths from respiratory causes. In the U.K., Hajat et
21	al. (2007) analyzed time-series data of daily infant mortality counts in 10 major cities to
22	quantify any associations with short-term changes in air pollution. When the results from
23	the 10 cities were combined there was no relationship between O_3 and postneonatal
24	mortality, even after restricting the analysis to just the summer months. In Ciudad Juarez,
25	Mexico, Romieu et al. (2004a) examined the daily number of deaths between 1997 and
26	2001, estimating the modifying effect of SES on the risk of postneonatal mortality.
27	Ambient O ₃ concentrations were not related to infant mortality overall, or in any of the
28	SES groups. In a follow-up study, Carbajal-Arroyo et al. (2011) evaluated the
29	relationship of 1-h daily max O_3 levels with postneonatal infant mortality in the
30	Mexico City Metropolitan Area between 1997 and 2005. Generally, short-term exposure
31	to O_3 was not significantly related to infant mortality. However, upon estimating the
32	modifying effect of SES on the risk of postneonatal mortality, the authors found that O_3
33	was statistically significantly related to respiratory mortality among those with low SES.
34	In a separate analysis, the effect of PM_{10} was evaluated with O_3 level quartiles. PM_{10}
35	alone was related to a significant increase in all-cause mortality. The magnitude of this

1	effect remained the same when only the days when O3 was in the lowest quartile were
2	included in the analyses. However, when only the days when O ₃ was in the highest
3	quartile were included in the analyses, the magnitude of the PM_{10} effect increased
4	dramatically (OR = 1.06 [95% CI: 0.909, 1.241] for PM_{10} on days with O_3 in lowest
5	quartile; OR = 1.26 [95% CI: 1.08, 1.47] for PM ₁₀ on days with O ₃ in the highest quartile.
6	These results suggest that while O ₃ alone may not have an effect on infant mortality, it
7	may serve to potentiate the observed effect of PM_{10} on infant mortality.
8	Tsai et al. (2006) used a case-crossover analysis to examine the relationship between
9	short-term exposure to air pollution and postneonatal mortality in Kaohsiung, Taiwan
10	during the period 1994-2000. The risk of postneonatal deaths was 1.023 (95% CI: 0.564,
11	1.858) per 10-ppb increase in 24-h avg O ₃ . The confidence interval for this effect
12	estimate is very wide, likely due to the small number of infants that died each day,
13	making it difficult to interpret this result. Several other studies conducted in Asia did not
14	find any association between O ₃ concentrations and infant mortality in the postneonatal
15	period. Ha et al. (2003) conducted a daily time-series study in Seoul, Korea to evaluate
16	the effect of short-term changes in ambient 8-h O ₃ concentrations on postneonatal
17	mortality. Son et al. (2008) examined the relationship between air pollution and
18	postneonatal mortality from all causes among firstborn infants in Seoul, Korea during
19	1999-2003. Yang et al. (2006) used a case-crossover analysis to examine the relationship
20	between air pollution exposure and postneonatal mortality in Taipei, Taiwan for the
21	period 1994-2000. The authors observed no associations between ambient levels of O_3
22	and postneonatal mortality.

7.4.10.5 Sudden Infant Death Syndrome

23	The strongest evidence for an association between ambient O ₃ concentrations and SIDS
24	comes from a study that evaluated the county-level relationship between SIDS and long-
25	term early-life exposure (first 2 months of life) to air pollutants across the U.S.
26	(Woodruff et al., 2008). The authors observed a 1.20 (95% CI: 1.09, 1.32) odds ratio for a
27	10-ppb increase in O_3 and deaths from SIDS. There was a monotonic increase in odds of
28	SIDS for each quartile of O_3 exposure compared with the lowest quartile (highest quartile
29	OR = 1.51; [95% CI: 1.17, 1.96]). In a multi-pollutant model including PM_{10} or $PM_{2.5}$,
30	CO and SO ₂ , the OR for SIDS and O_3 was not substantially lower than that found in the
31	single-pollutant model. When examined by season, the relationship between SIDS deaths
32	and O_3 was generally consistent across seasons with a slight increase for those babies
33	born in the summer. When stratified by birth weight, the OR for LBW babies was 1.27
34	(95% CI: 0.95, 1.69) per 10-ppb increase in O_3 and the OR for normal weight babies was
35	1.16 (95% CI: 1.01, 1.32) per 10-ppb increase in O ₃ .

1	Conversely, two additional studies reported no association between ambient levels of O_3
2	and SIDS. Ritz et al. (2006) linked birth and death certificates for infants who died
3	between 1989 and 2000 to evaluate the influence of outdoor air pollution on infant death
4	in the South Coast Air Basin of California. The authors examined short- and long-term
5	exposure periods 2 weeks, 1 month, 2 months, and 6 months before each case subject's
6	death and reported no association between ambient levels of O_3 and SIDS. <u>Dales et al.</u>
7	(2004) used time-series analyses to compare the daily mortality rates for SIDS and short-
8	term air pollution concentrations in 12 Canadian cities during the period of 1984-1999.
9	Increased daily rates of SIDS were associated with previous day increases in the levels of
10	SO_2 , NO_2 , and CO , but not O_3 or $PM_{2.5}$.
11	Table 7-8 provides a brief overview of the epidemiologic studies of infant mortality.

12 These studies have focused on short-term exposure windows (e.g., 1-3 days) and long-13 term exposure windows (e.g., up to 6 months). Collectively, they provide no evidence for

Study	Location	Mean O₃ (ppb)	Exposure Assessment	Effect Estimate ^a (95% CI):
<u>Pereira et al. (1998</u>)	Sao Paulo, Brazil	1-h max: 33.8	Citywide avg	L0-2: 1.00 (0.99, 1.01)
<u>Diaz et al. (2004</u>)	Madrid, Spain	24-h avg: 11.4	Citywide avg	NR
<u>Loomis et al. (1999</u>)	Mexico City, Mexico	24-h avg: 44.1 1-h max: 163.5	1 monitor	L0: 0.99 (0.97, 1.02) L1: 0.99 (0.96, 1.01) L2: 1.00 (0.98, 1.03) L3: 1.03 (1.00, 1.05) L4: 1.01 (0.98, 1.03) L5: 1.02 (0.99, 1.04) L0-2: 1.02 (0.99, 1.05)
<u>Ritz et al. (2006)</u>	Southern California	24-h avg: 21.9-22.1	Nearest Monitor	2 weeks before death: 1.03 (0.93, 1.14) 1 mo before death: NR 2 mo before death: 0.93 (0.89, 0.97) 6 mo before death: NR
<u>Hajat et al. (2007</u>)	10 Cities in the UK	24-h avg: 20.5-42.6	Citywide avg	L0-2: 1.00 (0.96, 1.06)
<u>Lin et al. (2004a</u>)	Sao Paulo, Brazil	24-h avg: 38.06	Citywide avg	L0: 1.00 (0.99, 1.01)
<u>Ha et al. (2003</u>)	Seoul, South Korea	8-h avg: 21.2	Citywide avg	L0: 0.93 (0.90, 0.96)
Romieu et al. (2004a)	Ciudad Juarez, Mexico	8-h avg: 43.43-55.12	Citywide avg	L1: 0.96 (0.90, 1.03) L2: 0.97 (0.91, 1.04) L0-1 cum: 0.96 (0.89, 1.04) L0-2 cum: 0.94 (0.87, 1.02)

Table 7-8Brief summary of infant mortality studies.

Study	Location	Mean O₃ (ppb)	Exposure Assessment	Effect Estimate ^a (95% CI):
Carbajal-Arroyo et al.	Mexico City,	1-h max:	Citywide avg	L0: 1.00 (0.99, 1.00)
<u>(2011</u>)	Mexico	103.0		L1: 0.99 (0.99, 0.99)
				L2: 0.99 (0.99, 1.00)
				L0-2: 0.99 (0.99, 1.00)
<u>Son et al. (2008</u>)	Seoul, South Korea	8-ha avg: 25.61	Citywide avg	L(NR): 0.984 (0.976, 0.992) ^b
<u>Tsai et al. (2006</u>)	Kaohsiung, Taiwan	24-h avg: 23.60	Citywide avg	L0-2 cum: 1.02 (0.56, 1.86)
Woodruff et al. (2008)	Nationwide, U.S.	24-h avg: 26.6	County wide avg	First 2 mo of life: 1.04 (0.98, 1.10)
<u>Yang et al. (2006</u>)	Taipei, Taiwan	24-h avg: 18.14	Citywide avg	L0-2 cum:1.00 (0.62, 1.61)
<u>Dales et al. (2004</u>)	12 Canadian	24-h: 31.77	Citywide avg	L0: NR
	cities			L1: NR
				L2: NR
				L3: NR
				L4: NR
				L5: NR
				Multiday lags of 2-6 days: NR

^aRelative risk of infant mortality per 10 ppb change in O_3

^bNo increment provided

L0 = Lag 0, L1= Lag 1, L2 = Lag 2, L3 = Lag 3, L4 = Lag 4, L5 = Lag 5, L6 = Lag 6

NR: No quantitative results reported

Table 7-9Summary of key reproductive and developmental toxicological
studies.

Study	Model	O ₃ (ppm)	Exposure Duration	Effects
<u>Sharkhuu et</u> <u>al. (2011</u>)	Pregnant mice; BALB/c; F; GD9-18; effects in offspring	0.4, 0.8, or 1.2	Continuously for 10 consecutive days	Dams: Decreased number of dams reaching parturition. Offspring: (1)-Decreased birth weights. (2)-Decreased rate of postnatal growth (body weight). (3)-impaired delayed type hypersensitivity.(4)-No effect on humoral immunity. (5)-Significantly affected allergic airway inflammation markers (eosinophilia, IgE) in female offspring sensitized early in life. 6-BALF LDH significantly elevated in female offspring.
<u>Bignami et</u> <u>al. (1994</u>)	Pregnant CD-1 dams (PD7-17)	0.4, 0.8 or 1.2	Continuous	Reproductive success was not affected by O_3 exposure (PD7- 17, proportion of successful pregnancies, litter size, ex ratio, frequency of still birth, or neonatal mortality). Ozone acted as a transient anorexigen in pregnant dams.
<u>Haro and</u> Paz (1993)	Rat dams, Exposure over the entirety of pregnancy;	1.0	12h/day during dark cycle	Decreased birth weight and postnatal body weight of offspring out to PND 90. Ozone-exposed pups manifested with inverted sleep-wake patterns or circadian rhythm phase-shift.
<u>López et al.</u> (2008)	Rats; Pregnant dams; GD1- GD18, GD20, or GD21.	1.0	(12 h/day, out to either GD18, GD20 or GD21)	O₃ induced delayed maturation of near term rodent bronchioles, with ultra-structural damage to bronchiolar epithelium.

Study	Model	O ₃ (ppm)	Exposure Duration	Effects
<u>Auten et al.</u> (2009)	C57BL/6 mouse pups	1.0	3 h/day, every other day, thrice weekly for 4 weeks	Postnatal O ₃ exposure significantly increased lung inflammatory cytokine levels; this was further exacerbated with gestational PM exposure.
Plopper et al. (2007)	Infant rhesus monkeys	0.5	Postnatal, PND30- 6month of age, 5 months of cyclic exposure, 5 days O ₃ followed by 9 days of filtered air, 8h/day.	Non-significant increases airway resistance and airway responsiveness with O_3 or inhaled allergen alone. Allergen + O_3 produced additive changes in both measures.
Fanucchi et al. (2006)	Infant male Rhesus monkeys, post- natal exposure	0.5	5 months of episodic exposure, age 1 month- age 6 months, 5 days O_3 followed by 9 days of filtered air, 8h/day.	Cellular changes and significant structural changes in the distal respiratory tract in infant rhesus monkeys exposed to O_3 postnatally.
<u>Dell'Omo et</u> <u>al. (1995</u>)	CD-1 Mouse dams and pups	0.6	6 days before breeding to weaning at PND21	Laterality changes in offspring: Ozone exposed pups showed a turning preference (left turns) distinct from air exposed controls (clockwise turns) as adults.
<u>Santucci et</u> <u>al. (2006</u>)	CD-1 Mouse dams	0.3 or 0.6	Dam exposure prior to mating through GD17.	Developmental O_3 caused increased defensive/submissive behavior in offspring. O_3 exposed offspring also had significant elevations of striatal BDNF and hippocampal NGF v. air exposed controls.
<u>Han et al.</u> (2011)	Rat; Sprague Dawley, M & F; PND13	0.6	3 h, BALF examined 10h after O_3 exposure	BALF polymorphonuclear leukocytes and total BALF protein were significantly elevated in O_3 exposed pups. Lung tissue from O_3 exposed pups had significant elevations of manganese superoxide dismutase (SOD) protein and significant decrements of extra-cellular SOD protein.
<u>Campos-</u> <u>Bedolla et al.</u> (2002)	Pregnant Rats; Sprague Dawley (GD5, GD10, or GD18)	3.0	1 h on one day of gestation, uteri collected 16-18 h later	Ozone inhalation modifies the contractile response of the pregnant uterus. The O_3 exposed pregnant uteri had significant increases in the maximum response to acetyl choline stimulation at GD5 and 10; they also had a significant increase in maximal response to oxytocin at GD 5.
<u>Kavlock et al.</u> (1980)	CD-1 mice; (pregnancy day 7-17)	0.4, 0.8 and 1.2	Continuous, pregnancy day 7-17	O_3 induced decrements in postnatal body weight gain. When O_3 was co-administered with sodium salicylate, O_3 synergistically increased the rate of pup resorption (1.0 ppm GD9-12).
<u>Jedlinska- Krakowska</u> <u>et al. (2006</u>)	5 month old male Wistar Hannover rats	3.0	0.5 ppm, 5h/day for 50 days	Histopathological evidence of impaired spermatogenesis (round spermatids/ 21 spermatocytes, giant spermatid cells, and focal epithelial desquamation with denudation to the 22 basement membrane). Vitamin E exposure concomitant with O_3 protected against pathological changes but Vitamin C did not.

7.4.11 Summary and Causal Determination

1	The 2006 O_3 AQCD concluded that the limited number of studies that investigated O_3
2	demonstrated no associations between O_3 and birth outcomes, with the possible exception
3	of birth defects. The current review included an expanded body of evidence on the
4	associations between O ₃ and reproductive and developmental effects. Recent
5	epidemiologic and toxicological studies provide evidence for an effect of prenatal
6	exposure to O_3 on pulmonary structure and function, including lung function changes in
7	the newborn, incident asthma, ultrastructural changes in bronchiole development,

1	
1	alterations in placental and pup cytokines, and increased pup airway hyper-reactivity.
2	Also, there is limited toxicological evidence for an effect of prenatal and early life
3	exposure on central nervous system effects, including laterality, brain morphology,
4	neurobehavioral abnormalities, and sleep aberration. Recent epidemiologic studies have
5	begun to explore the effects of O_3 on sperm quality, and provide limited evidence for
6	decrements in sperm concentration, while there is limited toxicological evidence for
7	testicular degeneration associated with O_3 .
8	While the collective evidence for many of the birth outcomes examined is generally
9	inconsistent (including birth defects), there are several well-designed, well-conducted
10	studies that indicate an association between O ₃ and adverse outcomes. For example, as
11	part of the southern California Children's Health Study, Salam et al. (2005) observed a
12	concentration-response relationship of decreasing birth weight with increasing O ₃
13	concentrations averaged over the entire pregnancy that was clearest above the 30-ppb
14	level (see Figure 7-4). Similarly, Hansen et al. (2008) utilized fetal ultrasonic
15	measurements and found a change in ultrasound measurements associated with O ₃ during
16	days 31-60 of gestation indicated that increasing O ₃ concentration decreased an
17	ultrasound measurement for women living within 2 km of the monitoring site.
18	The weight of evidence does not indicate that prenatal or early life O_3 concentrations are
19	associated with infant mortality. Collectively, there is limited though positive
20	toxicological evidence for O_3 -induced developmental effects, including effects on
21	pulmonary structure and function and central nervous system effects. Limited
22	epidemiologic evidence for an effect on prenatal O ₃ exposure on respiratory development
23	provides coherence with the effects observed in toxicological studies. There is also
24	limited epidemiologic evidence for an association with O_3 concentration and decreased
25	sperm concentration. A recent toxicological study provides limited evidence for a
26	possible biological mechanism (histopathology showing impaired spermatogenesis) for
27	such an association. Additionally, though the evidence for an association between O_3
28	concentrations and adverse birth outcomes is generally inconsistent, there are several
29	influential studies that indicate an association with reduced birth weight and restricted
30	fetal growth.
31	Some of the key challenges to interpretation of these study results include the difficulty in
32	assessing exposure as most studies use existing monitoring networks to estimate
33	individual exposure to ambient air pollution (see Section 4.6); the inability to control for
34	potential confounders such as other risk factors that affect birth outcomes (e.g., smoking);
35	evaluating the exposure window (e.g., trimester) of importance; integrating the results
36	from both short- and long-term exposure periods; integrating the results across a variety
50	from ooth short- and long-term exposure periods, integrating the results across a variety

 of reproductive and developmental outcomes; and limited evidence on the physiological mechanism of these effects.
 Taking into consideration the positive evidence for developmental and reproductive outcomes from toxicological and epidemiological studies, and the few influential birth outcome studies, the evidence is suggestive of a causal relationship between exposures to O₃ and reproductive and developmental effects.

7.5 Central Nervous System Effects

7.5.1 Effects on the Brain and Behavior

7 The 2006 O₃ AQCD included toxicological evidence that acute exposures to O₃ are 8 associated with alterations in neurotransmitters, motor activity, short and long term 9 memory, and sleep patterns. Additionally, histological signs of neurodegeneration have 10 been observed. Reports of headache, dizziness, and irritation of the nose with O₃ 11 exposure are common complaints in humans, and some behavioral changes in animals 12 may be related to these symptoms rather than indicative of neurotoxicity. Research in the 13 area of O₃-induced neurotoxicity has notably increased over the past few years, and 14 recent studies examining the effects of long-term exposure have demonstrated progressive damage in various regions of the brains of rodents in conjunction with altered 15 16 behavior. Evidence from epidemiologic studies has been more limited. A recently 17 published epidemiologic study examined the association between O_3 concentration and 18 neurobehavioral effects. Chen and Schwartz (2009) utilized data from the NHANES III 19 cohort to study the relationship between O_3 concentrations (mean annual O_3) 20 concentration 26.5 ppb) and neurobehavioral effects among adults aged 20-59 years. 21 Annual O_3 concentration was determined using inverse distance weighting for county of 22 residence and adjacent counties (for more information on inverse distance weighting and 23 other methods for exposure assessment, see Section 4.5.1 and 4.6). The authors observed an association between annual O₃ concentration and tests measuring coding ability 24 25 (symbol-digit substitution test) and attention/short-term memory (serial-digit learning 26 test). Each 10-ppb increase in annual O₃ concentration corresponded to an aging-related 27 cognitive performance decline of 3.5 yr for coding ability and 5.3 years for 28 attention/short-term memory. These associations persisted in both crude and adjusted 29 models. There was no association between O_3 concentration and reaction time tests. The 30 authors concluded that overall, there is an association between long-term O₃ 31 concentration and reduced performance on neurobehavioral tests.

1	A number of recent toxicological studies demonstrate various perturbations in neurologic
2	function or histology with long-term exposure to O ₃ , including changes similar to those
3	observed in neurodegenerative disorders such as Parkinson's and Alzheimer's disease
4	pathologies in relevant regions of the brain (Table 7-10). The central nervous system is
5	very sensitive to oxidative stress, due in part to its high content of polyunsaturated fatty
6	acids, high rate of oxygen consumption, and low antioxidant enzyme capacity. Oxidative
7	stress has been identified as one of the pathophysiological mechanisms underlying
8	neurodegenerative disease (Simonian and Coyle, 1996), and it is believed to play a role in
9	altering hippocampal function, which causes cognitive deficits with aging (Vanguilder
10	and Freeman, 2011). A particularly common finding in studies of O ₃ -exposed rats is lipid
11	peroxidation in the brain, especially in the hippocampus, which is important for higher
12	cognitive function including contextual memory acquisition. Performance in passive
13	avoidance learning tests is impaired when the hippocampus is injured. For example, in a
14	subchronic study, exposure of rats to 0.25 ppm O_3 (4 h/day) for 15-90 days caused a
15	complex array of responses, including a time-dependent increase in lipid peroxidation
16	products and immunohistochemical changes in the hippocampus that were correlated
17	with decrements in passive avoidance behavioral tests (Rivas-Arancibia et al., 2010).
18	Changes included increased numbers of activated microglia, a sign of inflammation, and
19	progressive neurodegeneration. Notably, continued exposure tends to bring about
20	progressive, cumulative damage, as shown by this study (Rivas-Arancibia et al., 2010)
21	and others (Santiago-López et al., 2010; Guevara-Guzmán et al., 2009; Angoa-Pérez et
22	<u>al., 2006</u>). The effects of O_3 on passive avoidance test performance were particularly
23	evident at 90 days for both short- and long-term memory. The greatest extent of cell loss
24	was also observed at this time point, whereas lipid peroxidation did not increase much
25	beyond 60 days of exposure.
26	The substantia nigra is another region of the brain affected by O ₃ , and seems particularly
27	sensitive to oxidative stress because the metabolism of dopamine, central to its function,
20	

2 28 is an oxidative process perturbed by redox imbalance. Oxidative stress has been 29 implicated in the premature death of substantia nigra dopamine neurons in Parkinson's 30 disease. Progressive damage has been found in the substantia nigra of male rats after 15, 31 30, and 60 days of exposure to 0.25 ppm O_3 for 4 h/day. Santiago-López et al. (2010) 32 observed a reduction dopaminergic neurons within the substantia nigra over time, with a 33 complete loss of normal morphology in the remaining cells and virtually no dopamine 34 immunoreactivity at 60 days. This was accompanied by an increase in p53 levels and 35 nuclear translocation, a process associated with programmed cell death. Similarly, 36 Angoa-Pérez et al. (2006) have shown progressive lipoperoxidation in the substantia 37 nigra and a decrease in nigral neurons in ovariectomized female rats exposed to 0.25 ppm 38 O₃, 4h/day, for 7 - 60 days. Lipid peroxidation effectively doubled between the 30 and 39 60 day time points. Total nigral cell number was also diminished to the greatest extent at

- 60 days, and cell loss was particularly evident in the tyrosine hydroxylase positive cell
 population (90%), indicating a selective loss of dopamine neurons or a loss of dopamine
 pathway functionality.
- 4 The olfactory bulb also undergoes oxidative damage in O₃-exposed animals, in some 5 cases altering olfactory-dependent behavior. Lipid peroxidation was observed in the 6 olfactory bulbs of ovariectomized female rats exposed to 0.25 ppm O_3 (4 h/day) for 30 or 7 60 days (Guevara-Guzmán et al., 2009). O₃ also induced decrements in a selective 8 olfactory recognition memory test, which were significantly greater at 60 days compared 9 to 30 days, and the authors note that early deficits in odor perception and memory are 10 components of human neurodegenerative diseases. The decrements in olfactory memory 11 did not appear to be due to damaged olfactory perception based on other tests early on, 12 but by 60 days deficits in olfactory perception had emerged.
- 13 Memory deficits and associated morphological changes can be attenuated by 14 administration of α -tocopherol (Guerrero et al., 1999), taurine (Rivas-Arancibia et al., 15 2000), and estradiol (Guevara-Guzmán et al., 2009; Angoa-Pérez et al., 2006), all of 16 which have antioxidant properties. In the study by Angoa-Pérez et al. (2006) described 17 above, estradiol seemed particularly effective at protecting against lipid peroxidation and 18 nigral cell loss at 60 days compared to shorter exposure durations. The same was true for 19 amelioration of decrements in olfactory recognition memory (Guevara-Guzmán et al., 20 2009), although protection against lipid peroxidation was similar for the 30 and 60 day 21 exposures.
- 22 CNS effects have also been demonstrated in adult mice whose only exposure to O₃ 23 occurred while in utero, a period particularly critical for brain development. Santucci et 24 al. (2006) investigated behavioral effects and gene expression after in utero exposure of 25 mice to 0.3 or 0.6 ppm O₃. Exposure began 30 days prior to mating and continued 26 throughout gestation. Testing of adult animals demonstrated increased 27 defensive/submissive behavior and reduced social investigation were observed in both the 28 0.3 and 0.6 ppm O₃ groups. Changes in gene expression of brain-derived neurotrophic 29 factor (BDNF, increased in striatum) and nerve growth factor (NGF, decreased in 30 hippocampus) accompanied these behavioral changes. BDNF and NGF are involved in 31 neuronal organization and the growth, maintenance, and survival of neurons during early 32 development and in adulthood. This study and two others using short-term exposures 33 demonstrate that CNS effects can occur as a result of in utero exposure to O₃, and 34 although the mode of action of these effects is not known, it has been suggested that 35 circulating lipid peroxidation products may play a role (Boussouar et al., 2009). 36 Importantly, these CNS effects occurred in rodent models after in utero only exposure to 37 (semi-) relevant concentrations of O₃.

Study	Model	O₃ (ppm)	Exposure Duration	Effects
<u>Angoa-Pérez et</u> <u>al. (2006</u>)	Rat; Wistar; F; Weight: 300 g; Ovariectomized	0.25	7 to 60 days, 4 h/day, 5 days/week	Long-term estradiol treatment protected against O ₃ -induced oxidative damage to nigral dopamine neurons, lipid peroxidation, and loss of tyrosine hydrolase-immunopositive cells.
<u>Guevara-</u> <u>Guzmán et al.</u> (2009)	Rat; Wistar; F; Weight: 264 g; Ovariectomized	0.25	30 and 60 days, 4h/day	Long-term estradiol treatment protected against O_3 -induced oxidative stress and decreases in α and β estrogen receptors and dopamine β -hydroxlyase in olfactory bulb, and deficits in olfactory social recognition memory and chocolate recognition.
<u>Rivas-Arancibia</u> <u>et al. (2010</u>)	Rat; Wistar; M; Weight: 250-300 g	0.25	15 to 90 days, 4h/day	Ozone produced significant increases in lipid peroxidation in the hippocampus, and altered the number of p53 positive immunoreactive cells, activated and phagocytic microglia, GFAP immunoreactive cells, double cortine cells, and short- and long-term memory- retention latency
<u>Santiago-López</u> <u>et al. (2010</u>)	Rat; Wistar; M; Weight: 250-300 g	0.25	15, 30, and 60 days, 4 h/day	Progressive loss of dopamine reactivity in the substantia nigra, along with morphological changes. Increased p53 levels and nuclear translocation.
<u>Santucci et al.</u> (2006)	Mice; CD-1; M; 18 weeks old	0.3; 0.6	Females continuously exposed from 30 days prior to breeding until GD17	Upon behavioral challenge with another male, there was a significant increase in defensive and freezing postures and decrease in the frequency of nose- sniffing. These behavioral changes were accompanied by a significant increase in BDNF in the striatum and a decrease of NGF in the hippocampus.

Table 7-10Central nervous system effects of long-term ozone exposure
in rats.

7.5.2 Summary and Causal Determination

1	The 2006 O_3 AQCD included toxicological evidence that acute exposures to O_3 are
2	associated with alterations in neurotransmitters, motor activity, short and long term
3	memory, and sleep patterns. Additionally, histological signs of neurodegeneration have
4	been observed. However, evidence regarding chronic exposure and neurobehavioral
5	effects was not available. Recent research in the area of O_3 -induced neurotoxicity has
6	included several long-term exposure studies. Notably, the first epidemiologic study to
7	examine the relationship between O_3 exposure and neurobehavioral effects observed an
8	association between annual O_3 levels and an aging-related cognitive performance decline
9	in tests measuring coding ability and attention/short-term memory. This observation is
10	supported by studies in rodents which demonstrate progressive oxidative stress and
11	damage in the brain and associated decrements in behavioral tests, including those
12	measuring memory, after subchronic exposure to 0.25 ppm O ₃ . Additionally,

1 2 3 neurobehavioral changes are evident in animals whose only exposure to O_3 occurred in utero. Collectively, the limited epidemiologic and toxicological evidence is coherent and suggestive of a causal relationship between O_3 exposure and CNS effects.

7.6 Carcinogenic and Genotoxic Potential of Ozone

7.6.1 Introduction

4	The radiomimetic and clastogenic qualities of O ₃ , combined with its ability to stimulate
5	proliferation of cells in the respiratory tract, have suggested that O_3 could act as a
6	carcinogen. However, toxicological studies of tumorigenesis in the rodent lung have
7	yielded mixed and often confusing results, and the epidemiologic evidence is equally
8	conflicted. The 2006 O_3 AQCD concluded that, "the weight of evidence from recent
9	animal toxicological studies and a very limited number of epidemiologic studies do not
10	support ambient O_3 as a pulmonary carcinogen" ¹ (<u>U.S. EPA, 2006b</u>).
11	Multiple epidemiologic studies reported in the 2006 O ₃ AQCD examined the association
12	between O_3 concentration and cancer. The largest of these studies, by <u>Pope et al. (2002</u>),
13	included 500,000 adults from the American Cancer Society's (ACS) Cancer Prevention II
14	study. In this study, no association was observed between O_3 concentration and lung
15	cancer mortality. The Adventist Health Study of Smog (AHSMOG) also examined the
16	association between O_3 concentration and lung cancer mortality (<u>Abbey et al., 1999</u>).
17	There was a positive association between O ₃ concentrations and lung cancer mortality
18	among men. No association was reported for women. Another study using the AHSMOG
19	cohort assessed the risk of incident lung cancer (Beeson et al., 1998). Among males, an
20	association with incidence of lung cancer was observed with increasing O_3
21	concentrations. When stratified by smoking status, the association persisted among never
22	smokers but was null for former smokers. No association was detected for females. The
23	Six Cities Study examined various air pollutants and mortality but did not specifically
24	explore the association between O3 concentrations and lung cancer mortality due to low
25	variability in O ₃ concentrations across the cities (Dockery et al., 1993). An ecologic study
26	performed in Sao Paulo City, Brazil examined the correlations between O3 concentrations
27	in four of the city districts and incident cancer of the larynx and lung reported in 1997
28	(<u>Pereira et al., 2005</u>). A correlation between the average number of days O_3
29	concentrations exceeded air quality standards from 1981 to 1990 and cancer incidence
30	was present for larynx cancer but not for lung cancer.

¹ The toxicological evidence is presented in detail in Table 6-18 on page 6-116 of the 1996 O_3 AQCD and Table AX5-13 on page AX5-43 of the 2006 O_3 AQCD.

1	Early toxicological research demonstrated lung adenoma ¹ acceleration in mice with daily
2	exposure to 1 ppm over 15 months (<u>Stokinger, 1962</u>). Later work demonstrated a
3	significant increase in lung tumor numbers in one strain of mouse (A/J) but not another
4	after exposure to 0.3-0.8 ppm O ₃ (Last et al., 1987; Hassett et al., 1985). The A/J mouse
5	strain is known to have a high incidence of spontaneous adenomas, and further studies
6	using this strain found a statistically significant increase in lung tumor incidence after a
7	9-month exposure to 0.5 ppm and incidence and multiplicity after a 5 month exposure to
8	0.12 ppm with a 4-month recovery period (<u>Witschi et al., 1999</u>). However, these findings
9	were discounted by the study authors due to the lack of a clear concentration-response,
10	and results from the Hassett et al. 1985 and Last et al. 1987 studies were retrospectively
10	deemed spurious based on what appeared to be unusually low spontaneous tumor
12	incidences in the control groups (<u>Witschi, 1991</u>). A study of carcinogenicity of O_3 by the
13	National Toxicology Program (NTP, 1994) reported increased incidences of
14	alveolar/bronchiolar adenoma or carcinoma (combined) in female B6C3F ₁ mice exposed
15	over 2 years to 1.0 ppm O_3 , but not 0.12 or .5 ppm. No effect was detected in male mice.
16	For a lifetime exposure to 0.5 or 1.0 ppm O_3 , an increase in the number of female mice
17	with adenomas (but not carcinomas or total neoplasms) was found. The number of total
18	neoplasms was also unaffected in male mice, but there was a marginally increased
19	incidence of carcinoma in males exposed to 0.5 and 1.0 ppm. Thus there was equivocal
20	evidence of carcinogenic activity in male mice and some evidence of carcinogenic
21	activity of O_3 in females. Experimental details of the NTP mouse study are available in
22	Table 6-19 on page 6-121 (U.S. EPA, 1996o) of the 1996 O_3 AQCD.
23	In Fischer-344/N rats (50 of each sex per group), neither a 2-year nor lifetime exposure to
24	O_3 ranging from 0.12 to 1.0 ppm was found to be carcinogenic (Boorman et al., 1994;
25	NTP, 1994). However, a marginally significant carcinogenic effect of 0.2 ppm O_3 was
26	reported in a study of male Sprague-Dawley rats exposed for 6 months ($n = 50$)
27	(Monchaux et al., 1996). These two studies also examined co-carcinogenicity of O_3 with
28	NNK^2 (Boorman et al., 1994) or a relatively high dose of radon (Monchaux et al., 1996),
29	finding no enhancement of NNK related tumors and a slight non-significant increase in
30	tumor incidence after combined exposure with radon, respectively. Another study
31	exploring co-carcinogenicity was conducted in hamsters. Not only was there no
32	enhancement of chemically induced tumors in the peripheral lung or nasal cavity, but
33	results suggested that O_3 could potentially delay or inhibit tumor development (Witschi et
55	results suggested that 03 could potentially delay of minor tunior development (Witsen et

results suggested that O_3 could potentially delay or inhibit tumor development (Witschi et al., 1993). Thus there is no concrete evidence that O_3 can act as a co-carcinogen.

34

¹ NOTE: Although adenomas are benign, over time they may progress to become malignant, at which point they are called adenocarcinomas. Adenocarcinoma is the predominant lung cancer subtype in most countries, and is the only lung cancer found in nonsmokers. From page 8-33 of the 1970 O_3 AQCD: "No true lung cancers have been reported, however, from experimental exposures to either O_3 alone or any other combination or ingredient of photochemical oxidants."

² 4-(N-nitrosomethylamino)-1-(3-pyridyl)-1-butanone

1	Immune surveillance is an important defense against cancer, and it should be noted that
2	natural killer (NK) cells, which destroy tumor cells in the lung, appear to be inhibited by
3	higher concentrations of O3 and either unaffected or stimulated at lower concentrations
4	(Section <u>6.2.5.4</u> , Infection and Adaptive Immunity). This aspect of tumorigenesis adds
5	yet another layer of complexity which may be reflected by conflicting results across
6	studies.
7	The following sections will examine epidemiologic studies of cancer incidence and
8	mortality and toxicological studies that have been published since the 2006 O_3 AQCD.
9	An epidemiologic study has been published with cancer as the outcome; most
10	epidemiologic studies examine markers of exposure.

7.6.2 Lung Cancer Incidence and Mortality

11	A recent re-analysis of the full ACS CPSII cohort by the Health Effects Institute is the
12	only epidemiologic study that has explored the association between O ₃ concentration and
13	cancer mortality since the last O ₃ AQCD. Krewski et al. (2009) conducted an extended
14	follow-up of the cohort (1982-2000). Mean O_3 concentration [obtained from the
15	Aerometric Information Retrieval System (AIRS) for 1980] were 22.91 ppb for the full
16	year and 30.15 ppb for the summer months (April-September). No association was
17	reported between lung cancer mortality and O_3 concentration (HR = 1.00 [95% CI:
18	0.96-1.04] per 10 ppb O ₃). Additionally, no association was observed when the analysis
19	was restricted to the summer months. There was also no association present in a sub-
20	analysis of the cohort examining the relationship between O_3 concentration and lung
21	cancer mortality in the Los Angeles area.
22	Since the 2006 O_3 AQCD, two toxicological studies have examined potential
22	Since the 2006 O_3 AQCD, two toxicological studies have examined potential
22 23	Since the 2006 O_3 AQCD, two toxicological studies have examined potential carcinogenicity of O_3 (<u>Kim and Cho, 2009a</u> , <u>b</u>). Looking across both studies, which used
22 23 24	Since the 2006 O_3 AQCD, two toxicological studies have examined potential carcinogenicity of O_3 (Kim and Cho, 2009a, b). Looking across both studies, which used the same mouse strain as the National Toxicology Program study described above (NTP,
22 23 24 25	Since the 2006 O_3 AQCD, two toxicological studies have examined potential carcinogenicity of O_3 (Kim and Cho, 2009a, b). Looking across both studies, which used the same mouse strain as the National Toxicology Program study described above (NTP, 1994), 0.5 ppm O_3 alone or in conjunction with chemical tumor inducers did not enhance
22 23 24 25 26	Since the 2006 O_3 AQCD, two toxicological studies have examined potential carcinogenicity of O_3 (Kim and Cho, 2009a, b). Looking across both studies, which used the same mouse strain as the National Toxicology Program study described above (NTP, 1994), 0.5 ppm O_3 alone or in conjunction with chemical tumor inducers did not enhance lung tumor incidence in males or females. However, a 10% incidence of oviductal
22 23 24 25 26 27	Since the 2006 O_3 AQCD, two toxicological studies have examined potential carcinogenicity of O_3 (Kim and Cho, 2009a, b). Looking across both studies, which used the same mouse strain as the National Toxicology Program study described above (NTP, 1994), 0.5 ppm O_3 alone or in conjunction with chemical tumor inducers did not enhance lung tumor incidence in males or females. However, a 10% incidence of oviductal carcinoma was observed in mice exposed to 0.5 ppm O_3 for 16 weeks. The implications
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7.6.3 DNA Damage

1	
1 2	The potential for genotoxic effects relating to O_3 exposure was predicted from the
2 3	radiomimetic properties of O_3 . The decomposition of O_3 in water produces OH and HO_2
	radicals, the same species that are generally considered to be the biologically active
4	products of ionizing radiation. Ozone has been observed to cause degradation of DNA in
5	a number of different models and bacterial strains. The toxic effects of O_3 have been
6	generally assumed to be confined to the tissues directly in contact with the gas, such as
7	the respiratory epithelium. Due to the highly reactive nature of O_3 , little systemic
8	absorption is predicted. Zelac et al. (1971a); (1971b), however, reported a significant
9	increase in chromosome aberrations in peripheral blood lymphocytes from Chinese
10	hamsters exposed to 0.2 ppm for 5 hours. Other in vivo exposure studies found increased
11	DNA strand breaks in respiratory cells from guinea pigs (Ferng et al., 1997) and mice
12	(Bornholdt et al., 2002) but only with exposure to higher concentrations of O_3 (1 ppm for
13	72 hours and 1 or 2 ppm for 90 minutes, respectively). In other studies there were no
14	observations of chromosomal aberrations in germ cells, but mutagenic effects have been
15	seen in offspring of mice exposed to 0.2 ppm during gestation (blepharophimosis or
16	dysplasia of the eyelids). The overall evidence for mutagenic activity from in vitro
17	studies is positive, and in the National Toxicology Program report described above, O ₃
18	was found to be mutagenic in Salmonella, with and without S9 metabolic activation. No
19	recent toxicological studies of DNA damage have become available since the 2006 O_3
20	AQCD.
21	A number of epidemiologic studies looked at the association between O_3 and DNA and
22	cellular level damages. These changes may be relevant to mechanisms leading to cancers
23	development and serve as early indicators of elevated risk of mutagenicity.
24	Two studies performed in California examined cytogenetic damage in relation to O_3
25	exposures. Huen et al. (2006) examined cytogenetic damage among African American
26	children and their mothers in Oakland, CA. Increased O_3 (mean monthly 8-h O_3
27	concentrations ranged from about 30 ppb in April to 14 ppb in November) was associated
28	with increased cytogenetic damage (micronuclei frequency among lymphocytes and
29	buccal cells) even after adjustment for household/personal smoking status and distance-
30	weighted traffic density. Chen et al. (2006a) recruited college students at the University
31	or California, Berkeley who reported never smoking and compared their levels of
32	cytogenetic damage (micronuclei frequency from buccal cells) in the spring and fall.
33	Cytogenetic damage was greater in the fall, which the authors attributed to the increase in
34	O ₃ over the summer. However, O ₃ levels over 2, 7, 10, 14, or 30 days (concentrations not
35	given) before collection of buccal cells did not correlate with cytogenetic damage.
36	Estimated lifetime O_3 exposure was also not correlated with cytogenetic damage.

1Additionally, the authors exposed a subset of the students (n = 15) to 200 ppb O3 for24 hours while the students exercised intermittently. Ozone was found to be associated3with an increase in cytogenetic damage in degenerated cells but not in normal cells49-10 days after exposure. Increased cytogenetic damage was also noted in peripheral5blood lymphocytes collected 18 hours after exposure.

6 A study performed in Mexico recruited 55 male workers working indoors (n = 27) or 7 outdoors (n = 28) in Mexico City or Puebla, Mexico in order to study the relationship 8 between O₃ and DNA damage (detected from peripheral blood samples using the Comet 9 assay) (Tovalin et al., 2006). The median estimated daily O_3 concentrations were 10 estimated to be 28.5 ppb for outdoor workers and 5.1 ppb for indoor workers in Mexico City and 36.1 ppb for outdoor workers and 19.5 ppb for indoor workers in 11 12 Puebla. Overall, a positive correlation between O₃ levels and DNA damage was 13 observed. However, when examining the relationship by city and workplace, only DNA 14 damage in outdoor workers in Mexico City remained correlated with O₃ levels.

15 Three studies examining the relationship between O₃ concentration and DNA-level 16 damage have been performed in Europe. The largest of these studies was the GenAir 17 case-control study, which was nested within the European Prospective Investigation into 18 Cancer and Nutrition (EPIC) study, and included individuals recruited between 1993 and 19 1998 from ten European countries. Only non-smokers (must not have smoked for at least 20 10 years prior to enrollment) were enrolled in the study. The researchers examined DNA 21 adduct levels (DNA bonded to cancer-causing chemicals) and their relationship with O_3 22 concentrations (concentrations not given) (Peluso et al., 2005). A positive association was 23 seen between DNA adduct levels and O₃ concentrations from 1990-1994 but not O₃ 24 concentrations from 1995-1999. In adjusted analyses with DNA adduct levels 25 dichotomized as high and low (detectable versus non-detectable), the OR was 1.97 26 (95% CI: 1.08, 3.58) when comparing the upper tertile of O_3 concentration to the lower 27 two tertiles. Two other European studies were conducted in Florence, Italy. The most 28 recent of these enrolled individuals from the EPIC study into a separate study between 29 March and September of 1999 (Palli et al., 2009). The purpose of the study was to 30 examine oxidative DNA damage (determined by Comet assay using blood lymphocytes) 31 in association with varying periods of O_3 exposure. The researchers observed that longer 32 periods of high O₃ concentrations (values not given) were more strongly correlated with 33 oxidative DNA damage than shorter periods of time (i.e., the rho [p-value] was 0.26 34 [0.03] for 0-10 days and 0.35 [0.002] for 0-90 days). This correlation was stronger among 35 men compared to women. The correlations for all time periods had p-values <0.05 for ex-36 and never-smokers. For current smokers, the correlation was only observed among time 37 periods \leq 25 days. When adjusted for age, gender, smoking history, traffic pollution 38 exposure, period of blood draw, and area of residence, the association between O_3

1	concentrations and oxidative DNA damage was positive for O_3 concentrations 0-60 days,
2	0-75 days, and 0-90 days prior to blood draw. Positive, statistically significant
3	associations were not observed among shorter time periods. The other study performed in
4	Florence recruited healthy volunteers who reported being non-smokers or light smokers
5	(Giovannelli et al., 2006). The estimated O_3 concentrations during the study ranged from
6	approximately 4-40 ppb for 3-day averages, 5-35 ppb for 7-day averages, and
7	7.5-32.5 ppb for 30-day averages. Ozone concentrations were correlated with DNA
8	strand breaks (measured from blood lymphocytes) over longer exposure periods (p-value:
9	0.002 at 30 days, p-value: 0.04 at 7 days; p-value: 0.17 at 3 days). This association was
10	robust to control for temperature, solar radiation, gender, and age. No association was
11	seen between O_3 concentrations and measures of oxidative DNA damage at 3, 7, or
12	30 days.

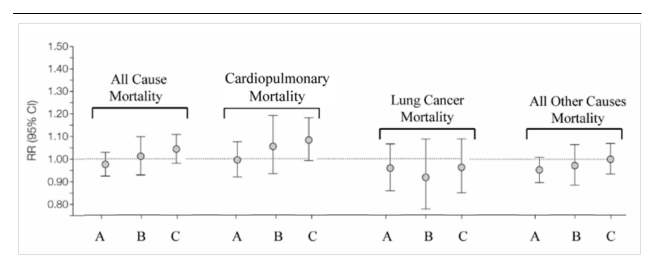
7.6.4 Summary and Causal Determination

13	The 2006 O_3 AQCD reported that evidence did not support ambient O_3 as a pulmonary
14	carcinogen. Since the 2006 O ₃ AQCD, very few epidemiologic and toxicological studies
15	have been published that examine O_3 as a carcinogen, but collectively, study results
16	indicate that O3 may contribute to DNA damage. O3 concentrations in most
17	epidemiologic studies were measured using air monitoring data. For more information on
18	long-term exposure assessment, see Section 4.6.3.2 Overall, the evidence is inadequate
19	to determine if a causal relationship exists between ambient O_3 exposures and
20	cancer.

7.7 Mortality

21	A limited number of epidemiologic studies have assessed the relationship between long-
22	term exposure to O_3 and mortality in adults. The 2006 O_3 AQCD concluded that an
23	insufficient amount of evidence existed "to suggest a causal relationship between chronic
24	O_3 exposure and increased risk for mortality in humans" (U.S. EPA, 2006b). In addition
25	to the infant mortality studies discussed in Section 7.4.10, additional studies have been
26	conducted among adults since the last review; an ecologic study that finds no association
27	between mortality and O ₃ , several re-analyses of the ACS cohort, one of which
28	specifically points to a relationship between long-term O ₃ exposure and an increased risk
29	of respiratory mortality, and a study of four cohorts of persons with potentially
30	predisposing conditions. These studies supplement the evidence from long-term cohort
31	studies characterized in previous reviews of O ₃ , and are summarized here briefly.

1	In the Harvard Six Cities Study (Dockery et al., 1993), adjusted mortality rate ratios were
2	examined in relation to long-term mean O ₃ concentrations in six cities: Topeka, KS; St.
3	Louis, MO; Portage, WI; Harriman, TN; Steubenville, OH; and Watertown, MA. Mean
4	O3 concentrations from 1977 to 1985 ranged from 19.7 ppb in Watertown to 28.0 ppb in
5	Portage. Long-term mean O ₃ concentrations were not found to be associated with
6	mortality in the six cities. However, the authors noted that "The small differences in O_3
7	levels among the (six) cities limited the power of the study to detect associations between
8	mortality and O_3 levels." In addition, while total and cardio-pulmonary mortality were
9	considered in this study, respiratory mortality was not specifically considered.
10	In a subsequent large prospective cohort study of approximately 500,000 U.S. adults,
11	Pope et al. (2002) examined the effects of long-term exposure to air pollutants on
12	mortality (American Cancer Society, Cancer Prevention Study II). All-cause,
13	cardiopulmonary, lung cancer and other mortality risk estimates for long-term O_3
14	exposure are shown in Figure 7-5. While consistently positive associations were not
15	observed between O_3 and mortality (effect estimates labeled A in Figure 7-5), the
16	mortality risk estimates were larger in magnitude when analyses considered more
17	accurate exposure metrics, increasing when the entire period was considered (effect
18	estimates labeled B in Figure 7-5) and becoming marginally significant when the
19	exposure estimate was restricted to the summer months (July to September; effect
20	estimates labeled C in Figure 7-5), especially when considering cardiopulmonary deaths.
21	In contrast, consistent positive and significant effects of PM _{2.5} were observed for both
22	lung cancer and cardio-pulmonary mortality.



	Years of Data Collection	Number of Metropolitan Areas	Number of Participants (in thousands)	1-h max O₃ Mean (SD)
Α	1980-1981	134	559	47.9 (11.0)
В	1982-1998	119	525	45.5 (7.3)
С	1982-1998 (July – Sept)	134	557	59.7 (12.8)

Source: Reprinted with permission of American Medical Association Pope et al. (2002).

Figure 7-5 Adjusted ozone-mortality relative risk estimates (95% CI) by time period of analysis per subject-weighted mean ozone concentration in the Cancer Prevention Study II by the American Cancer Society.

1	A study by Abbey et al. (1999) examined the effects of long-term air pollution exposure,
2	including O_3 , on all-cause (n = 1,575), cardiopulmonary (n = 1,029), nonmalignant
3	respiratory ($n = 410$), and lung cancer ($n = 30$) mortality in the long-term prospective
4	Adventist Health Study of Smog (AHSMOG) of 6,338 nonsmoking, non-Hispanic white
5	individuals living in California. A particular strength of this study was the extensive
6	effort devoted to assessing long-term air pollution exposures, including interpolation to
7	residential and work locations from monitoring sites over time and space. No associations
8	with long-term O_3 exposure were observed for all cause, cardiopulmonary, and
9	nonmalignant respiratory mortality. In a follow-up, Chen et al. (2005) utilized data from
10	the AHSMOG study and reported no evidence of associations between long-term O_3
11	exposure (mean O ₃ concentration 26.2 ppb) and fatal coronary heart disease. Thus, no
12	association of chronic O ₃ exposure with mortality outcomes has been detected in this
13	study.
14	Lipfert et al. (2003); (2000) reported positive effects on all-cause mortality for peak O_3
15	exposures (95th percentile levels) in the U.S. Veterans Cohort study of approximately

16 50,000 middle-aged men recruited with a diagnosis of hypertension. The actual analysis

1	involved smaller subcohorts based on exposure and mortality follow-up periods. Four
2	separate exposure periods were associated with three mortality follow-up periods. For
3	
	concurrent exposure periods, peak O ₃ was positively associated with all-cause mortality,
4	with a 9.4% (95% CI: 0.4, 18.4) excess risk per mean 95th percentile O_3 less estimated
5	background level (not stated). "Peak" refers, in this case, to the 95th percentile of
6	the hourly measurements, averaged by year and county. In a further analysis, Lipfert et al.
7	(2003) reported the strongest positive association for concurrent exposure to peak O_3 for
8	the subset of subjects with low diastolic blood pressure during the 1982 to 1988 period.
9	Two more recent studies of this cohort focused specifically on traffic density (Lipfert et
10	al., 2006a; 2006b). Lipfert et al. (2006b) concluded that: "Traffic density is seen to be a
11	significant and robust predictor of survival in this cohort, more so than ambient air
12	quality, with the possible exception of O_3 ," reporting a significant O_3 effect even with
13	traffic density included in the model: $RR = 1.080$ per 40 ppb peak O ₃ (95% CI: 1.019,
14	1.146). However, in Lipfert et al. (2006a), which considers only the EPA Speciation
15	Trends Network (STN) sites, O3 drops to non-significant predictor of total mortality for
16	this cohort. The authors acknowledge that: "Peak O ₃ has been important in analyses of
17	this cohort for previous periods, but in the STN data set, this variable has limited range
18	and somewhat lower values and its small coefficient of variation results in a relatively
19	large standard error." The restriction to subjects near STN sites likely reduced the power
20	of this analysis, though the size of the remaining subjects considered was not reported in
21	this paper. In addition, these various Veterans Cohort studies considered only total
22	mortality, and did not consider mortality on a by-cause basis.

- 23 An ecological study in Brisbane, Australia used a geospatial approach to analyze the 24 association of long-term exposure to gaseous air pollution with cardio-respiratory 25 mortality, in the period 1996-2004 (Wang et al., 2009c). A generalized estimating 26 equations model was employed to investigate the impact of NO_2 , O_3 and SO_2 , but PM 27 was not addressed. The results indicated that long-term exposure to O_3 was not associated 28 with cardio-respiratory mortality, but the fact that this study considered only one city, and 29 that the range of O_3 exposure across that city (23.7-35.6 ppb) was low and slight in 30 variation in comparison to the range of other pollutants across the city, limited study 31 power. In addition, confounding factors (e.g., smoking) could not be addressed at the 32 individual level in this ecological study. Respiratory mortality was not evaluated 33 separately.
- 34A recent study by Zanobetti and Schwartz examined whether year-to-year variations in358-h mean daily O3 concentrations for the summer (May-September) around their city-36specific long-term trend were associated with year-to-year variations in mortality around37its long-term trend. This association was examined among Medicare participants with38potentially predisposing conditions, including COPD, diabetes, CHF, and MI, defined as

1	patients discharged alive after an emergency admission for one of these four conditions.
2	The analyses was repeated in 105 cities using available data from 1985 through 2006, and
3	the results were combined using methods previously employed by these authors
4	(Zanobetti et al., 2008; Zanobetti and Schwartz, 2007). This study design eliminated
5	potential confounding by factors that vary across city, which is a common concern in
6	most air pollution cohort studies, and also avoided both confounding by cross-sectional
7	factors that vary by city and the short-term factors that confound daily time-series studies,
8	but are not present in annual analyses. The average 8-h mean daily summer O_3
9	concentrations ranged from 15.6 ppb (Honolulu, HI) to 71.4 ppb (Bakersfield, CA) for
10	the 105 cities. The authors observed associations between yearly fluctuations in summer
11	O ₃ concentrations and mortality in each of the four cohorts; the hazard ratios (per 10 ppb
12	increment) were 1.12 (95% CI: 1.06, 1.17) for the CHF cohort, 1.19 (95% CI 1.12, 1.25)
13	for the MI cohort, 1.14 (95% CI: 1.10, 1.21) for the diabetes cohort, and 1.14 (95% CI:
14	1.08, 1.19) for the COPD cohort. A key advantage to this study is that fluctuations from
15	summer to summer in O ₃ concentrations around long-term level and trend in a specific
16	city are unlikely to be correlated with most other predicators of mortality risk; except for
17	temperature, which was controlled for in the regression. Key limitations of the study were
18	the inability to control for $PM_{2.5}$, since it was not reliably measured in these cities until
19	1999, and the inability to separate specific causes of death (e.g., respiratory,
20	cardiovascular), since Medicare does not provide the underlying cause of death.
21	In the most recent follow-up analyses of the ACS cohort (Jerrett et al., 2009; Smith et al.,
22	<u>2009a</u>), the effects of long-term exposure to O_3 were evaluated alone, as well as in
23	copollutant models with $PM_{2.5}$ and components of $PM_{2.5}$. Jerrett et al. (2009) utilized the
24	ACS cohort with data from 1977 through 2000 (mean O_3 concentration ranged from 33.3
25	to 104.0 ppb) and subdivided cardiopulmonary deaths into respiratory and cardiovascular,
26	separately, as opposed to combined into one category, as was done by Pope et al. (2002).
27	Increases in exposure to O_3 were associated with an elevated risk of death from
28	cardiopulmonary, cardiovascular, ischemic heart disease, and respiratory causes.
29	Consistent with study hypotheses, inclusion of PM2.5 concentrations measured in
30	1999-2000 (the earliest years for which it was available) as a copollutant attenuated the
31	association with O_3 for all end points except death from respiratory causes, for which a
32	significant association persisted (Table 7-11). The association between increased O_3
33	concentrations and increased risk of death from respiratory causes was insensitive to the
34	use of a random-effects survival model allowing for spatial clustering within the
35	metropolitan area and state of residence, and adjustment for several ecologic variables
36	considered individually. Subgroup analyses showed that temperature and region of
37	country, but not sex, age at enrollment, body-mass index, education, or $PM_{2.5}$
38	concentration, modified the effects of O_3 on the risk of death from respiratory causes
39	(i.e., risks were higher at higher temperature, and in the Southeast, Southwest, and Upper
57	(nei, fisks were ingher at ingher temperature, and in the boutheast, bouth est, and epper

1	Midwest). Ozone threshold analyses indicated that the threshold model was not a better
2	fit to the data (p >0.05) than a linear representation of the overall O_3 -mortality
3	association. Overall, this new analysis indicates that long-term exposure to $PM_{2.5}$
4	increases risk of cardiac death, while long-term exposure to O3 is specifically associated
5	with an increased risk of respiratory death, and suggests that combining cardiovascular
6	and respiratory causes of mortality into one category for analysis may obscure any effect
7	that O ₃ may have on respiratory-related causes of mortality.

Table 7-11Relative risk (and 95% CI) of death attributable to a 10-ppb change
in the ambient ozone concentration.

Cause of Death	O ₃ (96 MSAs) ^a	O ₃ (86 MSAs) ^a	O ₃ +PM _{2.5} (86 MSAs) ^a
Any Cause	1.001 (0.996, 1.007)	1.001 (0.996, 1.007)	0.989 (0.981, 0.996)
Cardiopulmonary	1.014 (1.007, 1.022)	1.016 (1.008, 1.024)	0.992 (0.982, 1.003)
Respiratory	1.029 (1.010, 1.048)	1.027 (1.007, 1.046)	1.040 (1.013, 1.067)
Cardiovascular	1.011 (1.003, 1.023)	1.014 (1.005, 1.023)	0.983 (0.971, 0.994)
Ischemic Heart Disease	1.015 (1.003, 1.026)	1.017 (1.006, 1.029)	0.973 (0.958, 0.988)

^aOzone concentrations were measured from April to September during the years from 1977 to 2000, with follow-up from 1982 to 2000; changes in the concentration of $PM_{2.5}$ of 10 µg/m³ were recorded for members of the cohort in 1999 and 2000. Source: Reprinted with permission of Massachusetts Medical Society (Jerrett et al., 2009).

8 In a similar analysis, Smith et al. (2009a) used data from 66 Metropolitan Statistical 9 Areas (MSAs) in the ACS cohort to examine the association of O_3 concentrations during 10 the warm season and all-cause and cardiopulmonary mortality. Mortality effects were 11 estimated in single pollutant and copollutant models, adjusting for two PM_{2.5} constituents, 12 sulfate, and EC. When all-cause mortality was investigated, there was a 0.8% (95% CI: 13 -0.31, 1.9) increase associated with a 10 ppb increase in O₃ concentration. This 14 association was diminished when sulfate or EC were included in the model. There was a 15 2.48% (95% CI: 0.74, 4.3) increase in cardiopulmonary mortality associated with a 16 10 ppb increase in O_3 concentration. The cardiopulmonary association was robust to 17 adjustment for sulfate, and diminished, though still positive, after adjustment for EC 18 (1.63% increase; 95% CI: -0.41, 3.7). Smith et al. (2009a) did not specifically separate 19 out cardiovascular and respiratory causes of death from the cardiopulmonary category, as 20 was done by Jerrett et al. (2009).

7.7.1 Summary and Causal Determination

21The The 2006 O3 AQCD concluded that an insufficient amount of evidence existed "to22suggest a causal relationship between chronic O3 exposure and increased risk for

- 1 mortality in humans" (U.S. EPA, 2006b). Several additional studies have been conducted 2 since the last review that evaluate cause-specific and total mortality. An ecologic study 3 conducted in Australia observed no association between cardiopulmonary mortality and 4 O_3 (Wang et al., 2009c). Two reanalyses of the ACS cohort were conducted; one 5 provides weak evidence for an association with cardiopulmonary mortality (Smith et al., 6 2009a) while the other specifically points to a relationship between long-term O_3 7 exposure and an increased risk of respiratory mortality (Jerrett et al., 2009). Most 8 recently, a study of four cohorts of Medicare enrollees with potentially predisposing 9 conditions observed associations between O₃ and total mortality among each of the 10 cohorts (Zanobetti and Schwartz, 2011).
- 11 When considering the entire body of evidence, there is limited support for an association 12 with long-term exposure to ambient O_3 and total mortality. There is inconsistent evidence 13 for an association between long-term exposure to ambient O_3 and cardiopulmonary 14 mortality, with several analyses from the ACS cohort reporting some positive 15 associations (Smith et al., 2009a; Pope et al., 2002) while other studies reported no 16 association (Wang et al., 2009c; Abbey et al., 1999; Dockery et al., 1993). The strongest 17 evidence for an association between long-term exposure to ambient O_3 concentrations 18 and mortality is derived from associations reported in the Jerrett et al. (2009) study for 19 respiratory mortality that remained robust after adjusting for PM_{2.5} concentrations. 20 Finally, a recent analysis reported associations of ambient O₃ concentrations and total 21 mortality in potentially at-risk populations in the Medicare Cohort (Zanobetti and 22 Schwartz, 2011), while earlier studies generally report no associations with total 23 mortality (Lipfert et al., 2006a; Lipfert et al., 2003; Pope et al., 2002; Abbey et al., 1999; 24 Dockery et al., 1993). Studies of cardiopulmonary and total mortality provide limited 25 evidence for an association with long-term exposure to ambient O_3 concentrations. The 26 study by Jerrett et al. (2009) observes an association between long-term exposure to 27 ambient O_3 concentrations and respiratory mortality remained robust after adjusting for 28 PM_{2.5} concentrations. Coherence and biological plausibility for this observation is 29 provided by evidence from epidemiologic, controlled human exposure, and animal 30 toxicological studies for the effects of short- and long-term exposure to O_3 on respiratory 31 effects (See Sections 6.2 and 7.2). Respiratory mortality is a relatively small portion of 32 total mortality [about 7.6% of all deaths in 2010 were due to respiratory causes (Murphy 33 et al., 2012), thus it is not surprising that the respiratory mortality signal may be difficult 34 to detect in studies of cardiopulmonary or total mortality. Based on the recent evidence 35 for respiratory mortality along with limited evidence for total and cardiopulmonary 36 mortality, the evidence is suggestive of a causal relationship between long-term O_3 37 exposures and total mortality.

7.8 Overall Summary

1 2 3 The evidence reviewed in this chapter describes the recent findings regarding the health effects of long-term exposure to ambient O_3 concentrations. <u>Table 7-12</u> provides an overview of the causal determinations for each of the health categories evaluated.

Table 7-12Summary of causal determinations for long-term exposures to
ozone.

Health Category	Causal Determination
Respiratory Effects	Likely to be a causal relationship
Cardiovascular Effects	Suggestive of a causal relationship
Reproductive and Developmental Effects	Suggestive of a causal relationship
Central Nervous System Effects	Suggestive of a causal relationship
Carcinogenicity and Genotoxicity	Inadequate to infer a causal relationship
Total Mortality	Suggestive of a causal relationship

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8 POPULATIONS POTENTIALLY AT INCREASED RISK FOR OZONE-RELATED HEALTH EFFECTS

1 Interindividual variation in human responses to air pollution exposure can result in some 2 groups being at increased risk for detrimental effects in response to ambient exposure to 3 an air pollutant. The NAAOS are intended to provide an adequate margin of safety for 4 both the population as a whole and those potentially at increased risk for health effects in 5 response to ambient air pollution (Preface to this ISA). To facilitate the identification of 6 populations and lifestages at greater risk for air pollutant related health effects, studies 7 have evaluated factors that may contribute to the susceptibility and/or vulnerability of an 8 individual to air pollutants. The definitions of susceptibility and vulnerability have been 9 found to vary across studies, but in most instances "susceptibility" refers to biological or 10 intrinsic factors (e.g., lifestage, sex, preexisting disease/conditions) while "vulnerability" 11 refers to non-biological or extrinsic factors (e.g., socioeconomic status [SES]) (U.S. EPA, 12 2010c, 2009d). In some cases, the terms "at-risk" and "sensitive" populations have been 13 used to encompass these concepts more generally. The main goal of this evaluation is to 14 identify and understand those factors that result in a population or lifestage being at 15 increased risk of an air pollutant-related health effect, not to categorize the factors. To 16 this end, previous ISAs and reviews (Sacks et al., 2011; U.S. EPA, 2010c, 2009d) have 17 used "susceptible populations" to encompass these various factors. In this chapter, 18 "at-risk" is the all-encompassing term used for groups with specific factors that increase 19 the risk of an air pollutant (e.g., O₃)-related health effects in a population.

20 Individuals, and ultimately populations, could experience increased risk for air pollutant 21 induced health effects via multiple avenues. A group with intrinsically increased risk 22 would have some factor(s) that increases risk for an effect through a biological 23 mechanism. In general, people in this category would have a steeper concentration-24 risk relationship, compared to those not in the category. Potential factors that are often 25 considered intrinsic include genetic background and sex. A group of people could also 26 have extrinsically increased risk, which would be through an external, non-biological 27 factor. Examples of extrinsic factors include SES and diet. Some groups are at risk of 28 increased internal dose at a given exposure concentration, which includes individuals 29 that have a greater dose of delivered pollutant because of breathing pattern. This 30 category would include persons who work outdoors or exercise outdoors. In addition, 31 some outdoor workers could have greater exposure (concentration x time), regardless 32 of the delivered dose. Finally, there are those who might be placed at increased risk 33 for experiencing a greater exposure by being exposed at a higher concentration. For

1	example, groups of people exposed to higher air pollutant concentrations due to less
2	availability/use of home air conditioners (i.e., more open windows on high O_3 days).
3	Some factors described above are multifaceted and may influence the risk of an air
4	pollutant related health effect through a combination of avenues. For example, SES may
5	affect access to medical care, which itself may contribute to the presence of preexisting
6	diseases and conditions considered as intrinsic factors. Additionally, children tend to
7	spend more time outdoors at higher levels of activity than adults, which leads to
8	increased intake dose and exposure, but they also have biological (i.e., intrinsic)
9	differences when compared to adults.
10	The emphasis of this chapter is to identify and understand the factors that potentially
11	increase the risk of O ₃ -related health effects, regardless of whether the increased risk is
12	due to intrinsic factors, extrinsic factors, increased dose/exposure or a combination, due
13	to the often connected pathways between factors. The following sections examine factors
14	that potentially lead to increased risk of O ₃ -related health effects and characterize the
15	overall weight of evidence for each factor. Most of the factors are related to greater health
16	effects given a specific dose but there is also discussion of increased internal dose and/or
17	exposure at a given concentration integrated throughout the sections (i.e., lifestage,
18	outdoor workers, and air conditioning use).

Approach to Classifying Potential At-Risk Factors

19 To identify factors that potentially lead to some populations being at greater risk to air 20 pollutant related health effects, the evidence across relevant scientific disciplines 21 (i.e., exposure sciences, dosimetry, controlled human exposure, toxicology, and 22 epidemiology) was evaluated. In this systematic approach, the collective evidence is used 23 to examine coherence of effects across disciplines and determine biological plausibility. 24 By first focusing on studies (i.e., epidemiologic or controlled human exposure) that 25 conduct stratified analyses it is possible to identify factors that may result in some 26 populations being at greater risk of an air pollutant related health effect. These types of 27 studies allow for an evaluation of populations exposed to similar air pollutant (e.g., O_3) 28 concentrations within the same study design. Experimental studies also provide important 29 lines of evidence in the evaluation of factors that may lead to increased risk of an air 30 pollutant related-health effect. Toxicological studies conducted using animal models of 31 disease and controlled human exposure studies that examine individuals with underlying 32 disease or genetic polymorphisms may provide evidence in the absence of stratified 33 epidemiologic analyses. Additionally these studies can provide support for coherence 34 with the health effects observed in epidemiologic studies as well as an understanding of 35 biological plausibility. The collective results across the scientific disciplines comprise the

1	overall weight of evidence that is used to determine whether a specific factor results in a
2	population being at increased risk of an air pollutant related health effect.
3	Building on the causal framework discussed in detail in the Preamble and used
4	throughout the ISA, conclusions are made regarding the strength of evidence for each
5	factor that may contribute to increased risk of an O_3 -related health effect based on the
6	evaluation and synthesis of evidence across scientific disciplines. The conclusions drawn
7	considered the "Aspects to Aid in Judging Causality" discussed in Table 1 of the
8	Preamble. The categories considered for evaluating the potential increased risk of an air
9	pollutant-related health effect are "adequate evidence," "suggestive evidence,"
10	"inadequate evidence," and "evidence of no effect." They are described in more detail in
11	<u>Table 8-1</u> .

Table 8-1 Classification of Evidence for Potential At-Risk Factors.

	Health Effects
Adequate evidence	There is substantial, consistent evidence within a discipline to conclude that a factor results in a population or lifestage being at increased risk of air pollutant-related health effect(s) relative to some reference population or lifestage. Where applicable this includes coherence across disciplines. Evidence includes multiple high-quality studies.
Suggestive evidence	The collective evidence suggests that a factor results in a population or lifestage being at increased risk of an air pollutant-related health effect relative to some reference population or lifestage, but the evidence is limited due to some inconsistency within a discipline or, where applicable, a lack of coherence across disciplines.
Inadequate evidence	The collective evidence is inadequate to determine if a factor results in a population or lifestage being at increased risk of an air pollutant-related health effect relative to some reference population or lifestage. The available studies are of insufficient quantity, quality, consistency and/or statistical power to permit a conclusion to be drawn.
Evidence of no effect	There is substantial, consistent evidence within a discipline to conclude that a factor does not result in a population or lifestage being at increased risk of air pollutant-related health effect(s) relative to some reference population or lifestage. Where applicable this includes coherence across disciplines. Evidence includes multiple high-quality studies.

12This chapter evaluates the various factors indicated in the literature that may result in a13population being at increased risk of an O_3 -related health effect. For further detail on the14epidemiologic, controlled human exposure, and toxicological studies included in this15chapter, see Chapters 5, 6, and 7.

8.1 Genetic Factors

16	The potential effects of air pollution on individuals with specific genetic characteristics
17	have been examined; studies often target polymorphisms in already identified candidate
18	susceptibility genes or in genes whose protein products are thought to be involved in the
19	biological mechanism underlying the health effect of an air pollutant (Sacks et al., 2011).
20	As a result, multiple studies that examined the effect of short- and long-term O_3 exposure
21	on respiratory function have focused on whether various gene profiles lead to an

1	increased risk of O_3 -related health effects. For more details on the function and mode of
2	action of the genetic factors discussed in this section, see Section $5.4.2.1$. Additionally, a
3	limited number of toxicological studies have examined the joint effects of nutrition and
4	genetics. Details on these toxicological studies of nutrition and genetics can be found in
5	Section <u>5.4.2.3</u> .
6	Multiple genes, including glutathione S-transferase Mu 1 (GSTM1) and tumor necrosis
7	factor- α (TNF- α) were evaluated in the 2006 O ₃ AQCD and found to have a "potential
8	role in the innate susceptibility to O_3 " (U.S. EPA, 2006b). Epidemiologic, controlled
9	human exposure, and toxicological studies performed since the 2006 O ₃ AQCD have
10	continued to examine the roles of GSTM1 and TNF- α in modifying O ₃ -related health
11	effects and have examined other gene variants that may also increase risk. Due to small
12	sample sizes, many controlled human exposure studies are limited in their ability to test
13	genes with low frequency minor alleles and therefore, some genes important for
14	O3-related health effects may not have been examined in these types of studies. A
15	summary of effect measure modification findings from epidemiologic and controlled
16	human exposure studies discussed in this section is included as <u>Table 8-2</u> .

Table 8-2Summaries of results from epidemiologic and controlled human
exposures studies of modification by genetic variants.

Gene variant	Comparison group	Health outcome /population	Effect modification of association for the gene variant	Reference
GSTM1 null	GSTM1 positive	Respiratory symptoms among asthmatic children	↑	Romieu et al. (2006)
GSTP1 Val/Val	GSTP1 lle/lle or lle/Val	Respiratory symptoms among asthmatic children	↑	_
GSTP1 IIe/IIe or IIe/Val	GSTP1 Val/Val	Lung function among asthmatic children	\downarrow	
GSTP1 Ile/Val or Val/Val	GSTP1 lle/lle	Lung function among adults	\downarrow	Alexeeff et al. (2008)
HMOX1 S/L or L/L	HMOX1 S/S	Lung function among adults	\downarrow	
NQO1 wildtype and GSTM1 null	Other combinations	Lung function among healthy adults with exercise	\downarrow	<u>Bergamaschi et al.</u> (2001)
NQO1 wildtype and GSTM1 null	Other combinations	Lung function among mild-to- moderate asthmatics with moderate exercise	=	Vagaggini et al. (2010
NQO1 wildtype and GSTM1 null	Other combinations	Inflammatory responses among mild-to-moderate asthmatics with moderate exercise	=	
GSTM1 null	GSTM1 positive	Lung function among healthy adults with intermittent moderate exercise	=	<u>Kim et al. (2011</u>)
GSTM1 null	GSTM1 positive	Inflammatory responses among healthy adults with intermittent moderate exercise	=	
GSTM1 null	GSTM1 positive	Lung function among asthmatic children	\downarrow	Romieu et al. (2004b)
GSTM1 null	GSTM1 positive	Lung function among healthy adults with intermittent moderate exercise	=	<u>Alexis et al. (2009</u>)
GSTM1 null	GSTM1 positive	Inflammatory changes among healthy adults with intermittent moderate exercise	↑	-

1 Epidemiologic studies that examined the effects of short-term exposure to O_3 on lung 2 function included analyses of potential gene-environment interactions. Romieu et al. 3 (2006) reported an association between O₃ and respiratory symptoms that were larger 4 among children with GSTM1 null or glutathione S-transferase P 1 (GSTP1) Val/Val 5 genotypes compared with children with GSTM1 positive or GSTP1 Ile/Ile or Ile/Val 6 genotypes, respectively. However, results suggested that O₃-associated decreases in lung 7 function may be greater among children with GSTP1 Ile/Ile or Ile/Val compared to 8 GSTP1 Val/Val. Alexeeff et al. (2008) reported greater O₃-related decreases in lung 9 function among GSTP1 Val/Val adults than those with GSTP1 Ile/Ile or GSTP1 Ile/Val 10 genotypes. In addition, they detected greater O₃-associated decreases in lung function for 11 adults with long GT dinucleotide repeats in heme-oxygenase-1 (HMOX1) promoters.

1	Several controlled human exposure studies have reported that genetic polymorphisms of
2	antioxidant enzymes may modulate pulmonary function and inflammatory responses to
3	O ₃ challenge. Healthy carriers of NAD(P)H quinone oxidoreductase 1 (NQO1) wild type
4	(wt) in combination with GSTM1 null genotype had greater decreases in lung function
5	parameters with exposure to O_3 (Bergamaschi et al., 2001). Vagaggini et al. (2010)
6	exposed mild-to-moderate asthmatics to O3 during moderate exercise. In subjects with
7	NQO1 wt and GSTM1 null, there was no evidence of changes in lung function or
8	inflammatory responses to O ₃ . Kim et al. (2011) also recently conducted a study among
9	young adults, about half of whom were GSTM1-null and half of whom were
10	GSTM1-sufficient. They detected no difference in the FEV ₁ responses to O ₃ exposure by
11	GSTM1 genotype and did not examine NQO1. In another study that examined GSTM1
12	but not NQO1, asthmatic children with GSTM1 null genotype (Romieu et al., 2004b)
13	were reported to have greater decreases in lung function in relation to O ₃ exposure.
14	Additionally, supplementation with antioxidants (Vitamins C and E) had a slightly more
15	beneficial effect among GSTM1 null children (for more on modification by diet, see
16	Section <u>8.4.1</u>).
17	
17	In a study of healthy volunteers with GSTM1 sufficient (n = 19; 24 \pm 3) and GSTM1 null
18	$(n = 16; 25 \pm 5)$ genotypes exposed to 400 ppb O ₃ for 2 hours with exercise, <u>Alexis et al.</u>
19	(2009) found genotype effects on inflammatory responses but not lung function responses
20	to O_3 . At 4 hours post- O_3 exposure, individuals with either GSTM1 genotype had
21	statistically significant increases in sputum neutrophils with a tendency for a greater
22	increase in GSTM1 sufficient than GSTM1 nulls. At 24 hours postexposure, neutrophils
23	had returned to baseline levels in the GSTM1 sufficient individuals. In the GSTM1 null
24	subjects, neutrophil levels increased from 4 to 24 hours and were significantly greater
25	than both baseline levels and levels at 24 hours in the GSTM1 sufficient individuals. In
26	addition, O ₃ exposure increased the expression of the surface marker CD14 in airway
27	neutrophils of GSTM1 null subjects compared with GSTM1 sufficient subjects. CD14
28	and TLR4 are co-receptors for endotoxin, and signaling through this innate immune
29	pathway has been shown to be important for a number of biological responses to O_3
30	exposure in toxicological studies (Garantziotis et al., 2010; Hollingsworth et al., 2010;
31	Hollingsworth et al., 2004; Kleeberger et al., 2000). Alexis et al. (2009) also
32	demonstrated decreased numbers of airway macrophages at 4 and 24 hours following O_3
33	exposure in GSTM1 sufficient subjects. Airway macrophages in GSTM1 null subjects
34	were greater in number and found to have greater oxidative burst and phagocytic
35	capability following O_3 exposure than those of GSTM1 sufficient subjects. Airway
36	macrophages and dendritic cells from GSTM1 null subjects exposed to O3 expressed
37	higher levels of the surface marker HLA-DR, again suggesting activation of the innate
38	immune system. Since there was no FA control in the Alexis et al. (2009) study, effects
39	of the exposure other than O_3 cannot be ruled out. In general, the findings between these

1studies are inconsistent. It is possible that different genes may be important for different2phenotypes. Additional studies, which include appropriate controls, are needed to clarify3the influence of genetic polymorphisms on O3 responsiveness in humans.

- 4 In general, toxicological studies have reported differences in cardiac and respiratory 5 effects after O₃ exposure among different mouse strains, which alludes to differential risk 6 among individuals due to genetic variability (Tankersley et al., 2010; Chuang et al., 2009; 7 Hamade and Tankersley, 2009; Hamade et al., 2008). Thus strains of mice which are 8 prone to or resistant to O_3 -induced effects have been used to systematically identify 9 candidate genes that may increase risk of O₃-related health effects. Genome wide linkage 10 analyses have identified quantitative trait loci for O₃-induced lung inflammation and 11 hyperpermeability on chromosome 17 (Kleeberger et al., 1997) and chromosome 4 12 (Kleeberger et al., 2000), respectively, using recombinant inbred strains of mice. More 13 specifically, these studies found that TNF (protein product is the inflammatory cytokine 14 TNF- α) and Tlr4 (protein product is TLR4, involved in endotoxin responses) were 15 candidate susceptibility genes (Kleeberger et al., 2000; Kleeberger et al., 1997). The TNF 16 receptors 1 and 2 have also been found to play a role in injury, inflammation, and airway 17 hyperreactivity in studies of O_3 -exposed knockout mice (Cho et al., 2001). In addition to 18 Tlr4, other innate immune pattern recognition signaling pathway genes, including Tlr2 19 and Myd88, appear to be important in responses to O_3 , as demonstrated by Williams et al. 20 (2007b). A role for the inflammatory cytokine IL-6 has been demonstrated in 21 gene-deficient mice with respect to inflammation and injury, but not AHR (Johnston et 22 al., 2005; Yu et al., 2002). Mice deficient in IL-10, an anti-inflammatory cytokine, 23 demonstrated increased pulmonary inflammation in response to O_3 exposure (Backus et 24 al., 2010). Thus genes related to innate immune signaling and pro- and anti-inflammatory 25 genes are important for O₃-induced responses.
- 26 Altered O_3 responses between mouse strains could be due to genetic variability in nuclear 27 factor erythroid 2-related factor 2 (Nrf-2), suggesting a role for genetic differences in 28 altering the formation of ROS (Hamade et al., 2010; Cho and Kleeberger, 2007). 29 Additionally, some studies have reported O₃-related effects to vary by Inf-1 and Inf-2 30 quantitative trait loci (Tankersley and Kleeberger, 1994) and a gene coding for Clara cell 31 secretory protein (CCSP) (Broeckaert et al., 2003; Wattiez et al., 2003). Other 32 investigations in inbred mouse strains found that differences in expression of certain 33 proteins, such as CCSP (Broeckaert et al., 2003) and MARCO (Dahl et al., 2007), are 34 responsible for phenotypic characteristics, such as epithelial permeability and scavenging 35 of oxidized lipids, respectively, which confer sensitivity to O₃.
- 36 Nitric oxide (NO), derived from activated macrophages, is produced upon exposure to O₃
 37 and is thought to participate in lung damage. Mice deficient in the gene for inducible

- 1 nitric oxide synthase (NOS2/NOSII/iNOS) are partially protected against lung injury 2 (Kleeberger et al., 2001), and it appears that O_3 -induced iNOS expression is tied to the 3 TLR4 pathway described above. Similarly, iNOS deficient mice do not produce reactive 4 nitrogen intermediates after O_3 exposure, in contrast to their wild-type counterparts, and 5 also produce less PGE2 comparatively (Fakhrzadeh et al., 2002). These gene-deficient 6 mice were protected from O_3 -induced lung injury and inflammation. In contrast, another 7 study using a similar exposure concentration but longer duration of exposure found that 8 iNOS deficient mice were more susceptible to O₃-induced lung damage (Kenyon et al., 9 2002). Therefore it is unclear whether inducible nitric oxide synthase plays a protective 10 role or mediates damage.
- 11 Voynow et al. (2009) have shown that NQO1 deficient mice, like their human 12 counterparts, are resistant to O₃-induced AHR and inflammation. NQO1 catalyzes the 13 reduction of quinones to hydroquinones, and is capable of both protective detoxification 14 reactions and redox cycling reactions resulting in the generation of reactive oxygen 15 species. Reduced production of inflammatory mediators and cells and blunted AHR were 16 observed in NQO1 null mice after exposure to 1 ppm O₃ for 3 hours. These results 17 correlated with those from in vitro experiments in which human bronchial epithelial cells 18 treated with an NQO1 inhibitor exhibited reduced inflammatory responses to exposure to 19 0.4 ppm O₃ for 5 hours. This study may provide biological plausibility for the increased 20 biomarkers of oxidative stress and increased pulmonary function decrements observed in 21 O₃-exposed individuals bearing both the wild-type NQO1 gene and the null GSTM1 gene 22 (Bergamaschi et al., 2001).
- 23 The role of TNF- α signaling in O₃-induced responses has been previously established 24 through depletion experiments, but a more recent toxicological study investigated the 25 effects of combined O₃ and PM exposure in transgenic TNF overexpressing mice. 26 Kumarathasan et al. (2005) found that subtle effects of these pollutants were difficult to 27 identify in the midst of the severe pathological changes caused by constitutive TNF- α 28 overexpression. However, there was evidence that TNF transgenic mice were more 29 susceptible to O₃/PM-induced oxidative stress, and they exhibited elevation of a serum 30 creatine kinase after pollutant exposure, which may suggest potential systemic or cardiac 31 related effects. Differential susceptibility to O_3 among inbred strains of animals does not seem to be dose dependent since absorption of ¹⁸O in various strains of mice did not 32 33 correlate with resistance or sensitivity (Vancza et al., 2009).
- 34Defects in DNA repair mechanisms may also confer increased risk of O3-related health35effects. Cockayne syndrome, a rare autosomal recessive disorder in humans, is36characterized by UV sensitivity abnormalities, neurological abnormalities, and premature37aging. The same genetic defect in mice (Csb^{-/-}) makes them sensitive to oxidative

1stressors, including O_3 . Kooter et al. (2007) demonstrated that $Csb^{-/-}$ mice product2significantly more TNF- α after exposure to 0.8 ppm O_3 than their wild-type condition3However, there were no statistically significant differences in other markers of4inflammation or lung injury between the two strains of mice.	unterparts.
5 Overall, for variants in multiple genes there is suggestive evidence for potentia	l
6 involvement in populations being potentially more at-risk than others to the eff	ects of O ₃
7 exposure on health. Controlled human exposure and epidemiologic studies hav	e reported
8 some evidence of O ₃ -related increases in respiratory symptoms or decreases in	lung
9 function with variants including GSTM1, GSTP1, HMOX1 and NQO1, although	gh the
10 results are not consistent across studies and gene variants. Future studies of the	se and
11 other genes in human populations will be important for determining the role of	each
12 genotype and its effect on risk as well as finding coherence across the disciplin	es. NQO1
13 deficient mice were found to be resistant to O ₃ -induced AHR and inflammation	l,
14 providing biological plausibility for results of studies in humans. Additionally,	studies of
15 rodents have identified a number of other genes that may affect O ₃ -related heal	th
16 outcomes, including genes related to innate immune signaling and pro- and	
17 anti-inflammatory genes, which have not been investigated in human studies.	

8.2 Preexisting Disease/Conditions

18	Individuals with certain preexisting diseases are likely to constitute an at-risk population.
19	This may be the result of individuals with a preexisting disease/condition having less
20	reserve than healthy individuals, so although the absolute change may be the same, the
21	health consequences are different. Previous O3 AQCDs concluded that some people with
22	preexisting pulmonary disease, especially asthma, are among those at increased risk of an
23	O_3 -related health effect. Extensive toxicological evidence indicates that altered
24	physiological, morphological and biochemical states typical of respiratory diseases may
25	render people at risk of an additional oxidative burden induced by O_3 exposure. In
26	addition, a number of epidemiologic studies found that some individuals with respiratory
27	diseases are at increased risk of O3-related effects. The majority of the studies identified
28	in previous AQCDs focused on whether preexisting respiratory diseases result in
29	increased risk of O ₃ -related health effects, with a limited number of studies examining
30	other preexisting diseases, such as cardiovascular.
31	Studies identified since the completion of the 2006 O_3 AQCD that examined whether
32	preexisting diseases and conditions lead to increased risk of O_3 -induced health effects
33	were identified and are summarized below. <u>Table 8-3</u> displays the prevalence rates of

some of these conditions categorized by age and region among adults in the U.S.

34

1	population; data for children, when available, are presented within the following sections.
2	Substantial proportions of the U.S. population are affected by these conditions and
3	therefore may represent a potentially large at-risk population. While these diseases and
4	conditions represent biological or intrinsic factors that could lead to increased risk, the
5	pathways to their development may have intrinsic or extrinsic origins.

Table 8-3Prevalence of respiratory diseases, cardiovascular diseases, and
diabetes among adults by age and region in the U.S.

				Adul	ts				
	N (in thousands)		Α	ge			Regi	on	
Chronic Disease/Condition		18-44	45-64	65-74	75+	Northeast	Midwest	South	West
Respiratory Diseases									
Asthma ^a	16,380	7.2	7.5	7.8	6.4	7.7	8.0	5.9	8.4
COPD									
Chronic Bronchitis	9,832	3.2	5.5	5.9	5.3	3.4	4.8	5.2	2.9
Emphysema	3,789	0.2	2.0	5.7	5.0	1.2	1.9	1.9	1.3
Cardiovascular Diseases									
All Heart Disease	26,628	4.6	12.3	26.7	39.2	11.3	12.7	12.2	9.9
Coronary Heart Disease	14,428	1.1	6.7	16.9	26.7	5.7	6.5	7.3	4.9
Hypertension	56,159	8.7	32.5	54.4	61.1	22.9	24.1	27.1	20.6
Diabetes	18,651	2.3	12.1	20.4	17.3	4.5	7.6	9.0	7.7

^aAsthma prevalence is reported for "still has asthma."

Source: Pleis et al. (2009); National Center for Health Statistics.

8.2.1 Influenza/Infections

6	Recent studies have indicated that underlying infections may increase the risk of O ₃ -
7	related health effects because O_3 exposure likely impairs host defenses, which may
8	increase the body's response to an infectious agent. However, there is little epidemiologic
9	or experimental evidence that infection or influenza itself renders an individual at greater
10	risk of an O ₃ -induced health effect. A study of hospitalizations in Hong Kong reported
11	that increased levels of influenza intensity resulted in increased excess risk of respiratory
12	disease hospitalizations related to O_3 exposure (<u>Wong et al., 2009</u>). In addition, a study of
13	lung function in asthmatic children reported decreases in lung function with increased
14	short-term O_3 exposure for those with upper respiratory infections but not for those
15	without infections (Lewis et al., 2005). Toxicological studies provide biological
16	plausibility for the increase in O ₃ -induced health effects observed in epidemiologic
17	studies that examined infections by way of studies that demonstrated that exposure to

1	0.08 ppm O_3 increased streptococcus-induced mortality, regardless of whether O_3
2	exposure preceded or followed infection (Miller et al., 1978; Coffin and Gardner, 1972;
3	Coffin et al., 1967). Overall, the epidemiologic and experimental evidence supports the
4	potential for increased risk to be conferred by an infection but the number of studies is
5	limited. There have only been a few epidemiologic studies and these studies examine
6	different outcomes (respiratory-related hospital admissions or lung function) and
7	different modifiers (influenza or respiratory infection). In some of the toxicological
8	studies, the O_3 exposure came before the infection. Therefore, evidence is inadequate to
9	determine if influenza/infections increase the risk of O3-related health effects.

8.2.2 Asthma

10	Previous O ₃ AQCDs identified individuals with asthma as a population at increased risk
11	of O ₃ -related health effects. Within the U.S., approximately 7.3% of adults have reported
12	currently having asthma (Pleis et al., 2009), and 9.5% of children have reported currently
13	having asthma (<u>Bloom et al., 2008</u>). For more detailed prevalence by age, see <u>Table 8-4</u> .

Table 8-4Prevalence of asthma by age in the U.S.

Age (years)	N (in thousands)	Percent
0-4	1,276	6.2
5-11	3,159	11.2
12-17	2,518	10.2
18-44	7,949	7.2
45-64	5,768	7.5
65-74	1,548	7.8
75+	1,116	6.4

^aAsthma prevalence is reported for "still has asthma"

Source: Statistics for adults: Pleis et al. (2009); statistics for children: Bloom et al. (2008); National Center for Health Statistics.

14	Multiple epidemiologic studies included within this ISA have evaluated the potential for
15	increased risk of O3-related health effects among individuals with asthma. A study of
16	lifeguards in Texas reported decreased lung function with short-term O3 exposure among
17	both individuals with and without asthma, however, the decrease was greater among
18	those with asthma (Thaller et al., 2008). A Mexican study of children ages 6-14 detected
19	an association between short-term O_3 exposure and wheeze, cough, and bronchodilator
20	use among asthmatics but not non-asthmatics, although this may have been the result of a
21	small non-asthmatic population (Escamilla-Nuñez et al., 2008). A study of modification

- 1 by airway hyperresponsiveness (AHR) (a condition common among asthmatics) reported 2 greater short-term O₃-associated decreases in lung function in elderly individuals with 3 AHR, especially among those who were obese (Alexeeff et al., 2007). However, no 4 evidence for increased risk was found in a study performed among children in Mexico 5 City that examined the effect of short-term O_3 exposure on respiratory health (Barraza-6 Villarreal et al., 2008). In this study, a positive association was reported for airway 7 inflammation among asthmatic children, but the observed association was similar in 8 magnitude to that of non-asthmatics. Similarly, a study of children in California reported 9 an association between O₃ concentration and exhaled nitric oxide fraction (FeNO) that 10 persisted both among children with and without asthma as well as those with and without 11 respiratory allergy (Berhane et al., 2011). Finally, Khatri et al. (2009) found no 12 association between short-term O₃ exposure and altered lung function for either asthmatic 13 or non-asthmatic adults, but did note a decrease in lung function among individuals with 14 allergies.
- 15 Evidence for difference in effects among asthmatics has been observed in studies that 16 examined the association between O_3 exposure and altered lung function by asthma 17 medication use. A study of children with asthma living in Detroit reported a greater association between short-term O₃ and lung function for corticosteroid users compared 18 19 with noncorticosteroid users (Lewis et al., 2005). Conversely, another study found 20 decreased lung function among noncorticosteroid users compared to corticosteroid users, 21 although in this study, a large proportion of non-users were considered to be persistent 22 asthmatics (Hernández-Cadena et al., 2009). Lung function was not related to short-term 23 O_3 exposure among corticosteroid users and non-users in a study taking place during the 24 winter months in Canada (Liu et al., 2009a). Additionally, a study of airway 25 inflammation reported a counterintuitive inverse association with O_3 of similar magnitude 26 for all groups of corticosteroid users and non-users (Qian et al., 2009).
- 27 Controlled human exposure studies that have examined the effects of O_3 on individuals 28 with asthma and healthy controls are limited. Based on studies reviewed in the 1996 and 29 $2006 O_3 AOCDs$, subjects with asthma appeared to be at least as sensitive to acute effects 30 of O_3 in terms of FEV₁ and inflammatory responses as healthy non-asthmatic subjects. 31 For instance, Horstman et al. (1995) observed that mild-to-moderate asthmatics, on 32 average, experienced double the O_3 -induced FEV₁ decrement of healthy subjects (19%) 33 versus 10%, respectively, p = 0.04). Moreover, a statistically significant positive 34 correlation between FEV_1 responses to O_3 exposure and baseline lung function was 35 observed in individuals with asthma, i.e., responses increased with severity of disease. 36 Kreit et al. (1989) performed a short duration study in which asthmatics also showed a 37 considerable larger average O_3 -induced FEV₁ decrement than the healthy controls (25% 38 vs. 16%, respectively) following exposure to O_3 with moderate-heavy exercise. Alexis et

1	al. (2000) and Jorres et al. (1996) also reported a tendency for slightly greater FEV ₁
2	decrements in asthmatics than healthy subjects. Minimal evidence exists suggesting that
3	individuals with asthma have smaller O_3 -induced FEV ₁ decrements than healthy subjects
4	(3% versus 8%, respectively) (Mudway et al., 2001). However, the asthmatics in that
5	study also tended to be older than the healthy subjects, which could partially explain their
5 6	
	lesser response since FEV_1 responses to O_3 exposure diminish with age. Individuals with
7	asthma also had more neutrophils in the BALF (18 hours postexposure) than similarly
8	exposed healthy individuals (Peden et al., 1997; Scannell et al., 1996; Basha et al., 1994).
9	Furthermore, a study examining the effects of O ₃ on individuals with atopic asthma and
10	healthy controls reported that greater numbers of neutrophils, higher levels of cytokines
11	and hyaluronan, and greater expression of macrophage cell-surface markers were
12	observed in induced sputum of atopic asthmatics compared with healthy controls
13	(<u>Hernandez et al., 2010</u>). Differences in O ₃ -induced epithelial cytokine expression were
14	noted in bronchial biopsy samples from asthmatics and healthy controls (Bosson et al.,
15	2003). Cell-surface marker and cytokine expression results, and the presence of
16	hyaluronan, are consistent with O_3 having greater effects on innate and adaptive
17	immunity in these asthmatic individuals (see Section <u>5.4.2.2</u>). In addition, studies have
18	demonstrated that O ₃ exposure leads to increased bronchial reactivity to inhaled allergens
19	in mild allergic asthmatics (Kehrl et al., 1999; Jorres et al., 1996) and to the influx of
20	eosinophils in individuals with pre-existing allergic disease (Vagaggini et al., 2002;
21	Peden et al., 1995). Taken together, these results point to several mechanistic pathways
22	which could account for the increased risk of O_3 -related health effects in subjects with
23	asthma (see Section $5.4.2.2$).
24	Toxicological studies provide biological plausibility for greater effects of O_3 among those
25	with asthma or AHR. In animal toxicological studies, an asthmatic phenotype is modeled

2 26 by allergic sensitization of the respiratory tract. Many of the studies that provide evidence 27 that O3 exposure is an inducer of AHR and remodeling utilize these types of animal 28 models. For example, a series of experiments in infant rhesus monkeys have shown these 29 effects, but only in monkeys sensitized to house dust mite allergen (Fanucchi et al., 2006; 30 Joad et al., 2006; Schelegle et al., 2003). Similarly, Funabashi et al. (2004) demonstrated 31 changes in pulmonary function in mice exposed to O_3 , and Wagner et al. (2007) 32 demonstrated enhanced inflammatory responses in rats exposed to O₃, but only in 33 animals sensitized to allergen. In general, it is the combined effects of O_3 and allergic 34 sensitization which result in measurable effects on pulmonary function. In a bleomycin 35 induced pulmonary fibrosis model, exposure to 250 ppb O₃ for 5 days increased 36 pulmonary inflammation and fibrosis, along with the frequency of bronchopneumonia in 37 rats. Thus, short-term exposure to O₃ may enhance damage in a previously injured lung 38 (Oyarzún et al., 2005).

1	In the 2006 O_3 AQCD, the potential for individuals with asthma to have greater risk of
2	O ₃ -related health effects was supported by a number of controlled human exposure
3	studies, evidence from toxicological studies, and a limited number of epidemiologic
4	studies. Overall, in the recent epidemiologic literature some, but not all, studies report
5	greater risk of health effects among individuals with asthma. Studies examining effect
6	measure modification of the relationship between short-term O_3 exposure and altered
7	lung function by corticosteroid use provided limited and inconsistent evidence of
8	O3-related health effects. Additionally, recent studies of behavioral responses have found
9	that studies do not take into account individual behavioral adaptations to forecasted air
10	pollution levels (such as avoidance and reduced time outdoors), which may underestimate
11	the observed associations in studies that examined the effect of O_3 exposure on
12	respiratory health (Neidell and Kinney, 2010). This could explain some inconsistency
13	observed among recent epidemiologic studies. The evidence from controlled human
14	exposure studies provides support for increased decrements in FEV_1 and greater
15	inflammatory responses to O_3 in individuals with asthma than in healthy individuals
16	without a history of asthma. The collective evidence for increased risk of O3-related
17	health effects among individuals with asthma from controlled human exposure studies is
18	supported by recent toxicological studies which provide biological plausibility for
19	heightened risk of asthmatics to respiratory effects due to O3 exposure. Evidence
20	indicating O ₃ -induced respiratory effects among individuals with asthma is further
21	supported by additional studies of O_3 -related respiratory effects (Section <u>6.2</u>). Overall,
22	there is adequate evidence for asthmatics to be a potentially at-risk population based on
23	the substantial, consistent evidence among controlled human exposure studies and
24	coherence from epidemiologic and toxicological studies.

8.2.3 Chronic Obstructive Pulmonary Disease (COPD)

25	In the U.S. over 4% of adults report having chronic bronchitis and almost 2% report
26	having emphysema, both of which are classified as COPD (Pleis et al., 2009).
27	A recent study reported no association between O ₃ exposure and lung function regardless
28	of whether the study participant had COPD or other preexisting diseases (asthma or IHD)
29	(<u>Lagorio et al., 2006</u>).
30	Peel et al. (2007) found that individuals with COPD were at increased risk of
31	cardiovascular ED visits in response to short-term O3 exposure compared to healthy
32	individuals in Atlanta, GA. The authors reported that short-term O ₃ exposure was
33	associated with higher odds of an emergency department (ED) visit for peripheral and
34	cerebrovascular disease among individuals with COPD compared to individuals without

1	COPD. However, preexisting COPD did not increase the odds of hospitalization for all
2	CVD outcomes (i.e., IHD, dysrhythmia, or congestive heart failure). In an additional
3	study performed in Taiwan, individuals with and without COPD had higher odds of
4	congestive heart failure associated with O ₃ exposure on warm days (Lee et al., 2008a). As
5	discussed in Section <u>6.3</u> , most studies reported no overall association between O_3
6	concentration and CVD morbidity.
7	Recent epidemiologic evidence indicates that persons with COPD may have increased
8	risk of O ₃ -related cardiovascular effects, but little information is available on whether
9	COPD leads to an increased risk of O ₃ -induced respiratory effects. Overall, this small
10	number of studies provides inadequate evidence to determine whether COPD results in
11	increased risk of O ₃ -related health effects.

8.2.4 Cardiovascular Disease (CVD)

12 Cardiovascular disease has become increasingly prevalent in the U.S., with about 12% of 13 adults reporting a diagnosis of heart disease (Table 8-3). A high prevalence of other 14 cardiovascular-related conditions has also been observed, such as hypertension which is 15 prevalent among approximately 24% of adults. In the 2006 O₃ AQCD, little evidence was 16 available regarding whether preexisting CVD contributed to increased risk of O₃-related 17 health effects. Recent epidemiologic studies have examined cardiovascular-related 18 diseases as modifiers of the O₃-outcome associations; however, no recent evidence is 19 available from controlled human exposure studies or toxicological studies.

20 Peel et al. (2007) compared the associations between short-term O_3 exposure and 21 cardiovascular ED visits in Atlanta, GA among multiple comorbid conditions. The 22 authors found no evidence of increased risk of cardiovascular ED visits in individuals 23 previously diagnosed with dysrhythmia, congestive heart failure, or hypertension 24 compared to healthy individuals. Similarly, a study in France examined the association 25 between O_3 concentrations and ischemic cerebrovascular events (ICVE) and myocardial 26 infarction (MI) and the influence of multiple vascular risk factors on any observed 27 associations (Henrotin et al., 2010). The association between O_3 exposure and ICVE was 28 elevated for individuals with multiple risk factors, specifically individuals with diabetes 29 or hypertension. For the association between O_3 and MI, increased odds were apparent 30 only for those with hypercholesterolemia. In a study conducted in Taiwan, a positive 31 association was observed for O_3 on warm days and congestive heart failure hospital 32 admissions, but the association did not differ between individuals with/without 33 hypertension or with/without dysrhythmia (Lee et al., 2008a). Another study in Taiwan 34 reported that the association between O₃ levels and ED visits for arrhythmias were greater 1on warm days among those with congestive heart failure compared to those without2congestive heart failure; however, the estimate and 95% CIs for those without congestive3heart failure is completely contained within the 95% CI of those with congestive heart4failure (Chiu and Yang, 2009).

5 Although not studied extensively, a study has examined the increased risk of O₃-related 6 changes in blood markers for individuals with CVD. There was a greater association 7 between O₃ exposure and some, but not all, blood inflammatory markers among 8 individuals with a history of CVD (Liao et al., 2005). Liao et al. (2005) found that 9 increased fibrinogen was positively associated with short-term O_3 exposure but this 10 association was present only among individuals with a history of CVD. No association 11 was observed among those without a history of CVD. However, for another biomarker 12 (vWF), CVD status did not modify the positive association with short-term O_3 exposure 13 (Liao et al., 2005).

14 Mortality studies provide some evidence for a potential increase in O₃-induced mortality 15 in individuals with preexisting atrial fibrillation and atherosclerosis. In a study of 48 U.S. 16 cities, increased risk of mortality with short-term O_3 exposure was observed only among 17 individuals with secondary atrial fibrillation (Medina-Ramón and Schwartz, 2008). No 18 association was observed for short-term O₃ exposure and mortality in a study of 19 individuals with diabetes with or without CVD prior to death; however, there was some 20 evidence of increased risk of mortality during the warm season if individuals had diabetes 21 and atherosclerosis compared to only having diabetes (Goldberg et al., 2006).

Finally, although not extensively examined, a study explored whether a preexisting CVD increased the risk of an O₃-induced respiratory effect. Lagorio et al. (2006) examined the effect of O₃ exposure on lung function among participants with a variety of preexisting diseases, including IHD. No association was observed regardless of whether the participant had IHD.

27 Overall, most short-term exposure studies did not report increased O₃-related 28 cardiovascular morbidity for individuals with preexisting CVD. However, as discussed in 29 Section 6.3, most studies reported no overall association between O_3 concentration and 30 CV morbidity. Thus, it is likely the association would be null regardless of the 31 stratification. A limited number of studies examined whether cardiovascular disease 32 modifies the association between O_3 and respiratory effects. There was some evidence 33 that cardiovascular disease increases the risk of O_3 -related mortality but again the number 34 of studies was limited. Currently, evidence is inadequate to classify CVD as a potential 35 at-risk factor for O_3 -related health effects. Future research among those with CVD 36 compared to those without will increase the understanding of potential increased risk of 37 O₃-related health effects among this group.

8.2.5 Diabetes

1	The literature has not extensively examined whether individuals with diabetes (about 8%
2	of U.S. adults) are potentially at increased risk of O3-related health effects. In a study of
3	short-term O3 exposure and cardiovascular ED visits in Atlanta, GA, no association was
4	observed for individuals with or without diabetes (Peel et al., 2007). A similar study
5	conducted in Taiwan reported a positive association between O3 exposure on warm days
6	and hospital admissions for congestive heart failure; however, no modification of the
7	association by diabetes was observed (Lee et al., 2008a). Finally, in a study of O_3
8	exposure and ED visits for arrhythmia in Taiwan, there was no evidence of effect
9	measure modification by diabetes on warm or cool days (Chiu and Yang, 2009).
10	Currently, the limited number of epidemiologic studies as well as the lack of controlled
11	human exposure or toxicological studies provides inadequate evidence to indicate
12	whether diabetes results in a potentially increased risk of O_3 -related health effects.

8.2.6 Hyperthyroidism

13	Hyperthyroidism has been identified in toxicological studies as a potential factor that may
14	lead to increased risk of O3-related health effects but has not yet been explored in
15	epidemiologic or controlled human exposure studies. Lung damage and inflammation due
16	to oxidative stress may be modulated by thyroid hormones. Compared to controls,
17	hyperthyroid rats exhibited elevated levels of BAL neutrophils and albumin after a 4-hour
18	exposure to O_3 , indicating O_3 -induced inflammation and damage. Hyperthyroidism did
19	not affect production of reactive oxygen or nitrogen species, but BAL phospholipids were
20	increased, indicating greater activation of Type II cells and surfactant protein production
21	compared to normal rats (Huffman et al., 2006). Thus, this study provides some
22	underlying evidence, which suggests that individuals with hyperthyroidism may represent
23	an at-risk population; however, overall the lack of additional studies provides inadequate
24	evidence to determine whether hyperthyroidism results in potentially increased risk of
25	O ₃ -related health effects.

8.3 Sociodemographic Factors

8.3.1 Lifestage

26 27 The 1996 and 2006 O_3 AQCDs identified children, especially those with asthma, and older adults as at-risk populations. These previous AQCDs reported clinical evidence that

children have greater spirometric responses to O₃ than middle-aged and older adults (U.S.
 EPA, 1996a). Similar results were observed for symptomatic responses and O₃ exposure.
 Among older adults, most studies reported in the 2006 O₃ AQCD reported greater effects
 of short-term O₃ exposure and mortality compared to other age groups (U.S. EPA,
 Evidence published since the 2006 O₃ AQCD, summarized below, further
 supports these findings.

8.3.1.1 Children

7 The 2000 Census reported that 28.6% of the U.S. population was under 20 years of age, 8 with 14.1% under the age of 10 (SSDAN CensusScope, 2010a). Children's respiratory 9 systems are undergoing lung growth until about 18-20 years of age and are therefore 10 thought to be intrinsically more at risk for O_3 -induced damage (U.S. EPA, 2006b). It is 11 generally recognized that children spend more time outdoors than adults, and therefore 12 would be expected to have higher exposure to O_3 than adults. The ventilation rates also 13 vary between children and adults, particularly during moderate/heavy activity. Children 14 aged 11 years and older and adults have higher absolute ventilation rates than children 15 aged 1 -11 years. However, children have higher ventilation rates relative to their lung 16 volumes, which tends to increase dose normalized to lung surface area. Exercise intensity 17 has a substantial effect on ventilation rate, with high intensity activities resulting in nearly 18 double the ventilation rate during moderate activity among children and those adults less 19 than 31 years of age. For more information on time spent outdoors and ventilation rate 20 differences by age group, see Section 4.4.1.

21 The 1996 O₃ AQCD, reported clinical evidence that children, adolescents, and young 22 adults (<18 years of age) appear, on average, to have nearly equivalent spirometric 23 responses to O_3 exposure, but have greater responses than middle-aged and older adults 24 (U.S. EPA, 1996a). Symptomatic responses (e.g., cough, shortness of breath, pain on 25 deep inspiration) to O_3 exposure, however, appear to increase with age until early 26 adulthood and then gradually decrease with increasing age (U.S. EPA, 1996a). For 27 subjects aged 18-36 years, McDonnell et al. (1999b) reported that symptom responses 28 from O₃ exposure also decrease with increasing age. Complete lung growth and 29 development is not achieved until 18-20 years of age in women and the early 20s for 30 men; pulmonary function is at its maximum during this time as well. Additionally, PBPK 31 modeling reported lung regional extraction of O₃ to be higher in infants compared to 32 adults. This is thought to be due to the smaller nasal and pulmonary regions' surface area 33 in children under the age of 5 years compared to the total airway surface area observed in 34 adults (Sarangapani et al., 2003).

1	Recent epidemiologic studies have examined different age groups and their risk to
2	O3-related respiratory hospital admissions and ED visits. A study in Cyprus of short-term
3	O3 concentrations and respiratory hospital admissions detected possible effect measure
4	modification by age with a larger association among individuals <15 years of age
5	compared with those >15 years of age. However, this difference was only apparent with a
6	2-day lag (Middleton et al., 2008). Similarly, a Canadian study of asthma-eD visits
7	reported the strongest O3-related associations among 5 to 14 year-olds compared to the
8	other age groups (ages examined 0-75+) (Villeneuve et al., 2007). Greater O ₃ -associated
9	risk in asthma-related ED visits were also reported among children (<15 years) as
10	compared to adults (15 to 64 years) in a study from Finland (Halonen et al., 2009). A
11	study of New York City hospital admissions demonstrated an increase in the association
12	between O_3 exposure and asthma-related hospital admissions for 6 to 18 year-olds
13	compared to those <6 years old and those >18 years old (Silverman and Ito, 2010). When
14	examining long-term O ₃ exposure and asthma hospital admissions among children,
15	associations were determined to be larger among children 1 to 2 years old compared to
16	children 2 to 6 years old (Lin et al., 2008b). A few studies reported positive associations
17	among both children and adults and no modification of the effect by age. A study
18	performed in Hong Kong examined O ₃ exposure and asthma-related hospital admissions
19	for ages 0 to 14, 15 to 65, and >65 (Ko et al., 2007). The researchers reported that the
20	association was greater among the 0 to 14 and 14 to 65 age groups compared to the >65
21	age group. Another study looking at asthma-related ED visits and O ₃ exposure in Maine
22	reported positive associations for all age groups (ages 2 to 65) (Paulu and Smith, 2008).
23	Effects of O ₃ exposure on asthma hospitalizations among both children and adults (<18
24	and \geq 18 years old) were demonstrated in a study in Washington, but only children (<18
25	years of age) had statistically significant results at lag day 0, which the authors wrote,
26	"suggests that children are more immediately responsive to adverse effects of O ₃
27	exposure" (Mar and Koenig, 2009).
29	
28	The evidence reported in epidemiologic studies is supported by recent toxicological
29	studies which observed O_3 -induced health effects in immature animals. Early life
30	exposures of multiple species of laboratory animals, including infant monkeys, resulted
31	in changes in conducting airways at the cellular, functional, ultra-structural, and
32	morphological levels. <u>Carey et al. (2007</u>) conducted a study of O_3 exposure in infant
33	rhesus macaques, whose respiratory tract closely resemble that of humans. Monkeys were
34	exposed either acutely for 5 days to 0.5 ppm O_3 , or episodically for 5 biweekly cycles
35	alternating 5 days of 0.5 ppm O_3 with 9 days of filtered air, designed to mimic human
36	exposure (70 days total). All monkeys acutely exposed to O_3 had moderate to marked

necrotizing rhinitis, with focal regions of epithelial exfoliation, numerous infiltrating neutrophils, and some eosinophils. The distribution, character, and severity of lesions in episodically exposed infant monkeys were similar to that of acutely exposed animals.

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1	Neither exposure protocol for the infant monkeys produced mucous cell metaplasia
2	proximal to the lesions, an adaptation observed in adult monkeys exposed continuously to
3	0.3 ppm O_3 in another study (<u>Harkema et al., 1987a</u>). Functional (increased airway
4	resistance and responsiveness with antigen $+ O_3$ co-exposure) and cellular changes in
5	conducting airways (increased numbers of inflammatory eosinophils) were common
6	manifestations of exposure to O_3 among both the adult and infant monkeys (Plopper et
7	al., 2007). In addition, the lung structure of the conducting airways in the infant monkeys
8	was stunted by O_3 and this aberrant development was persistent 6 months postexposure.
9	This developmental endpoint was not, of course, studied in the adult monkey experiments
10	(Fanucchi et al., 2006). Thus, some functional and biochemical effects were similar
11	between the infant and adult monkeys exposed to O ₃ , but because the study designs did
12	not include concentration-response experiments, it is not possible to determine whether
13	the infant monkeys were more at risk for the effects of O_3 .

- 14Similarly, rat fetuses exposed to O3 in utero had ultrastructural changes in bronchiolar15epithelium when examined near the end of gestation (López et al., 2008). In addition,16exposure of mice to mixtures of air pollutants early in development affected pup lung17cytokine levels (TNF, IL-1, KC, IL-6, and MCP-1) (Auten et al., 2009). In utero exposure18of animals to PM augmented O3-induced airway hyper-reactivity in these pups as19juveniles.
- 20 Age may affect the inflammatory response to O_3 exposure. In comparing neonatal mice to 21 adult mice, increased bronchoalveolar lavage (BAL) neutrophils were observed in four 22 strains of neonates 24 hours after exposure to 0.8 ppm O_3 for 5 hours (Vancza et al., 23 2009). Three of these strains also exhibited increased BAL protein, although the two 24 endpoints were not necessarily consistently correlated in a given strain. In some strains, 25 however, adults were responsive, indicating a strain-age interaction. Measurement of ¹⁸O 26 determined that the observed strain- and age-dependent differences were not due to 27 absorbed O₃ dose. Using electron microscopy, Bils (1970) studied the lungs of mice of 28 different ages (4 days or 1 to 2 months) exposed to 0.6 to 1.3 ppm O₃ for 6 to 7 h/day for 29 1 to 2 days and noted swelling of the alveolar epithelial lining cells without intra-alveolar 30 edema. Swelling of endothelial cells and occasional breaks in the basement membrane 31 were observed. These effects were most evident in younger mice exposed for 2 days. 32 Toxicological studies reported that the difference in effects among younger lifestage test 33 animals may be due to age-related changes in endogenous antioxidants and sensitivity to 34 oxidative stress. A recent study demonstrated that 0.25 ppm O₃ exposure differentially 35 altered expression of metalloproteinases in the skin of young (8 weeks old) and aged 36 (18 months old) mice, indicating age-related susceptibility to oxidative stress (Fortino et 37 al., 2007). Valacchi et al. (2007) found that aged mice had more Vitamin E in their 38 plasma but less in their lungs compared to young mice, which may affect their pulmonary

1	antioxidant defenses. Servais et al. (2005) found higher levels of oxidative damage
2	indicators in immature (3 weeks old) and aged (20 months old) rats compared to adult
3	rats, the latter which were relatively resistant to an intermittent 7-day exposure to
4	0.5 ppm O ₃ . Immature rats exhibited a higher ventilation rate, which may have increased
5	exposure. Additionally, a series of toxicological studies reported an association between
6	O_3 exposure and bradycardia that was present among young but not older mice (<u>Hamade</u>
7	et al., 2010; Tankersley et al., 2010; Hamade and Tankersley, 2009; Hamade et al., 2008).
8	Regression analysis revealed an interaction between age and strain on heart rate, which
9	implies that aging may affect heart rate differently among mouse strains (Tankersley et
10	al., 2010). The authors proposed that the genetic differences between the mice strains
11	could be altering the formation of ROS, which tends to increase with age, thus
12	modulating the changes in cardiopulmonary physiology after O ₃ exposure.
13	The previous and recent human clinical and toxicological studies reported evidence of
13 14	The previous and recent human clinical and toxicological studies reported evidence of increased risk from O_3 exposure for younger ages, which provides coherence and
14	increased risk from O ₃ exposure for younger ages, which provides coherence and
14 15	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was
14 15 16	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for
14 15 16 17	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation
14 15 16 17 18	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation of these studies is limited by the lack of consistency in comparison age groups and
14 15 16 17 18 19	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation of these studies is limited by the lack of consistency in comparison age groups and outcomes examined. Toxicological studies observed O_3 -induced health effects in
14 15 16 17 18 19 20	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation of these studies is limited by the lack of consistency in comparison age groups and outcomes examined. Toxicological studies observed O_3 -induced health effects in immature animals, including infant monkeys, though the effects were not consistently
14 15 16 17 18 19 20 21	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation of these studies is limited by the lack of consistency in comparison age groups and outcomes examined. Toxicological studies observed O_3 -induced health effects in immature animals, including infant monkeys, though the effects were not consistently greater in young animals than adults. However, overall, the epidemiologic, controlled
14 15 16 17 18 19 20 21 22	increased risk from O_3 exposure for younger ages, which provides coherence and biological plausibility for the findings from epidemiologic studies. Although there was some inconsistency, generally, the epidemiologic studies reported larger associations for respiratory hospital admissions and ED visits for children than adults. The interpretation of these studies is limited by the lack of consistency in comparison age groups and outcomes examined. Toxicological studies observed O_3 -induced health effects in immature animals, including infant monkeys, though the effects were not consistently greater in young animals than adults. However, overall, the epidemiologic, controlled human exposure, and toxicological studies provide substantial and consistent evidence

8.3.1.2 Older Adults

25	Older adults may be at greater risk of health effects associated with O ₃ exposure through
26	a variety of intrinsic pathways. In addition, older adults may differ in their exposure and
27	internal dose. Older adults were outdoors for a slightly longer proportion of the day than
28	adults aged 18-64 years. Older adults also have somewhat lower ventilation rates than
29	adults aged 31 - less than 61 years. For more information on time spent outdoors and
30	ventilation rate differences by age group, see Section $4.4.1$. The gradual decline in
31	physiological processes that occur with aging may lead to increased risk of O3-related
32	health effects (U.S. EPA, 2006a). Respiratory symptom responses to O_3 exposure appears
33	to increase with age until early adulthood and then gradually decrease with increasing age
34	(U.S. EPA, 1996a), which may put older adults at increased risk by withstanding
35	continued O3 exposure and thus not seeking relief and avoiding exposure. In addition,

- 1 older adults, in general, have a higher prevalence of preexisting diseases, with the 2 exception of asthma, compared to younger age groups and this may also lead to increased 3 risk of O₃-related health effects (see Table 8-3 that gives preexisting rates by age). With 4 the number of older Americans increasing in upcoming years (estimated to increase from 5 12.4% of the U.S. population to 19.7% between 2000 to 2030, which is approximately 6 35 million and 71.5 million individuals, respectively) this group represents a large 7 population potentially at risk of O_3 -related health effects (SSDAN CensusScope, 2010a; 8 U.S. Census Bureau, 2010).
- 9 The majority of recent studies reported greater effects of short-term O_3 exposure and 10 mortality among older adults, which is consistent with the findings of the 2006 O_3 11 AQCD. A study conducted in 48 cities across the U.S. reported larger effects among 12 adults \geq 65 years old compared to those <65 years (Medina-Ramón and Schwartz, 2008). 13 Further investigation of this study population revealed a trend of O₃-related mortality risk 14 that gets larger with increasing age starting at age 50 (Zanobetti and Schwartz, 2008a). A 15 study of 7 urban centers in Chile reported similar results, with greater effects in adults 16 \geq 65 years old, however the effects were smaller among those \geq 85 years old compared to 17 those in the 75 to 84 years old age range (Cakmak et al., 2007). More recently, a study 18 conducted in the same area reported similar associations between O_3 exposure and 19 mortality in adults aged <64 years old and 65 to 74 years old, but the risk was increased 20 among older age groups (Cakmak et al., 2011). A study performed in China reported 21 greater effects in populations \geq 45 years old (compared to 5 to 44 year-olds), with 22 statistically significant effects present only among those ≥ 65 years old (Kan et al., 2008). 23 An Italian study reported higher risk of all-cause mortality associated with increased O_3 24 concentrations among individuals \geq 85 years old as compared to those 35 to 84 years old. 25 Those 65 to 74 and 75 to 84 years old did not show a greater increase in risk compared to 26 those aged 35 to 64 years (Stafoggia et al., 2010). The Air Pollution and Health: A 27 European and North American Approach (APHENA) project examined the association 28 between O_3 exposure and mortality for those <75 and \geq 75 years of age. In Canada, the 29 associations for all-cause and cardiovascular mortality were greater among those 30 \geq 75 years old in the summer-only and all-year analyses. Age groups were not compared 31 in the analysis for respiratory mortality in Canada. In the U.S., the association for 32 all-cause mortality was slightly greater for those <75 years of age compared to those ≥ 75 33 years old in summer-only analyses. No consistent pattern was observed for CVD 34 mortality. In Europe, slightly larger associations for all-cause mortality were observed in 35 those <75 years old in all-year and summer-only analyses. Larger associations were 36 reported among those <75 years for CVD mortality in all-year analyses, but the reverse 37 was true for summer-only analyses (Katsouyanni et al., 2009).

1	Multiple epidemiologic studies of O ₃ exposure and hospital admissions were stratified by
2	age groups. A positive association was reported between short-term O_3 exposure and
3	respiratory hospital admissions for adults ≥ 65 years old but not for those adults aged 15
4	to 64 years (Halonen et al., 2009). In the same study, no association was observed
5	between O_3 concentration and respiratory mortality among those ≥ 65 years old or those
6	15 to 64 years old; however, an inverse association between O ₃ concentration and
7	cardiovascular mortality was present among individuals ≥ 65 years old but not among
8	individuals <65 years old. This inverse association among those \geq 65 years old persisted
9	when examining hospital admissions for coronary heart disease. A study of CVD-related
10	hospital visits in Bangkok, Thailand reported an increase in percent change for hospital
11	visits with previous day and cumulative 2-day O_3 levels among those ≥ 65 years old,
12	whereas no association was present for individuals less than 65 years of age (Buadong et
13	al., 2009). No association was observed for current day or cumulative 3-day averages in
14	any age group. A study examining O_3 and hospital admissions for CVD-related health
15	effects reported no association for individuals aged 15 to 64 or individuals aged ≥ 65
16	years, although one lag-time did show an inverse effect for coronary heart disease among
17	elderly that was not present among 15 to 64 year-olds (Halonen et al., 2009). However, as
18	discussed in the Section on CVD hospital admissions ($6.3.2.7$), results were inconsistent
19	and often null so it is plausible that no association would be observed regardless of age.
20	No modification by age (40 to 64 year-olds versus >64 years old) was observed in a study
21	from Brazil examining O_3 levels and COPD ED visits (<u>Arbex et al., 2009</u>).
22	Biological plausibility for differences by age is provided by toxicological studies. O_3

Biological plausibility for differences by age is provided by toxicological studies. O₃ 22 23 exposure resulted in an increase in left ventricular chamber dimensions at end diastole 24 (LVEDD) in young and old mice, whereas decreases in left ventricular posterior wall 25 thickness at end systole (PWTES) were only observed among older mice (Tankersley et 26 al., 2010). Other toxicological studies also indicate increased risk in older animals for 27 additional endpoints, including neurological and immune. The hippocampus, one of the 28 main regions affected by age-related neurodegenerative diseases, may be more sensitive 29 to oxidative damage in aged rats. In a study of young (47 days) and aged (900 days) rats 30 exposed to 1 ppm O₃ for 4 hours, O₃-induced lipid peroxidation occurred to a greater 31 extent in the striatum of young rats, whereas it was highest in the hippocampus in aged 32 rats (Rivas-Arancibia et al., 2000). In young mice, healing of skin wounds is not 33 significantly affected by O_3 exposure (Lim et al., 2006). However, exposure to 0.5 ppm 34 O₃ for 6 h/day significantly delays wound closure in aged mice.

35Although some outcomes reported mixed findings regarding an increase in risk for older36adults, recent epidemiologic studies report consistent positive associations between short-37term O3 exposure and mortality in older adults. The evidence from mortality studies is38consistent with the results reported in the 2006 O3 AQCD and is supported by

1	toxicological studies providing biological plausibility for increased risk of effects in older
2	adults. Also, older adults may be experiencing increased exposure compared to younger
3	adults. Overall, adequate evidence is available indicating that older adults are potentially
4	at increased risk of O3-related health effects based on the substantial and consistent
5	evidence within epidemiologic studies on O3 exposure and mortality and the coherence
6	with toxicological studies.

8.3.2 Sex

7	The distribution of males and females in the U.S. is similar. In 2000, 49.1% of the U.S.
8	population was male and 50.9% were female. However, this distribution does vary by age
9	with a greater prevalence of females ≥ 65 years old compared to males (<u>SSDAN</u>
10	CensusScope, 2010a). The 2006 O_3 AQCD did not report evidence of differences
11	between the sexes in health responses to O_3 exposure (U.S. EPA, 2006b). Recent
12	epidemiologic studies have evaluated the effects of short-term and long-term exposure to
13	O_3 on multiple health endpoints stratified by sex.
14	A study in Maine that examined short-term O ₃ concentrations and asthma ED visits
15	detected greater effects among males ages 2 to 14 years and among females ages 15 to 34
16	years compared to males and females in the same age groups (no difference was detected
17	for males and females aged 35 to 64) (Paulu and Smith, 2008). A Canadian study
18	reported no associations between short-term O_3 and respiratory infection hospital
19	admissions for either boys or girls under the age of 15 (Lin et al., 2005), whereas another
20	Canadian study reported a slightly higher but non-statistically significant increase in
21	respiratory hospital admissions for males (mean ages 47.6 to 69.0 years) (Cakmak et al.,
22	2006b). A recent study from Hong Kong examining individuals of all ages reported no
23	effect measure modification by sex for overall respiratory disease hospital admissions,
24	but did detect a greater excess risk of hospital admissions for COPD among females
25	compared to males (Wong et al., 2009). Similarly a study in Brazil found higher effect
26	estimates for COPD ED visits among females compared to males (Arbex et al., 2009).
27	Higher levels of respiratory hospital admissions with greater O ₃ concentrations was also
28	observed for females in a study of individuals living in Cyprus (Middleton et al., 2008).
29	A study of lung function unrelated to hospital admissions and ED visits was conducted
30	among lifeguards in Texas and reported decreased lung function with increased O_3
31	exposure among females but not males (Thaller et al., 2008). This study included
32	individuals aged 16 to 27 years, and the majority of participants were male. A New York
33	study found no evidence of effect measure modification of the association between
34	long-term O_3 exposure and asthma hospital admissions among males and females
35	between 1 and 6 years old (Lin et al., 2008b).

- 1 In addition to examining the potential modification of O_3 associations with respiratory 2 outcomes by sex, studies also examined cardiovascular-related outcomes specifically 3 hospital admissions and ED visits. All of these studies reported no effect modification by 4 sex with some studies reporting null associations for both males and females (Wong et 5 al., 2009; Middleton et al., 2008; Villeneuve et al., 2006a) and one study reporting a 6 positive associations for both sexes (Cakmak et al., 2006a). A French study examining 7 the associations between O_3 concentrations and risk of ischemic strokes (not limited to 8 ED visits or hospital admissions) reported no association for either males or females with 9 lags of 0, 2, or 3 days (Henrotin et al., 2007). A positive association was reported for 10 males with a lag of 1 day, but this association was null for females. The authors noted 11 that men in the study had much higher rates of current and former smoking than women 12 (67.4% versus 9.3%). Additionally, cardiovascular hospital admissions and ED visits 13 overall have demonstrated inconsistent and null results (Section 6.3.2.7). The lack of 14 effect measure modification by sex may be indicative of the lack of association, not the 15 lack of effect of sex.
- 16 A biomarker study investigating the effects of O_3 concentrations on high-sensitivity 17 C-reactive protein (hs-CRP), fibrinogen, and white blood cell (WBC) count, reported 18 observations for various lag times ranging from 0 to 7 days (Steinvil et al., 2008). Most 19 of the associations were null for males and females although one association between O_3 20 and fibrinogen was positive for males and null for females (lag day 4); however, this 21 positive association was null or negative when other pollutants were included in the 22 model. One study examining correlations between O₃ levels and oxidative DNA damage 23 examined results stratified by sex. In this study Palli et al. (2009) reported stronger 24 correlations for males than females, both during short-term exposure (less than 30 days) 25 and long-term exposure (0-90 days). However, the authors commented that this 26 difference could have been partially explained by different distributions of exposure to 27 traffic pollution at work.
- 28 A few studies have examined the association between short-term O_3 concentrations and 29 mortality stratified by sex and, in contrast with studies of other endpoints, were more 30 consistent in reporting elevated risks among females. These studies, conducted in the 31 U.S. (Medina-Ramón and Schwartz, 2008), Italy (Stafoggia et al., 2010), and Asia (Kan 32 et al., 2008), reported larger effect estimates in females compared to males. In the U.S. 33 study, the elevated risk of mortality among females was greater specifically among those 34 \geq 60 years old (Medina-Ramón and Schwartz, 2008). However, a recent study in Chile 35 reported similar associations between O₃ exposure and mortality among both men and 36 women (Cakmak et al., 2011). A long-term O_3 exposure study of respiratory mortality 37 stratified their results by sex and reported relative risks of 1.01 (95% CI: 0.99, 1.04) for 38 males and 1.04 (95% CIs 1.03, 1.07) for females (Jerrett et al., 2009).

1	Experimental response provided a further understanding of the underlying mechanisms
1 2	Experimental research provided a further understanding of the underlying mechanisms that may explain a possible differential risk in O_3 -related health effects among males and
2 3	
3 4	females. Several studies have suggested that physiological differences between sexes
	may predispose females to greater effects from O_3 . In females, lower plasma and nasal
5	lavage fluid (NLF) levels of uric acid (most prevalent antioxidant), the initial defense
6 7	mechanism of O_3 neutralization, may be a contributing factor (<u>Housley et al., 1996</u>).
7	Consequently, reduced absorption of O_3 in the upper airways of females may promote its
8	deeper penetration. Dosimetric measurements have shown that the absorption distribution
9	of O_3 is independent of sex when absorption is normalized to anatomical dead space
10	(Bush et al., 1996). Thus, a differential removal of O_3 by uric acid seems to be minimal.
11	In general, the physiologic response of young healthy females to O ₃ exposure appears
12	comparable to the response of young males (<u>Hazucha et al., 2003</u>). A few studies have
13	examined changes in O ₃ responses during various menstrual cycle phases. Lung function
14	response to O ₃ was enhanced during the follicular phase of the menstrual cycle compared
15	to the luteal phase in a small study of women (Fox et al., 1993). However, Seal et al.
16	(1996) later reported no effect of menstrual cycle phase in their analysis of responses
17	from 150 women, but conceded that the methods used by Fox et al. (1993) more precisely
18	defined the menstrual cycle phase. Another study also reported no difference in responses
19	among females during the follicular and luteal phases of their cycle (Weinmann et al.,
20	1995c). Additionally, in this study the responses in women were comparable to those
21	reported for men in the study. In a toxicological study, small differences in effects by sex
22	were seen in adult mice with respect to pulmonary inflammation and injury after a 5-h
23	exposure to 0.8 ppm O_3 , and although adult females were generally more at risk, these
24	differences were strain-dependent, with some strains exhibiting greater risk in males
25	(Vancza et al., 2009). The most obvious sex difference was apparent in lactating females,
26	which incurred the greatest lung injury or inflammation among several of the strains.
27	Overall, results have varied, with recent evidence for increased risk for O ₃ -related health
28	effects present for females in some studies and males in other studies. Most studies
29	examining the associations O_3 and mortality report females to be at greater risk than
30	males, but minimal evidence is available regarding a difference between the sexes for
31	other outcomes. Inconsistent findings were reported on whether effect measure
32	modification exists by sex for respiratory and cardiovascular hospital admissions and ED
33	visits, although there is some indication that females are at increased risk of O ₃ -related
34	respiratory hospital admissions and ED visits. While O3-related effects may occur in both
35	men and women, there is suggestive evidence exists indicating that females are at
36	potentially increased risk of O ₃ -related health effects as there are consistent findings
37	among epidemiologic studies of mortality.

8.3.3 Socioeconomic Status

1 SES is often represented by personal or neighborhood SES, which is comprised of a 2 variety of components such as educational attainment, household income, health 3 insurance status, and other such factors. SES is often indicative of such things as access 4 to healthcare, quality of housing, and pollution gradient to which people are exposed. 5 One or a combination of these components could modify the risk of O₃-related health 6 effects. Based on the 2000 Census data, 12.4% of Americans live in poverty (poverty 7 threshold for family of four was \$17,463) (SSDAN CensusScope, 2010c). Although 8 included below, studies stratifying by SES that are conducted outside the U.S. may not be 9 comparable to those studies from within the U.S. Having low SES in another country 10 may be different than having low SES in the U.S. based on SES definitions, population 11 composition, and/or conditions in that country.

12 Multiple epidemiologic studies have reported individuals of low SES to have increased 13 risk for the effects of short-term O₃ exposure on respiratory hospital admissions and ED 14 visits. In New York State, larger associations between long-term O₃ exposure and asthma 15 hospital admissions were observed among children of mothers who did not graduate from 16 high school, whose births were covered by Medicaid/self-paid, or who were living in 17 poor neighborhoods compared to children whose mothers graduated from high school, 18 whose births were covered by other insurance, or who were not living in poor 19 neighborhoods, respectively (Lin et al., 2008b). In addition, a study conducted across 10 20 cities in Canada found the largest association between O_3 exposure and respiratory 21 hospital admissions was among those with an educational level less than grade 9, but no 22 consistent trend in the effect was seen across quartiles of income (Cakmak et al., 2006b). 23 A Canadian study reported inverse effects of O₃ on respiratory hospital admissions and 24 ED visits for all levels of SES, measured by average census tract household income 25 (Burra et al., 2009). A study performed in Korea examined the association between O_3 26 concentrations and asthma hospital admissions and reported larger effect estimates in 27 areas of moderate and low SES compared with areas of high SES (SES was based on 28 average regional insurance rates) (Lee et al., 2006).

The examination of the potential effects of SES on O₃-related cardiovascular health
effects is relatively limited. A study conducted in Canada reported the association
between short-term O₃ and ED visits for cardiac disease by quartiles of
neighborhood-level education and income. No effect measure modification was apparent
for either measure of SES (Cakmak et al., 2006a). However, this may be due to the lack
of association present between O₃ and ED visits for cardiac disease regardless of SES.
Several studies were conducted that examined the modification of the relationship

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between short-term O₃ concentrations and mortality by SES. A U.S. multicity study

1	reported that communities with a higher proportion of the population unemployed had
2	higher O ₃ -related mortality effect estimates (Bell and Dominici, 2008). A study in seven
3	urban centers in Chile reported on modification of the association between O ₃ exposure
4	and mortality using multiple SES markers (Cakmak et al., 2011). Increased risk was
5	observed among the categories of low SES for all measures (personal educational
6	attainment, personal occupation, community income level). Additionally, the APHENA
7	study, which examined the association between O_3 and mortality by percentage
8	unemployed, reported a higher percent change in mortality with increased percent
9	unemployed but this varied across the regions included in the study (U.S., Canada,
10	Europe) (Katsouyanni et al., 2009). A Chinese study reported that the greatest effects
11	between O_3 concentrations and mortality at lag day 0 were among individuals living in
12	areas of high social deprivation (i.e., low SES), but this association was not consistent
13	across lag days (at other lag times, the middle social deprivation index category had the
14	greatest association) (Wong et al., 2008). However, another study in Asia comparing low
15	to high educational attainment populations reported no evidence of greater mortality
16	effects (total, CVD, or respiratory) (Kan et al., 2008). Additionally, a study in Italy
17	reported no difference in risk of mortality among census-block level derived income
18	levels (Stafoggia et al., 2010). A study of infant mortality in Mexico reported no
19	association between O3 concentrations and infant mortality among any of the three levels
20	of SES determined using a socioeconomic index based on residential areas (Romieu et
21	<u>al., 2004a</u>). Another study in Mexico reported a positive association between O_3 levels at
22	lag 0 and respiratory-related infant mortality in only the low SES group (determined
23	based on education, income, and household conditions across residential areas), but no
24	association was observed in any of the SES groups with other lags (Carbajal-Arroyo et
25	<u>al., 2011</u>).
26	Studies of O3 concentrations and reproductive outcomes have also examined associations
27	by SES levels. A study in California reported greater decreases in birth weight associated
28	with full pregnancy O_3 concentration for those with neighborhood poverty levels of at
29	least 7% compared with those in neighborhoods with less than 7% poverty (the authors
30	do not provide information on how categories of the SES variable were determined)
31	(Morello-Frosch et al., 2010). No dose response was apparent and those with

(<u>Morello-Frosch et al., 2010</u>). No dose response was apparent and those with 32 neighborhood poverty levels of 7-21% had greater decreases observed for the association 33 than those living in areas with poverty rates of at least 22%. An Australian study reported 34 an inverse association between O₃ exposure during days 31-60 of gestation and 35 abdominal circumference during gestation (Hansen et al., 2008). The interaction with 36 SES (area-level measured socioeconomic disadvantage) was examined and although the 37 inverse association remained statistically significant in only the highest SES quartile, 38 there were large confidence interval overlaps among estimates for each quartile so no 39 difference in the association for the quartiles was apparent.

1	Evidence from a controlled human exposure study that examined O ₃ effects on lung
2	function does not provide support for greater O3-related health effects in individuals of
3	lower SES. In a follow-up study (Seal et al., 1993) on modification by race, Seal et al.
4	(1996) reported that, of three SES categories, individuals in the middle SES category
5	showed greater concentration-dependent decline in percent-predicted FEV_1 (4-5% at
6	400 ppb O ₃) than in low and high SES groups. The authors did not have an "immediately
7	clear" explanation for this finding and controlled human exposure studies are typically
8	not designed to answer questions about SES.
9	Overall, most studies of individuals have reported that individuals with low SES and
10	those living in neighborhoods with low SES are more at risk for O_3 -related health effects,
11	resulting in increased risk of respiratory hospital admissions and ED visits. Inconsistent
12	results have been observed in the few studies examining effect modification of
13	associations between O ₃ exposure and mortality and reproductive outcomes. Also, a
14	controlled human exposure study does not support evidence of increased risk of
15	respiratory morbidity among individuals with lower SES. Overall, evidence is suggestive
16	of SES as a factor affecting risk of O_3 -related health outcomes based on collective
17	evidence from epidemiologic studies of respiratory hospital admissions but inconsistency
18	among epidemiologic studies of mortality and reproductive outcomes. Further studies are
19	needed to confirm this relationship, especially in populations within the U.S.

8.3.4 Race/Ethnicity

20	Based on the 2000 Census, 69.1% of the U.S. population identified as non-Hispanic
21	whites. Approximately 12.1% of people reported their race/ethnicity as non-Hispanic
22	black and 12.6% reported being Hispanic (SSDAN CensusScope, 2010b).
23	Only a few studies examined the associations between short-term O ₃ concentrations and
24	mortality and reported higher effect estimates among blacks (Medina-Ramón and
25	Schwartz, 2008) and among communities with larger proportions of blacks (Bell and
26	Dominici, 2008). Another study examined long-term exposure to O ₃ concentrations and
27	asthma hospital admissions among children in New York State. These authors reported
28	no statistically significant difference in the odds of asthma hospital admissions for blacks
29	compared to other races but did detect higher odds for Hispanics compared to
30	non-Hispanics (<u>Lin et al., 2008b</u>).
31	Additionally, recent epidemiologic studies have stratified by race when examining the
32	association between O ₃ concentration and birth outcomes. A study conducted in Atlanta,
33	GA reported decreases in birth weight with increased third trimester O ₃ concentrations
34	among Hispanics but not among non-Hispanic whites (Darrow et al., 2011b). An inverse

- association was also present for non-Hispanic blacks but was not statistically significant. A California study reported that the greatest decrease in birth weight associated with full pregnancy O₃ concentration was among non-Hispanic whites (<u>Morello-Frosch et al.</u>, <u>2010</u>). This inverse association was also apparent, although not as strong, for Hispanics and non-Hispanic blacks. Increased birth weight was associated with higher O₃ exposure among non-Hispanic Asians and Pacific Islanders but these results were not statistically significant.
- 8 Similar to the epidemiologic studies, a controlled human exposure study suggested 9 differences in lung function responses by race (Seal et al., 1993). The independent effects 10 of sex-race group and O₃ concentration on lung function were positive, but the interaction 11 between sex-race group and O_3 concentration was not statistically significant. The 12 findings indicated some overall difference between the sex-race groups that was 13 independent of O_3 concentration (the concentration-response curves for the four sex-race 14 groups are parallel). In a multiple comparison procedure on data collapsed across all O_3 15 concentrations for each sex-race group, both black men and black women had larger 16 decrements in FEV_1 than did white men. The authors noted that the O₃ dose per unit of 17 lung tissue would be greater in blacks and females than whites and males, respectively. That this difference in tissue dose might have affected responses to O₃ cannot be ruled 18 19 out. The college students recruited for the Seal et al. (1993) study were probably from 20 better educated and more SES advantaged families, thus reducing potential for these 21 variables to be confounding factors. Que et al. (2011) also examined pulmonary 22 responses to O_3 exposure in blacks of African American ancestry and in whites. On 23 average, the black males experienced the greatest decrements in FEV₁ following O₃ 24 exposure. This decrease was larger than the decrement observed among black females, 25 white males, and white females.
- 26Overall, the results of recent studies indicate that there may be race-related increase in27risk of O3-related health effects for some outcomes, although the overall understanding of28potential effect measure modification by race is limited by the small number of studies.29Additionally, these results may be confounded by other factors, such as SES. Overall,30evidence is inadequate to determine if O3-related health effects vary by race because of31the insufficient quantity of studies and lack of consistency within disciplines.

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8.4 Behavioral and Other Factors

8.4.1 Diet

1	Diet was not examined as a factor potentially affecting risk in previous O ₃ AQCDs, but
2	recent studies have examined modification of the association between O ₃ and health
3	effects by dietary factors. Because O ₃ mediates some of its toxic effects through oxidative
4	stress, the antioxidant status of an individual is an important factor that may contribute to
5	increased risk of O3-related health effects. Supplementation with Vitamins C and E has
6	been investigated in a number of studies as a means of inhibiting O ₃ -mediated damage.
7	Epidemiologic studies have examined effect measure modification by diet and found
8	evidence that certain dietary components are related to the effect O3 has on respiratory
9	outcomes. In a recent study the effects of fruit/vegetable intake and Mediterranean diet
10	was examined (Romieu et al., 2009). Increases in these food patterns, which have been
11	noted for their high Vitamins C and E and omega-3 fatty acid content, protected against
12	O ₃ -related decreases in lung function among children living in Mexico City. Another
13	study examined supplementation of the diets of asthmatic children in Mexico with
14	Vitamins C and E (Sienra-Monge et al., 2004). Associations were detected between
15	short-term O_3 exposure and nasal airway inflammation among children in the placebo
16	group but not in those receiving the supplementation. The authors concluded that
17	"Vitamin C and E supplementation above the minimum dietary requirement in asthmatic
18	children with a low intake of Vitamin E might provide some protection against the nasal
19	acute inflammatory response to ozone."
20	The epidemiologic evidence is supported by controlled human exposure studies, which
21	have shown that the first line of defense against oxidative stress is antioxidants-rich
22	extracellular lining fluid (ELF) which scavenge free radicals and limit lipid peroxidation.
23	Exposure to O_3 depletes the antioxidant level in nasal ELF probably due to scrubbing of
24	O_3 (Mudway et al., 1999a); however, the concentration and the activity of antioxidant
25	enzymes either in ELF or plasma do not appear to be related to O_3 responsiveness
26	(e.g., pulmonary function and inflammation) (Samet et al., 2001; Avissar et al., 2000;
27	Blomberg et al., 1999). Carefully controlled studies of dietary antioxidant
28	supplementation have demonstrated some protective effects of α -tocopherol (a form of
29	Vitamin E) and ascorbate (Vitamin C) on spirometric measures of lung function after O_3
30	exposure but not on the intensity of subjective symptoms and inflammatory response
31	including cell recruitment, activation and a release of mediators (Samet et al., 2001;
32	Trenga et al., 2001). Dietary antioxidants have also afforded partial protection to

asthmatics by attenuating postexposure bronchial hyperresponsiveness (<u>Trenga et al.</u>, <u>2001</u>).

- 3 Toxicological studies provide evidence of biological plausibility to the epidemiologic and 4 controlled human exposure studies. Wagner et al. (2009); (2007) found reductions in 5 O_3 -exacerbated nasal allergy responses in rats with γ -tocopherol treatment (a form of 6 Vitamin E). O₃-induced inflammation and mucus production were also inhibited by 7 γ -tocopherol. Supplementation with Vitamins C and E partially ameliorated 8 inflammation, oxidative stress, and airway hyperresponsiveness in guinea pigs exposed 9 subchronically to 0.12 ppm O₃ ppm (<u>Chhabra et al., 2010</u>). Inconsistent results were 10 observed in other toxicological studies of Vitamin C deficiency and O₃-induced 11 responses. Guinea pigs deficient in Vitamin C displayed only minimal injury and 12 inflammation after exposure to O₃ (Kodavanti et al., 1995). A recent study in mice 13 demonstrated a protective effect of β -carotene in the skin, where it limited the production 14 of proinflammatory markers and indicators of oxidative stress induced by O_3 exposure 15 (Valacchi et al., 2009). Deficiency of Vitamin A, which has a role in regulating the 16 maintenance and repair of the epithelial layer, particularly in the lung, appears to enhance 17 the risk of O_3 -induced lung injury (Paquette et al., 1996). Differentially susceptible 18 mouse strains that were fed a Vitamin A sufficient diet were observed to have different 19 tissue concentrations of the vitamin, potentially contributing to their respective 20 differences in O₃-related outcomes. In addition to the studies of antioxidants, one 21 toxicological study examined protein deficiency. Protein deficiency alters the levels of 22 enzymes and chemicals in the brain of rats involved with redox status; exposure to 23 0.75 ppm O_3 has been shown to differentially affect Na⁺/K⁺ ATPase, glutathione, and 24 lipid peroxidation, depending on the nutritional status of the animal, but the significance 25 of these changes is unclear (Calderón Guzmán et al., 2006). There may be a protective 26 effect of overall dietary restriction with respect to lung injury, possibly related to 27 increased Vitamin C in the lung surface fluid (Kari et al., 1997). 28 There is adequate evidence that individuals with reduced intake of Vitamins E and C are
- potentially at risk for O₃-related health effects based on substantial, consistent evidence
 both within and among disciplines. The evidence from epidemiologic studies is supported
 by controlled human exposure and toxicological studies.

8.4.2 Obesity

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32	Obesity, defined as a BMI of 30 kg/m ² or greater, is an issue of increasing importance in
33	the U.S., with self-reported rates of obesity of 26.7% in 2009, up from 19.8% in 2000

1 (Sherry et al., 2010). BMI may affect O_3 -related health effects through multiple avenues, 2 such as, inflammation in the body, increased preexisting disease, and poor diet.

- 3 A few studies have been performed examining the association between BMI and 4 O₃-related changes in lung function. An epidemiologic study reported decreased lung 5 function with increased short-term O_3 exposure for both obese and non-obese subjects; 6 however, the magnitude of the reduction in lung function was greater for those subjects 7 who were obese (Alexeeff et al., 2007). Further decrements in lung function were noted 8 for obese individuals with AHR. Controlled human exposure studies have also detected 9 differential effects of O₃ exposure on lung function for individuals with varying BMIs. In 10 a retrospective analysis of data from 541 healthy, nonsmoking, white males between the ages of 18-35 years from 15 studies conducted at the U.S. EPA Human Studies Facility in 11 12 Chapel Hill, North Carolina, McDonnell et al. (2010) found that increased body mass 13 index (BMI) was found to be associated with enhanced FEV_1 responses. The BMI effect 14 was of the same order of magnitude but in the opposite direction of the age effect 15 whereby FEV₁ responses diminish with increasing age. In a similar analysis, Bennett et 16 al. (2007) found enhanced FEV_1 decrements following O₃ exposure with increasing BMI 17 in a group of healthy, nonsmoking, women (BMI range 15.7 to 33.4), but not among 18 healthy, nonsmoking men (BMI range 19.1 to 32.9). In the women, greater O_3 -induced 19 FEV_1 decrements were seen in individuals that were overweight/obese (BMI >25) 20 compared to normal weight (BMI from 18.5 to 25), and in normal weight compared to 21 underweight (BMI <18.5). Even disregarding the five underweight women, a greater O_3 22 response in the overweight/obese category (BMI >25) was observed compared with the 23
- 24 Studies in genetically and dietarily obese mice have shown enhanced pulmonary 25 inflammation and injury with acute O_3 exposure, but responses to longer exposures at a 26 lower concentration appear to differ. A recent study found that obese mice are actually 27 resistant to O₃-induced pulmonary injury and inflammation and reduced lung compliance 28 following exposure to 0.3 ppm O_3 for 72 hours, regardless of whether obesity was 29 genetic- or diet-induced (Shore et al., 2009).

normal weight group (BMI from 18.5 to 24.9).

30 Multiple epidemiologic, human clinical, and toxicological studies have reported 31 suggestive evidence for increased O₃-related respiratory health effects among obese 32 individuals. Future research of the effect modification of the relationship between O_3 and 33 other health-related outcomes besides respiratory health effects by BMI and studies 34 examining the role of physical conditioning will advance understanding of obesity as a 35 factor potentially increasing an individual's risk.

8.4.3 Smoking

1	Previous O_3 AQCDs have concluded that smoking does not increase the risk of O_3 -related
2	health effects; in fact, in controlled human exposure studies, smokers have been found to
3	be less responsive to O_3 than non-smokers. Data from recent interviews conducted as part
4	of the 2008 National Health Interview Survey (NHIS) (Pleis et al., 2009) have shown the
5	rate of smoking among adults \geq 18 years old to be approximately 20% in the U.S.
6	Approximately 21% of individuals surveyed were identified as former smokers.
7	Baccarelli et al. (2007) performed a study of O_3 concentrations and plasma homocysteine
8	levels (a risk factor for vascular disease). They found no interaction of smoking (smokers
9	versus non-smokers) for the associations between O_3 concentrations and plasma
10	homocysteine levels. Another study examined the association between O_3 and resting
11	heart rate and also reported no interaction with smoking status (current smokers versus
12	current non-smokers) (Ruidavets et al., 2005a).
13	A study examining correlations between O_3 levels and oxidative DNA damage examined
14	results stratified by current versus never and former smokers (Palli et al., 2009). Ozone
15	was positively associated with DNA damage for short-term and long-term exposures
16	among never/former smokers. For current smokers, short-term O_3 concentrations were
17	inversely associated with DNA damage; however, the number of current smokers in the
18	study was small (n = 12).
 19 20 21 22 23 24 25 26 27 	The findings of Palli et al. (2009) were consistent with those from controlled human exposure studies that have confirmed that smokers are less responsive to O ₃ exposure than non-smokers. Spirometric and plethysmographic pulmonary function decline, nonspecific AHR, and inflammatory responses of smokers to O ₃ exposure were all weaker than those reported for non-smokers. Similarly, the time course of development and recovery from these effects, as well as their reproducibility, was not different from non-smokers. Chronic airway inflammation with desensitization of bronchial nerve endings and an increased production of mucus may plausibly explain the pseudo-protective effect of smoking (Frampton et al., 1997a; Torres et al., 1997).
28	These findings for smoking are consistent with the conclusions from previous AQCDs.
29	An epidemiologic study of O_3 -associated DNA damage reported smokers to be less at
30	risk for O_3 -related health effects. In addition, both epidemiologic studies of short-term
31	exposure and CVD outcomes found no evidence of effect measure modification by
32	smoking. No toxicological studies provide biological support for O_3 -related effects.
33	Overall, evidence of potential differences in O_3 -related health effects by smoking status is
34	inadequate due to insufficient coherence and a limited number of studies.

8.4.4 Outdoor Workers

1	Studies included in the 2006 O_3 AQCD reported that individuals who participate in
2	outdoor activities or work outside to be a population at increased risk based on
3	consistently reported associations between O_3 exposure and respiratory health outcomes
4	in these groups (U.S. EPA, 2006b). Outdoor workers are exposed to ambient O_3
5	concentrations for a greater period of time than individuals who spend their days indoors.
6	As discussed in Section $4.3.3$ of this ISA, outdoor workers sampled during the work shift
7	had a higher ratio of personal exposure to fixed-site monitor concentrations than health
8	clinic workers who spent most of their time indoors. Additionally, an increase in dose to
9	the lower airways is possible during outdoor exercise due to both increases in the amount
10	of air breathed (i.e., minute ventilation) and a shift from nasal to oronasal breathing
11	(Sawyer et al., 2007; Nodelman and Ultman, 1999; Hu et al., 1994). For further
12	discussion of the association between FEV_1 responses to O_3 exposure and minute
13	ventilation, refer to Section <u>6.2.3.1</u> of the 2006 O_3 AQCD. A recent study has explored
14	the potential effect measure modification of O_3 exposure and DNA damage by
15	indoor/outdoor workplace (Tovalin et al., 2006). In a study of indoor and outdoor
16	workers in Mexico, individuals who worked outdoors in Mexico City had a slight
17	association between O ₃ exposure and DNA damage (measured by comet tail length
18	assay), whereas no association was observed for indoor workers. However, workers in
19	another Mexican city, Puebla, demonstrated no association between O ₃ levels and DNA
20	damage, regardless of whether they worked indoors or outdoors.
21	Previous studies have shown that increased exposure to O ₃ due to outdoor work leads to
22	increased risk of O_3 -related health effects, specifically decrements in lung function (U.S.
23	EPA, 2006b). Recent evidence from a stratified analysis does not indicate that increased
24	O ₃ exposure due to outdoor work leads to DNA damage. However, the strong evidence
25	from the 2006 O ₃ AQCD which demonstrated increased exposure, dose, and ultimately
26	risk of O ₃ -related health effects in this population supports that there is adequate evidence

26risk of O_3 -related health effects in this population supports that there is adequate evidence27available to indicate that increased exposure to O_3 through outdoor work potentially28increases the risk of O_3 -related health effects.

8.4.5 Air Conditioning Use

29	Air conditioning use is an important component of O_3 exposure, as use of central air
30	conditioning will limit exposure to O_3 by blocking the penetration of O_3 into the indoor
31	environment (see Section $4.3.2$). Air conditioning use is a difficult effect measure
32	modifier to examine in epidemiologic studies because it is often estimated using regional
33	prevalence data and may not reflect individual-level use. More generally, air conditioning

1	prevalence is associated with temperature of a region; those areas with higher
2	temperatures have a greater prevalence of households with air conditioning. Despite these
3	limitations, a few studies have examined effect measure modification by prevalence of air
4	conditioning use in an area. Studies examining multiple cities across the U.S. have
5	assessed whether associations between O3 concentrations and hospital admissions and
6	mortality varied among areas with high and low prevalence of air conditioning. Medina-
7	Ramon et al. (2006) conducted a study during the warm season and observed a greater
8	association between O_3 levels and pneumonia-hospital admissions among areas with a
9	lower proportion of households having central air conditioning compared to areas with a
10	larger proportion of households with air conditioning. However, a similar observation
11	was not observed when examining COPD hospital admissions complicating the
12	interpretation of the results cfrom this study. Bell and Dominici (2008) found evidence of
13	increased risk of O ₃ -related mortality in areas with a lower prevalence of central air
14	conditioning in a study of 98 U.S. communities. Conversely, Medina-Ramón and
15	Schwartz (2008) found that among individuals with atrial fibrillation, a lower risk of
16	mortality was observed for areas with a lower prevalence of central air conditioning.
17	The limited number of studies that examined whether air conditioning use modifies the
18	association between O_3 exposure and health has not provided consistent evidence across

19health endpoints. Therefore, the limited and inconsistent results across epidemiologic20studies has provided inadequate evidence to determine whether a lower prevalence of air21conditioning use leads to a potential increased risk of O3-related health effects.

8.5 Summary

22	In this section, epidemiologic, controlled human exposure, and toxicological studies have
23	been evaluated and indicate that various factors may lead to increased risk of O3-related
24	health effects (<u>Table 8-5</u>).
25	The populations and lifestages identified in this section that have "adequate" evidence for
26	potentially increased O ₃ -related health effects are individuals with asthma, younger and
27	older age groups, individuals with reduced intake of certain nutrients, and outdoor
28	workers, based on consistency in findings across studies and evidence of coherence in
29	results from different scientific disciplines. Asthma as a factor potentially affecting risk
30	was supported by controlled human exposure and toxicological studies, as well as some
31	evidence from epidemiologic studies. Generally, studies of age groups reported positive
32	associations for respiratory hospital admissions and ED visits among children. Biological
33	plausibility for this increased risk is supported by toxicological and clinical research.
34	Also, children have higher exposure and dose due to increased time spent outdoors and

1	ventilation rate. Most studies comparing age groups reported greater effects of short-term
2	O3 exposure on mortality among older adults, although studies of other health outcomes
3	had inconsistent findings regarding whether older adults were at increased risk. Older
4	adults may also withstand greater O3 exposure and not seek relief as quickly as younger
5	adults. Multiple epidemiologic, controlled human exposure, and toxicological studies
6	reported that reduced Vitamins E and C intake are associated with risk of O3-related
7	health effects. Previous studies have shown that increased exposure to O_3 due to outdoor
8	work leads to a potentially increased risk of O3-related health effects and it is clear that
9	outdoor workers have higher exposures, and possibly greater internal doses, of O_3 , which
10	may lead to increased risk of O_3 -related health effects.

Table 8-5Summary of evidence for potential increased risk of ozone-related
health effects.

Evidence Classification	Potential At Risk Factor
Adequate evidence	Asthma (Section 8.2.2)
	Children (Section <u>8.3.1.1</u>)
	Older adults (Section 8.3.1.2)
	Diet (Section 8.4.1)
	Outdoor workers (Section 8.4.4)
Suggestive evidence	Genetic factors (Section 8.1)
	Sex (Section 8.3.2)
	SES (Section 8.3.3)
	Obesity (Section 8.4.2)
Inadequate evidence	Influenza/Infection (Section 8.2.1)
	COPD (Section 8.2.3)
	CVD (Section <u>8.2.4</u>)
	Diabetes (Section 8.2.5)
	Hyperthyroidism (Section <u>8.2.6</u>)
	Race/ethnicity (Section 8.3.4)
	Smoking (Section 8.4.3)
	Air conditioning use (Section 8.4.5)
Evidence of no effect	

11	In some cases, it is difficult to determine a factor that results in potentially increased risk
12	of effects. For example, previous assessments have included controlled human exposure
13	studies in which some healthy individuals demonstrate greater O ₃ -related health effects
14	compared to other healthy individuals. Intersubject variability has been observed for lung
15	function decrements, symptomatic responses, pulmonary inflammation, AHR, and altered
16	epithelial permeability in healthy adults exposed to O_3 (Que et al., 2011; Holz et al.,
17	<u>2005</u> ; <u>McDonnell, 1996</u>). These responses to O_3 exposure in healthy individuals tend to
18	be reproducible within a given individual over a period of several months indicating
19	differences in the intrinsic responsiveness (Holz et al., 2005; Hazucha et al., 2003; Holz
20	<u>et al., 1999;</u> McDonnell et al., 1985b).

1	Limitations include the challenge of evaluating effect measure modification in
2	epidemiologic studies with widespread populations with variation in numerous factors.
3	For a number of the factors described below, there are few available studies. Many
4	toxicological and controlled human exposure studies are the only ones that have
5	examined certain factors and therefore have not been replicated. In considering
6	epidemiologic studies conducted in other countries, it is possible that those populations
7	may differ in SES or other demographic indicators, thus limiting generalizability to a
8	U.S. population. Additionally, many epidemiologic studies that stratify by factors of
9	interest have small sample sizes, which can decrease precision of effect estimates.
10	These challenges and limitations in evaluating the factors that can increase risk for
11	experiencing O ₃ -related health effects may contribute to conclusions that evidence for
12	some factors, such as genetic factors, sex, SES, and obesity provided "suggestive"
13	evidence of potentially increased risk. In addition, for a number of factors listed in
14	Table 8-5 the evidence was inadequate to draw conclusions about potential increase in
15	risk of effects. Overall, the factors most strongly supported as contributing to potentially
16	increased risk of O3-related effects among various populations and lifestages were related
17	to asthma, age group (children and older adults), dietary factors, and working outdoors.

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9 ENVIRONMENTAL EFFECTS: OZONE EFFECTS ON VEGETATION AND ECOSYSTEMS

9.1 Introduction

1	This chapter synthesizes and evaluates the relevant science to help form the scientific
2	foundation for the review of a vegetation- and ecologically-based secondary NAAQS for
3	O ₃ . The secondary NAAQS are based on welfare effects. The Clean Air Act (CAA)
4	definition of welfare effects includes, but is not limited to, effects on soils, water,
5	wildlife, vegetation, visibility, weather, and climate, as well as effects on materials,
6	economic values, and personal comfort and well-being. The effects of O_3 as a greenhouse
7	gas and its direct effects on climate are discussed in Chapter 10 of this document.
8	The intent of the ISA, according to the CAA, is to "accurately reflect the latest scientific
9	knowledge expected from the presence of [a] pollutant in ambient air" (42 U.S.C.7408
10	and 42 U.S.C.7409. This chapter of the ISA includes scientific research from
11	biogeochemistry, soil science, plant physiology, and ecology conducted at multiple levels
12	of biological organization (e.g., organ, organism, population, community, ecosystem).
13	Key information and judgments formerly found in the AQCDs regarding O3 effects on
14	vegetation and ecosystems are found in this chapter. This chapter of the O ₃ ISA serves to
15	update and revise Chapter 9 and AX9 of the 2006 O_3 AQCD (<u>U.S. EPA, 2006b</u>).
1.6	
16	Numerous studies of the effects of O ₃ on vegetation and ecosystems were reviewed in the
17	Numerous studies of the effects of O_3 on vegetation and ecosystems were reviewed in the 2006 O_3 AQCD. That document concluded that the effects of ambient O_3 on vegetation
17 18	2006 O_3 AQCD. That document concluded that the effects of ambient O_3 on vegetation and ecosystems appear to be widespread across the U.S., and experimental studies
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outside of North America that contribute to the understanding of O_3 effects on general ecosystem processes are discussed in the chapter.

3 Sections of this chapter first discuss exposure methods, followed by effects on vegetation 4 and ecosystems at various levels of biological organization and ends with policy-relevant 5 discussions of exposure indices and exposure-response. Figure 9-1 is a simplified 6 illustrative diagram of the major pathway through which O₃ enters plants and the major 7 endpoints O_3 may affect. First, Section <u>9.2</u> presents a brief overview of various 8 methodologies that have been, and continue to be, central to quantifying O_3 effects on 9 vegetation (see AX9.1 of the 2006 O_3 AQCD for more detailed discussion) (U.S. EPA, 10 <u>2006b</u>). Section <u>9.3</u> through Section <u>9.4</u> begin with a discussion of effects at the cellular 11 and subcellular level followed by consideration of the O_3 effects on plant and ecosystem 12 processes (Figure 9-1). In Section 9.3, research is reviewed from the molecular to the 13 biochemical and physiological levels in impacted plants, offering insight into the mode of 14 action of O₃. Section 9.4 provides a review of the effects of O₃ exposure on major 15 endpoints at the whole plant scale including growth, reproduction, visible foliar injury 16 and leaf gas exchange in woody and herbaceous plants in the U.S., as well as a brief 17 discussion of O_3 effects on agricultural crop yield and quality. Section 9.4 also integrates 18 the effects of O_3 on individual plants in a discussion of available research for assessing 19 the effect of O₃ on ecosystems, along with available studies that could inform 20 assessments of various ecosystem services (See Section 9.4.1.2). The development of 21 indices of O_3 exposure and dose modeling is discussed in Section 9.5. Finally, exposure-22 response relationships for a number of tree species, native vegetation, and crop species 23 and cultivars are reviewed, tabulated, and compared in Section 9.6 to form the basis for 24 an assessment of the potential risk to vegetation from current ambient levels of O₃.

1

2

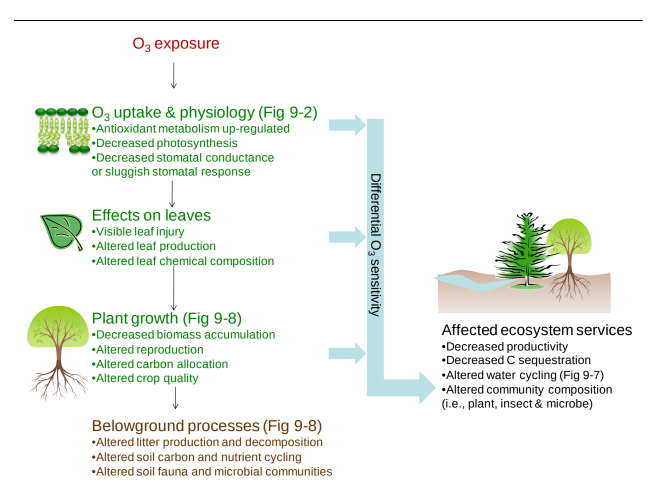


Figure 9-1 An illustrative diagram of the major pathway through which ozone enters plants and the major endpoints that ozone may affect in plants and ecosystems.

9.2 Experimental Exposure Methodologies

9.2.1 Introduction

1A variety of methods for studying plant response to O3 exposures have been developed2over the last several decades. The majority of methodologies currently used have been3discussed in detail in the 1996 O3 AQCD and 2006 O3 AQCD. This section will serve as4a short overview of the methodologies and the reader is referred to the previous O35AQCDs for more in-depth discussion.

9.2.2 "Indoor," Controlled Environment, and Greenhouse Chambers

1	The earliest experimental investigations of the effects of O ₃ on plants utilized simple
2	glass or plastic-covered chambers, often located within greenhouses, into which a flow of
3	O ₃ -enriched air or oxygen could be passed to provide the exposure. The types, shapes,
4	styles, materials of construction, and locations of these chambers have been numerous.
5	Hogsett et al. (1987a) have summarized the construction and performance of more
6	elaborate and better instrumented chambers since the 1960s, including those installed in
7	greenhouses (with or without some control of temperature and light intensity).
8	One greenhouse chamber approach that continues to yield useful information on the
9	relationships of O_3 uptake to both physiological and growth effects employs continuous
10	stirred tank reactors (CSTRs) first described by Heck et al. (1978). Although originally
11	developed to permit mass-balance studies of O ₃ flux to plants, their use has more recently
12	widened to include short-term physiological and growth studies of $O_3 \times CO_2$ interactions
13	(Loats and Rebbeck, 1999; Reinert et al., 1997; Rao et al., 1995; Reinert and Ho, 1995;
14	Heagle et al., 1994a), and validation of visible foliar injury on a variety of plant species
15	(Kline et al., 2009; Orendovici et al., 2003). In many cases, supplementary lighting and
16	temperature control of the surrounding structure have been used to control or modify the
17	environmental conditions (<u>Heagle et al., 1994a</u>).
18	Many investigations have utilized commercially available controlled environment
19	chambers and walk-in rooms adapted to permit the introduction of a flow of O_3 into the
20	controlled air-volume. Such chambers continue to find use in genetic screening and in
21	physiological and biochemical studies aimed primarily at improving the understanding of
22	modes of action. For example, some of the studies of the O_3 responses of common
23	plantain (Plantago major) populations have been conducted in controlled environment
24	chambers (Whitfield et al., 1996; Reiling and Davison, 1994).
25	More recently, some researchers have been interested in attempting to investigate direct
26	O_3 effects on reproductive processes, separate from the effects on vegetative processes
27	(Black et al., 2010). For this purpose, controlled exposure systems have been employed
28	to expose the reproductive structures of annual plants to gaseous pollutants independently
29	of the vegetative component (Black et al., 2010; Stewart et al., 1996).

9.2.3 Field Chambers

30	In general, field chamber studies are dominated by the use of various versions of the open
31	top chamber (OTC) design, first described by Heagle et al. (1973) and Mandl et al.
32	(1973). The OTC method continues to be a widely used technique in the U.S. and Europe

- 1 for exposing plants to varying levels of O_3 . Most of the new information confirms earlier 2 conclusions and provides additional support for OTC use in assessing plant species and in 3 developing exposure-response relationships. Chambers are generally ~3 meters in 4 diameter with 2.5 meter-high walls. Hogsett et al. (1987b) described in detail many of the 5 various modifications to the original OTC designs that appeared subsequently, e.g., the 6 use of larger chambers for exposing small trees (Kats et al., 1985) or grapevines (Mandl 7 et al., 1989), the addition of a conical baffle at the top to improve ventilation (Kats et al., 8 1976), a frustum at the top to reduce ambient air incursions, and a plastic rain-cap to 9 exclude precipitation (Hogsett et al., 1985). All versions of OTCs included the discharge 10 of air via ports in annular ducting or interiorly perforated double-layered walls at the base 11 of the chambers to provide turbulent mixing and the upward mass flow of air.
- 12 Chambered systems, including OTCs, have several advantages. For instance, they can 13 provide a range of treatment levels including charcoal-filtered (CF), clean-air control, and 14 several above ambient concentrations for O_3 experiments. Depending on experimental 15 intent, a replicated, clean-air control treatment is an essential component in many 16 experimental designs. The OTC can provide a consistent, definable exposure because of 17 the constant wind speed and delivery systems. Statistically robust concentration-response (C-R) functions can be developed using such systems for evaluating the implications of 18 19 various alternative air quality scenarios on vegetation response. Nonetheless, there are 20 several characteristics of the OTC design and operation that can lead to exposures that 21 might differ from those experienced by plants in the field. First, the OTC plants are 22 subjected to constant air flow turbulence, which, by lowering the boundary layer 23 resistance to diffusion, may result in increased uptake. This may lead to an 24 overestimation of effects relative to areas with less turbulence (Krupa et al., 1995; Legge 25 et al., 1995). However, other research has found that OTC's may slightly change vapor 26 pressure deficit (VPD) in a way that may decrease the uptake of O_3 into leaves (Piikki et 27 al., 2008a). As with all methods that expose vegetation to modified O_3 concentrations in 28 chambers, OTCs create internal environments that differ from ambient air. This so-called 29 "chamber effect" refers to the modification of microclimatic variables, including reduced 30 and uneven light intensity, uneven rainfall, constant wind speed, reduced dew formation, 31 and increased air temperatures (Fuhrer, 1994; Manning and Krupa, 1992). However, in at 32 least one case where canopy resistance was quantified in OTCs and in the field, it was 33 determined that gaseous pollutant exposure to crops in OTCs was similar to that which 34 would have occurred at the same concentration in the field (Unsworth et al., 1984a, b). 35 Because of the standardized methodology and protocols used in National Crop Loss 36 Assessment Network (NCLAN) and other programs, the database can be assumed to be 37 internally consistent.

1	While it is clear that OTCs can alter some aspects of the microenvironment and plant
	· · ·
2	growth, it is important to establish whether or not these differences affect the relative
3	response of a plant to O ₃ . As noted in the 1996 O ₃ AQCD, evidence from a number of
4	comparative studies of OTCs and other exposure systems suggested that responses were
5	essentially the same regardless of exposure system used and chamber effects did not
6	significantly affect response. In studies that included exposure to ambient concentrations
7	of O_3 in both OTCs, and open-air, chamberless control plots, responses in the OTCs were
8	the same as in open-air plots. Examples include studies of tolerant and sensitive white
9	clover clones (Trifolium repens) to ambient O ₃ in greenhouse, open top, and ambient
10	plots (Heagle et al., 1996), Black Cherry (Prunus serotina) (Neufeld et al., 1995), and
11	three species of conifers (Neufeld et al., 2000). Experimental comparisons between
12	exposure methodologies are reviewed in Section $9.2.6$.
13	Another type of field chamber called a "terracosm" has been developed and used in
14	recent studies (Lee et al., 2009a). Concern over the need to establish realistic plant-litter-
15	soil relationships as a prerequisite to studies of the effects of O_3 and CO_2 enrichment on
16	ponderosa pine (Pinus ponderosa) seedlings led Tingey et al. (1996) to develop closed,
17	partially environmentally controlled, sun-lit chambers ("terracosms") incorporating
18	lysimeters (1 meter deep) containing forest soil in which the appropriate horizon structure
19	was retained.
20	Other researchers have recently published studies using another type of out-door chamber
21	called recirculating Outdoor Plant Environment Chambers (OPECs) (Flowers et al.,
22	<u>2007</u>). These closed chambers are approximately 2.44 meters $\times 1.52$ meters with a growth
23	volume of approximately 3.7 m ³ in each chamber. These chambers admit 90% of full
24	sunlight and control temperature, humidity and vapor pressure (Fiscus et al., 1999).

9.2.4 Plume and FACE-Type Systems

25	Plume systems are chamberless exposure facilities in which the atmosphere surrounding
26	plants in the field is modified by the injection of pollutant gas into the air above or
27	around them from multiple orifices spaced to permit diffusion and turbulence, so as to
28	establish relatively homogeneous conditions as the individual plumes disperse and mix
29	with the ambient air. They can only be used to increase the O_3 levels in the ambient air.
30	The most common plume system used in the U.S. is a modification of the free-air carbon
31	dioxide/ozone enrichment (FACE) system (Hendrey et al., 1999; Hendrey and Kimball,
32	<u>1994</u>). Although originally designed to provide chamberless field facilities for studying
33	the CO ₂ effects of climate change, FACE systems have been adapted to include the
34	dispensing of O_3 (Karnosky et al., 1999). This method has been employed in Illinois

1 (SoyFACE) to study soybeans (Morgan et al., 2004; Rogers et al., 2004) and in 2 Wisconsin (Aspen FACE) to study trembling aspen (Populus tremuloides), birch (Betula 3 papyrifera) and maple (Acer saccharum) (Karnosky et al., 1999). Volk et al. (2003) 4 described a similar system for exposing grasslands that uses 7-m diameter plots. Another 5 similar FACE system has been used in Finland (Saviranta et al., 2010; Oksanen, 2003). 6 The FACE systems in the U.S. discharge the pollutant gas (O₃ and/or CO₂) through 7 orifices spaced along an annular ring (or torus) or at different heights on a ring of vertical 8 pipes. Computer-controlled feedback from the monitoring of gas concentration regulates 9 the feed rate of enriched air to the dispersion pipes. Feedback of wind speed and 10 directional information ensures that the discharges only occur upwind of the treatment plots, and that discharge is restricted or closed down during periods of low wind speed or 11 12 calm conditions. The diameter of the arrays and their height (25-30 m) in some FACE 13 systems requires large throughputs of enriched air per plot, particularly in forest tree 14 systems. The cost of the throughputs tends to limit the number of enrichment treatments, 15 although Hendrey et al. (1999) argued that the cost on an enriched volume basis is 16 comparable to that of chamber systems. 17 A different FACE-type facility has been developed for the Kranzberg Ozone Fumigation 18 Experiment (KROFEX) in Germany beginning in 2000 (Nunn et al., 2002; Werner and 19 Fabian, 2002). The experiment aims to study the effects of O_3 on mature stands of beech 20 (*Fagus sylvatica*) and spruce (*Picea abies*) trees in a system that functions independently 21 of wind direction. The enrichment of a large volume of the ambient air immediately 22 above the canopy takes place via orifices in vertical tubes suspended from a horizontal 23 grid supported above the canopy. 24 Although plume systems make virtually none of the modifications to the physical 25 environment that are inevitable with chambers, their successful use depends on selecting 26 the appropriate numbers, sizes, and orientations of the discharge orifices to avoid 27 "hot-spots" resulting from the direct impingement of jets of pollutant-enriched air on 28

plant foliage (Werner and Fabian, 2002). Because mixing is unassisted and completely 29 dependent on wind turbulence and diffusion, local gradients are inevitable especially in 30 large-scale systems. FACE systems have provisions for shutting down under low wind 31 speed or calm conditions and for an experimental area that is usually defined within a 32 generous border in order to strive for homogeneity of the exposure concentrations within 33 the treatment area. They are also dependent upon continuous computer-controlled 34 feedback of the O₃ concentrations in the mixed treated air and of the meteorological 35 conditions. Plume and FACE systems also are unable to reduce O₃ levels below ambient 36 in areas where O_3 concentrations are phytotoxic.

9.2.5 Ambient Gradients

1 2 3 4 5 6 7 8	Ambient O ₃ gradients that occur in the U.S. hold potential for the examination of plant responses over multiple levels of exposure. However, few such gradients can be found that meet the rigorous statistical requirements for comparable site characteristics such as soil type, temperature, rainfall, radiation, and aspect (Manning and Krupa, 1992); although with small plants, soil variability can be avoided by the use of plants in large pots. The use of soil monoliths transported to various locations along natural O ₃ gradients is another possible approach to overcome differences in soils; however, this approach is also limited to small plants.
9	Studies in the 1970s used the natural gradients occurring in southern California to assess
10	yield losses of alfalfa and tomato (Oshima et al., 1977; Oshima et al., 1976). A transect
11	study of the impact of O_3 on the growth of white clover and barley in the U.K. was
12	confounded by differences in the concurrent gradients of SO_2 and NO_2 pollution
13	(Ashmore et al., 1988). Studies of forest tree species in national parks in the eastern U.S.
14	(Winner et al., 1989) revealed increasing gradients of O_3 and visible foliar injury with
15	increased elevation.
16	Several studies have used the San Bernardino Mountains Gradient Study in southern
17	California to study the effects of O_3 and N deposition on forests dominated by ponderosa
18	and Jeffrey pine (Jones and Paine, 2006; Arbaugh et al., 2003; Grulke, 1999; U.S. EPA,
19	<u>1977</u>). However, it is difficult to separate the effects of N and O_3 in some instances in
20	these studies (Arbaugh et al., 2003). An O_3 gradient in Wisconsin has been used to study
21	foliar injury in a series of trembling aspen clones (Populus tremuloides) differing in O ₃
22	sensitivity (<u>Maňkovská et al., 2005; Karnosky et al., 1999</u>). Also in the Midwest, an
23	east-west O_3 gradient around southern Lake Michigan was used to look at growth and
24	visible foliar injury in (P. serotina) and common milkweed (Asclepias syriaca) (Bennett
25	<u>et al., 2006</u>).
26	More recently, studies have been published that have used natural gradients to study a
27	variety of endpoints and species. For example, Gregg et al. (2003) studied cottonwood
28	(<i>Populus deltoides</i>) saplings grown in an urban to rural gradient of O_3 by using seven
29	locations in the New York City area. The secondary nature of the reactions of O_3
30	formation and NO _X titration reactions within the city center resulted in significantly
31	higher cumulative O ₃ exposures in more rural sites. Potential modifying factors such as
32	soil composition, moisture, or temperature were either controlled or accounted for in
33	analysis. As shown in Section <u>9.6.3.3</u> , the response of this species to O_3 exposure was
34	much stronger than most species. The natural gradient exposures were reproduced in
35	parallel using OTCs, and yielded similar results. Also, the U.S. Forest Service Forest
36	Inventory and Analysis (FIA) program uses large-scale O ₃ exposure patterns across the

continental U.S. to study occurrences of foliar injury due to O₃ exposure (<u>Smith et al.</u>,
 <u>2003</u>) (Section <u>9.4.2</u>). Finally, <u>McLaughlin et al.</u> (<u>2007a</u>); <u>2007b</u>) used spatial and
 temporal O₃ gradients to study forest growth and water use in the southern Appalachians.
 These studies found varying O₃ exposures between years and between sites.

9.2.6 Comparative Studies

5 All experimental approaches used to expose plants to O_3 have strengths and weaknesses. 6 One potential weakness of laboratory, greenhouse, or field chamber studies is the 7 potential effect of the chamber on micrometeorology. In contrast, plume, FACE and 8 gradient systems are limited by the very small number of possible exposure levels 9 (almost always no more than two), small replication and the inability to reduce O_3 levels 10 below ambient. In general, experiments that aim at characterizing the effect of a single 11 variable, e.g., exposure to O_3 , must not only manipulate the levels of that variable, but 12 also control potentially interacting variables and confounders, or else account for them. 13 However, while increasing control of environmental variables makes it easier to discern 14 the effect of the variable of interest, it must be balanced with the ability to extend 15 conclusions to natural, non-experimental settings. More naturalistic exposure systems, on 16 the other hand, let interacting factors vary freely, resulting in greater unexplainable 17 variability. The various exposure methodologies used with O_3 vary in the balance each 18 strikes between control of environmental inputs, closeness to the natural environment, 19 noisiness of the response data, and ability to make general inferences.

20 Studies have examined the comparability of results obtained though the various exposure 21 methodologies. As noted in the 1996 O₃ AQCD, evidence from the comparative studies 22 of OTCs and from closed chamber and O₃-exclusion exposure systems on the growth of 23 alfalfa (Medicago sativa) by Olszyk et al. (1986) suggested that, since significant 24 differences were found for fewer than 10% of the growth parameters measured, the 25 responses were, in general, essentially the same regardless of exposure system used, and 26 chamber effects did not significantly affect response. In 1988, Heagle et al. (1988) 27 concluded: "Although chamber effects on yield are common, there are no results showing 28 that this will result in a changed yield response to O_3 ." A study of the effects of an 29 enclosure examined the responses of tolerant and sensitive white clover clones (Trifolium 30 *repens*) to ambient O_3 in a greenhouse, open-top chamber, and ambient (no chamber) 31 plots (<u>Heagle et al., 1996</u>). For individual harvests, greenhouse O_3 exposure reduced the 32 forage weight of the sensitive clone 7 to 23% more than in OTCs. However, the response 33 in OTCs was the same as in ambient plots. Several studies have shown very similar 34 response of yield to O_3 for plants grown in pots or in the ground, suggesting that even

1	such a significant change in environment does not alter the proportional response to O ₃ ,
2	providing that the plants are well watered (Heagle et al., 1983; Heagle, 1979).
3	A few recent studies have compared results of O3 experiments between OTCs, FACE
4	experiments, and gradient studies. For example, a series of studies undertaken at Aspen
5	FACE (Isebrands et al., 2001; Isebrands et al., 2000) showed that O ₃ symptom expression
6	was generally similar in OTCs, FACE, and ambient O ₃ gradient sites, and supported the
7	previously observed variation among trembling aspen clones using OTCs (Maňkovská et
8	al., 2005; Karnosky et al., 1999). In the SoyFACE experiment in Illinois, soybean
9	(Pioneer 93B15 cultivar) yield loss data from a two-year study was published (Morgan et
10	al., 2006). This cultivar is a recent selection and, like most modern cultivars, has been
11	selected under an already high current O_3 exposure. It was found to have average
12	sensitivity to O ₃ compared to 22 other cultivars tested at SoyFACE. In this experiment,
13	ambient hourly O_3 concentrations were increased by approximately 20% and measured
14	yields were decreased by 15% in 2002 as a result of the increased O_3 exposure (Morgan
15	et al., 2006). To compare these results to chamber studies, Morgan et al. (2006)
16	calculated the expected yield loss from a linear relationship constructed from chamber
17	data using seven-hour seasonal averages (Ashmore, 2002). They calculated an 8%
18	expected yield loss from the 2002 O_3 exposure using that linear relationship. As reported
19	in Section <u>9.2.5</u> , <u>Gregg et al. (2006</u> , <u>2003</u>) found similar O ₃ effects on cottonwood
20	sapling biomass growth along an ambient O_3 gradient in the New York City area and a
21	parallel OTC study.
22	Finally, EPA conducted comparisons of exposure-response model predictions based on
23	OTC studies, and more recent FACE observations. These comparisons include yield of
24	annual crops, and biomass growth of trees. They are presented in Section 9.6.3 of this

9.3 Mechanisms Governing Vegetation Response to Ozone

9.3.1 Introduction

document.

25

26	This section focuses on the effects of O ₃ stress on plants and their responses to that stress
27	on the molecular, biochemical and physiological levels. First, the pathway of O_3 uptake
28	into the leaf and the initial chemical reactions occurring in the substomatal cavity and
29	apoplast will be described (Section $9.3.2$); additionally, direct effects of O_3 on the
30	stomatal apparatus will be discussed. Once O_3 has entered the substomatal cavity and
31	apoplast, it is thought that the cell must be able to detect the presence of O_3 or its

1	breakdown products in order to initiate the rapid changes in signaling pathways and gene
2	expression that have been measured in O3-treated plants. While it remains unclear exactly
3	how O ₃ and/or its breakdown products are detected in the apoplast and how that leads to
4	signaling of oxidative stress in plants, much progress has been made in examining several
5	different mechanisms that may contribute to detecting the presence of O ₃ and its
6	breakdown products, and also initiating a signal transduction cascade, which will be
7	described in Section 9.3.3.1. The next section focuses on changes in gene and protein
8	expression measured in plants exposed to O ₃ , with particular emphasis on results from
9	transcriptome (all RNA molecules produced in a cell) and proteome (all proteins
10	produced in a cell) analyses (Section <u>9.3.3.2</u>). Subsequently, the role of phytohormones
11	such as salicylic acid (SA), ethylene (ET), jasmonic acid (JA), and abscisic acid (ABA)
12	and their interactions in both signal transduction processes and in determining plant
13	response to O_3 is discussed in Section <u>9.3.3.3</u> . After O_3 uptake, some plants can respond
14	to the oxidative stress with detoxification to minimize damage. These mechanisms of
15	detoxification, with particular emphasis on antioxidant enzymes and metabolites, are
16	reviewed in Section 9.3.4. The next section focuses on changes in primary and secondary
17	metabolism in plants exposed to O ₃ , looking at photosynthesis, respiration and several
18	secondary metabolites, some of which may also act as antioxidants and protect the plant
19	from oxidative stress (Section $9.3.5$). For many of these topics, information from the
20	2006 O ₃ AQCD has been summarized, as this information is still valid and supported by
21	more recent findings. For other topics, such as genomics and proteomics, which have
22	arisen due to the availability of new technologies, the information is based solely on new
23	publications with no reference to the 2006 O_3 AQCD.
24	As Section <u>9.3</u> focuses on mechanisms underlying effects of O_3 on plants and their

2 25 response to it, the conditions that are used to study these mechanisms do not always 26 reflect conditions that a plant may be exposed to in an agricultural setting or natural 27 ecosystem. The goal of many of these studies is to generate an O₃ effect in a relatively 28 short period of time and not always to simulate ambient O₃ exposures. Therefore, plants 29 are often exposed to unrealistically high O₃ concentrations for several hours or days 30 (acute exposure), and only in some cases to ambient or slightly elevated O₃ 31 concentrations for longer time periods (chronic exposure). Additionally, the plant species 32 utilized in these studies are often not agriculturally important or commonly found as part 33 of natural ecosystems. Model organisms such as Arabidopsis thaliana are used frequently 34 as they are easy to work with, and mutants or transgenic plants are easy to develop or 35 have already been developed. Furthermore, the Arabidopsis genome has been sequenced, 36 and much is known about the molecular basis of many biochemical and cellular 37 processes.

1	Many of the studies described in this section focus on changes in the expression of genes
2	in O3-treated plants. Changes in gene expression (i.e., either upregulation or
3	downregulation of gene expression) do not always translate into changes in protein
4	quantity and/or activity, as there are many levels of post-transcriptional and post-
5	translational modifications which impact protein quantity and activity. Many studies do
6	not evaluate whether the observed changes in gene expression lead to changes at the
7	protein level and, therefore, it is not always clear if the changes in gene expression
8	represent a meaningful biological response to O_3 exposure. However, with the advent of
9	proteomics, some very recent studies have evaluated changes in protein expression for
10	large numbers of proteins in O ₃ treated plants, and the findings from these studies support
11	the previous results regarding changes in gene expression studies as a result of O_3
12	exposure. The next step in the process is to determine the implications of the measured
13	changes occurring at the cellular level to whole plants and ecosystems, which is an
14	important topic of study which has not been widely addressed.
15	The most noteworthy new body of research since the 2006 O_3 AQCD is on the
16	understanding of molecular mechanisms underlying how plants are affected by O ₃ ; many
17	of the recent studies reviewed here focus on changes in gene expression in plants exposed
18	to elevated O ₃ . The findings summarized in the 2006 O ₃ AQCD included decreases in

- 18to elevated O3. The findings summarized in the 2006 O3 AQCD included decreases in19transcript levels of photosynthesis associated genes, and increases in transcript levels of20genes encoding for pathogenesis-related proteins, enzymes needed for ethylene synthesis,21antioxidant enzymes and defense genes such as phenylalanine ammonia lyase in plants22exposed to O3. These findings have been supported by the new studies, and the advent of23new technologies has allowed for a more comprehensive understanding of the24mechanisms governing how plants are affected by O3.
- 25In summary, these new studies have increased knowledge of the molecular, biochemical26and cellular mechanisms occurring in plants in response to O3 by often using artificial27exposure conditions and model organisms. This information adds to the understanding of28the basic biology of how plants are affected by oxidative stress in the absence of any29other potential stressors. The results of these studies provide important insights, even30though they may not always directly translate into effects observed in other plants under31more realistic exposure conditions.

9.3.2 Ozone Uptake into the Leaf

32	Appendix AX9.2.3 of the 2006 O_3 AQCD clearly described the process by which O_3
33	enters plant leaves through open stomata (U.S. EPA, 2006b). This information continues
34	to be valid and is only summarized here.

1	Stomata provide the principal pathway for O_3 to enter and affect plants (Massman and
2	Grantz, 1995; Fuentes et al., 1992; Reich, 1987; Leuning et al., 1979). Ozone moves into
3	the leaf interior by diffusing through open stomata, and environmental conditions which
4	promote high rates of gas exchange will favor the uptake of the pollutant by the leaf.
5	Factors that may limit uptake include boundary layer resistance and the size of the
6	stomatal aperture (Figure 9-2) (U.S. EPA, 2006b). Once inside the substomatal cavity, O_3
7	is thought to rapidly react with the aqueous apoplast to form breakdown products known
8	as reactive oxygen species (ROS), such as hydrogen peroxide (H_2O_2), superoxide (O_2^-),
9	hydroxyl radicals (HO) and peroxy radicals (HO ₂) (Figure 9-3). Hydrogen peroxide is
10	not only a toxic breakdown product of O ₃ , but has been shown to function as a signaling
11	molecule, which is activated in response to both biotic and abiotic stressors. The role of
12	H ₂ O ₂ in signaling was described in detail in the 2006 O ₃ AQCD. Additional organic
13	molecules present in the apoplast or cell wall, such as those containing double bonds or
14	sulfhydryls that are sensitive to oxidation, could also be converted to oxygenated
15	molecules after interacting with O_3 (Figure 9-4). These reactions are not only pH
16	dependent, but are also influenced by the presence of other molecules in the apoplast
17	(U.S. EPA, 2006b). The 2006 O_3 AQCD provided a comprehensive summary of these
18	possible interactions of O_3 with other biomolecules (U.S. EPA, 2006b). It is in the
19	apoplast that initial detoxification reactions by antioxidant metabolites and enzymes take
20	place, and these initial reactions are critical to reduce concentrations of the oxidative
21	breakdown products of O_3 ; these reactions are described in more detail in Section <u>9.3.4</u> of
22	this document.

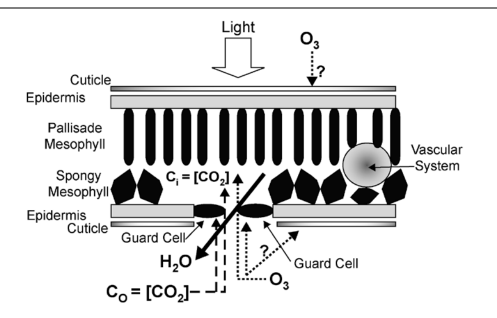
9.3.2.1 Changes in Stomatal Function

23	Ozone-induced changes in stomatal conductance have been reviewed in detail in previous
24	O ₃ AQCDs. The findings summarized in these documents demonstrate that stomatal
25	conductance is often reduced in plants exposed to O ₃ , resulting either from a direct
26	impact of O_3 on the stomatal complex which causes closure, or as a response to
27	increasing CO_2 concentrations in the substomatal cavity as carbon fixation is reduced.
28	Although the nature of these effects depends upon many different factors, including the
29	plant species, concentration and duration of the O3 exposure, and prevailing
30	meteorological conditions, stomatal conductance is often negatively affected by plant
31	exposure to O_3 (Wittig et al., 2007). Decreases in conductance have been shown to result
32	from direct as well as indirect effects on stomata (Wittig et al., 2007). Results from the
33	use of Arabidopsis mutants and new technologies, which allow for analysis of guard cell
34	function in whole plants rather than in isolated guard cells or epidermal peels, suggest
35	that O_3 may also have a direct impact on stomatal guard cells, leading to alterations in

1 stomatal conductance. The use of a new simultaneous O_3 exposure/gas exchange device 2 has demonstrated that exposure of Arabidopsis ecotypes Col-0 and Ler to 150 ppb O_3 3 resulted in a 60-70% decline in stomatal conductance within 9-12 minutes of beginning 4 the exposure. Twenty to thirty minutes later, stomatal conductance had returned to its 5 initial value, even with continuing exposure to O_3 , indicating a rapid direct effect of O_3 6 on stomatal function (Kollist et al., 2007). This transient decrease in stomatal 7 conductance was not observed in the abscisic acid insensitive (ABI2) Arabidopsis 8 mutant. As the ABI2 protein is thought to regulate the signal transduction process 9 involved in stomatal response downstream of ROS production, the authors suggest that 10 the transient decrease in stomatal conductance in the Col-0 and Ler ecotypes results from 11 the biological action of ROS in transducing signals, rather than direct physical damage to 12 guard cells by ROS (Kollist et al., 2007). This rapid transient decrease in stomatal 13 conductance was also not observed when exposing the Arabidopsis mutant slac1 (slow 14 anion channel-associated 1) to 200 ppb O_3 (Vahisalu et al., 2008). The SLAC1 protein 15 was shown to be essential for guard cell slow anion channel functioning and for stomatal 16 closure in response to O₃. Based on additional studies using a variety of Arabidopsis 17 mutants impaired in various aspects of stomatal function, Vahisalu et al. (2008) suggest 18 that the presence of ROS in the guard cell apoplast (formed either by O₃ breakdown or 19 through ROS production from NADPH oxidase activity) leads to the activation of a 20 signaling pathway in the guard cells, which includes SLAC1, and results in stomatal 21 closure.

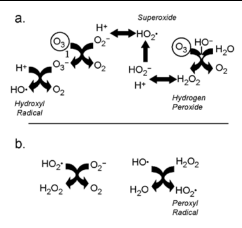
22 A review by McAinsh et al. (2002) discusses the role of calcium as a part of the signal 23 transduction pathway involved in regulating stomatal responses to pollutant stress. A 24 number of studies in this review provide some evidence that exposure to O_3 increases the cytosolic free calcium concentration ($[Ca^{2+}]$ cyt) in guard cells, which may result in an 25 26 inhibition of the plasma membrane inward-rectifying K^+ channels in guard cells, which 27 allow for the K^+ uptake needed for stomatal opening (McAinsh et al., 2002; Torsethaugen 28 et al., 1999). This would compromise the ability of the stomata to respond to various 29 stimuli, including light, CO₂ concentration and drought. Pei et al. (2000) reported that the presence of H₂O₂ activated Ca²⁺ -permeable channels, which mediate increases in 30 31 $[Ca^{2+}]$ cyt in guard cell plasma membranes of Arabidopsis. They also determined that abscisic acid (ABA) induced H₂O₂ production in guard cells, leading to ABA-induced 32 stomatal closure via activation of the membrane Ca^{2+} channels. Therefore, it is possible 33 that H_2O_2 , a byproduct of O_3 breakdown in the apoplast, could disrupt the Ca²⁺-ABA 34 35 signaling pathway that is involved in regulating stomatal responses (McAinsh et al., 36 2002). The studies described here provide some evidence to suggest that O_3 and its 37 breakdown products can directly affect stomatal functioning by impacting the signal 38 transduction pathways which regulate guard cells. Stomatal sluggishness has been 39 described as a delay in stomatal response to changing environmental conditions in

1sensitive species exposed to higher concentrations and/or longer-term O3 exposures2(Paoletti and Grulke, 2010, 2005; McAinsh et al., 2002). It is possible that the signaling3pathways described above could be involved in mediating this stomatal sluggishness in4some plant species under certain O3 exposure conditions (Paoletti and Grulke, 2005;5McAinsh et al., 2002).



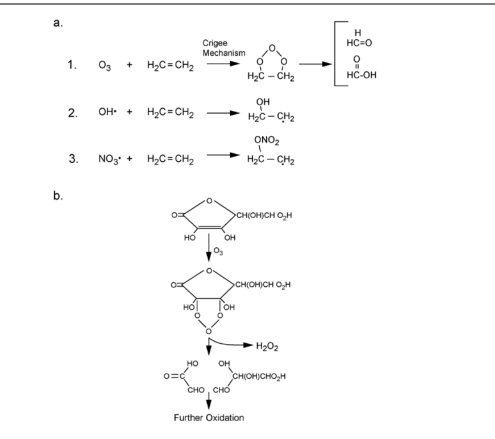
Note: While details among species vary, the general overview remains the same. Light that drives photosynthesis generally falls upon the upper (adaxial) leaf surface. Carbon dioxide and ozone enter through the stomata on the lower (abaxial) leaf surface, while water vapor exits through the stomata (transpiration).

Figure 9-2 The microarchitecture of a dicot leaf.



Note: (a) Ozone reacts at the double bonds to form carbonyl groups. (b) Under certain circumstances, peroxides are generated.

Figure 9-3 **Possible reactions of ozone within water.**



Note: (a) The typical Crigee mechanism is shown in which several reaction paths from the initial product are shown. (b) Typical reaction of ascorbic acid with ozone. Source: Adapted from Mudd (1996).

Figure 9-4 The Crigee mechanism of ozone attack of a double bond.

9.3.3 Cellular to Systemic Responses

9.3.3.1 Ozone Detection and Signal Transduction

1	New technologies allowing for large-scale analysis of oxidative stress-induced changes in
2	gene expression have facilitated the study of signal transduction processes associated
3	with the perception and integration of responses to the stress. Many of these studies have
4	been conducted using Arabidopsis or tobacco plants, for which a variety of mutants are
5	available and/or which can be easily genetically modified to generate either loss-of-
6	function or over-expressing genotypes. Several comprehensive review articles provide an
7	overview of what is known of O ₃ -induced signal transduction processes and how they
8	may help to explain differential sensitivity of plants to the pollutant (Ludwikow and
9	Sadowski, 2008; Baier et al., 2005; Kangasjarvi et al., 2005). Additionally, analysis of
10	several studies of transcriptome changes has also allowed for the compilation of these
11	data to determine an initial time-course for O ₃ -induced activation of various signaling
12	compounds (<u>Kangasjarvi et al., 2005</u>).
13	A number of different mechanisms for detection of O ₃ by plants have been proposed;
14	however, there is still much that is not known about this process. Some of the earliest
15	events that occur in plants exposed to O_3 have been described in the guard cells of
16	stomata. Reactive oxygen species were observed in the chloroplasts of guard cells in the
17	O ₃ tolerant Col-0 Arabidopsis thaliana ecotype plants within 5 minutes of plant exposure
18	to 350 ppb O_3 (Joo et al., 2005). Reactive oxygen species from the breakdown of O_3 in
19	the apoplast are believed to activate GTPases (G-proteins), which, in turn, activate
20	several intracellular sources of ROS, including ROS derived from the chloroplasts.
21	G-proteins are also believed to play a role in activating membrane-bound NADPH
22	oxidases to produce ROS and, as a result, propagate the oxidative burst to neighboring
23	cells (Joo et al., 2005). Therefore, G-proteins are recognized as important molecules
24	involved in plant responses to O_3 and may play a role in detecting the presence of ROS
25	from the breakdown of O_3 in the apoplast (<u>Kangasjarvi et al., 2005</u> ; <u>Booker et al., 2004a</u>).
26	A change in the redox state of the plant and the oxidation of sensitive molecules in itself
27	may represent a means of perception and signaling of oxidative stress in plants.
28	Disulfide-thiol conversions in proteins and the redox state of the glutathione pool may be
29	important components of redox detection and signal transduction (Foyer and Noctor,
30	<u>2005a</u> , <u>b</u>).
31	Calcium (Ca^{2+}) has also been implicated in the transduction of signals to the nucleus in
32	response to oxidative stress. The influx of Ca^{2+} from the apoplast into the cell occurs
33	early during plant exposure to O_3 , and it is thought to play a role in regulating the activity

- 1of protein kinases, which are discussed below (Baier et al., 2005; Hamel et al., 2005).2Calcium channel blockers inhibited O_3 -induced activation of protein kinases in tobacco3suspension cells exposed to 500 ppb O_3 for 10 minutes, indicating that the opening of4 Ca^{2+} channels is an important upstream signaling event or that the (as yet unknown)5upstream process has a requirement for Ca^{2+} (Samuel et al., 2000).
- Further transmission of information regarding the presence of ROS to the nucleus 6 7 involves mitogen-activated protein kinases (MAPK), which phosphorylate proteins and 8 activate various cellular responses (Hamel et al., 2005). Mitogen-activated protein 9 kinases are induced in several different plant species in response to O_3 exposure, 10 including tobacco (Samuel et al., 2005), Arabidopsis (Ludwikow et al., 2004), the shrub 11 Phillyrea latifolia (Paolacci et al., 2007) and poplar (Hamel et al., 2005). Disruption of 12 these signal transduction pathways by over-expressing or suppressing MAPK activity in 13 different Arabidopsis and tobacco lines resulted in increased plant sensitivity to O_3 (Miles 14 et al., 2005; Samuel and Ellis, 2002). Additionally, greater O₃ tolerance of several 15 Arabidopsis ecotypes was correlated with greater upregulation of MAPK signaling 16 pathways upon O₃ exposure than in more sensitive Arabidopsis ecotypes (Li et al., 17 2006b; Mahalingam et al., 2006; Overmyer et al., 2005), indicating that determination of 18 plant sensitivity and plant response to O_3 may, in part, be determined not only by whether 19 these pathways are turned on, but also by the magnitude of the signals moving through 20 these communication channels.
- 21In conclusion, experimental evidence suggests that there are likely several different22mechanisms by which the plant detects the presence of O3 or its breakdown products.23These mechanisms may vary by species or developmental stage of the plant, or may24co-exist and be activated by different exposure conditions. Calcium and protein kinases25are likely involved in relaying information about the presence of the stressor to the26nucleus and other cellular compartments as a first step in determining whether and how27the plant will respond to the stress.

9.3.3.2 Gene and Protein Expression Changes in Response to Ozone

The advent of DNA microarray technology has allowed for the study of gene expression in cells on a large scale. Rather than assessing changes in gene expression of individual genes, DNA microarrays facilitate the evaluation of entire transcriptomes, providing a comprehensive picture of simultaneous alterations in gene expression. In addition, these studies have provided more insight into the complex interactions between molecules, how those interactions lead to the communication of information in the cell (or between

1	neighboring cells), and which role these interactions play in determining tolerance or
2	sensitivity and how a plant may respond to stresses such as O ₃ (Ludwikow and Sadowski,
3	<u>2008</u>). Transcriptome analysis of O_3 -treated plants has been performed in several species,
4	including Arabidopsis thaliana (Li et al., 2006b; Tosti et al., 2006; Heidenreich et al.,
5	2005; Mahalingam et al., 2005; Tamaoki et al., 2003), pepper (Capsicum annuum) (Lee
6	and Yun, 2006), clover (Medicago truncatula) (Puckette et al., 2008), Phillyrea latifolia
7	(Paolacci et al., 2007), poplar (Street et al., 2011), and European beech (Fagus sylvatica)
8	(Olbrich et al., 2010; Olbrich et al., 2009; Olbrich et al., 2005). In some cases,
9	researchers compared transcriptomes of two or more cultivars, ecotypes or mutants that
10	differed in their sensitivity to O ₃ (Puckette et al., 2008; Rizzo et al., 2007; Lee and Yun,
11	<u>2006; Li et al., 2006b; Tamaoki et al., 2003</u>). Species, O_3 exposure conditions
12	(concentration, duration of exposure) and sampling times varied considerably in these
13	studies. However, functional classification of the genes that were either upregulated or
14	downregulated by plant exposure to O_3 exhibited common trends. Genes involved in
15	plant defense, signaling and those associated with the synthesis of plant hormones and
16	secondary metabolism were generally upregulated, while those related to photosynthesis
17	and general metabolism were typically downregulated in O ₃ -treated plants (Puckette et
18	al., 2008; Lee and Yun, 2006; Li et al., 2006b; Tosti et al., 2006; Olbrich et al., 2005;
19	<u>Tamaoki et al., 2003</u>).
20	
20	Analysis of the transcriptome has been used to evaluate differences in gene expression
21	between sensitive and tolerant plants in response to O_3 exposure. In pepper, 67% of the
22	180 genes studied that were affected by O_3 were differentially regulated in the sensitive
23	and tolerant cultivars. At both 0 hours and 48 hours after a 3-day exposure at 150 ppb, O_3
24	responsive genes were either upregulated or downregulated more markedly in the
25	sensitive than in the tolerant cultivar (Lee and Yun, 2006). Transcriptome analysis also
26	revealed differences in timing and magnitude of changes in gene expression between
27	sensitive and tolerant clovers. Acute exposure (300 ppb O ₃ for 6 hours) led to the
28	production of an oxidative burst in both clovers (Puckette et al., 2008). However, the
29	sensitive-Jemalong cultivar exhibited a sustained ROS burst and a concomitant
30	downregulation of defense response genes at 12 hours after the onset of exposure, while
31	the tolerant JE 154 accession showed much more rapid and large-scale transcriptome

33Arabidopsis ecotypes WS and Col-0 were exposed to $1.2 \times \text{ambient O}_3$ concentrations for348-12 days at the SoyFACE site (Li et al., 2006b). The sensitive WS ecotype showed a far35greater number of changes in gene expression in response to this low-level O₃ exposure36than the tolerant Col-0 ecotype. In a different study, exposure of the WS ecotype to37300 ppb O₃ for 6 hours showed a rapid induction of genes leading to cell death, such as

changes than the Jemalong cultivar (Puckette et al., 2008).

32

1 2	proteases, and downregulation or inactivation of cell signaling genes, demonstrating an ineffective defense response in this O_3 sensitive ecotype (<u>Mahalingam et al., 2006</u>).
3	The temporal response of plants to O_3 exposure was evaluated in the Arabidopsis Col-0
4	ecotype during a 6-hour exposure at 350 ppb O_3 and for 6 hours after the exposure was
5	completed. Results of this study, shown in Figure 9-5, indicate that genes associated with
6	signal transduction and regulation of transcription were in the class of early upregulated
7	genes, while genes associated with redox homeostasis and defense/stress response were
8	in the class of late upregulated genes (Mahalingam et al., 2005).
9	A few studies have been conducted to evaluate transcriptome changes in response to
10	longer term chronic O3 exposures in woody plant species. Longer term exposures resulted
11	in the upregulation of genes associated with secondary metabolites, including
12	isoprenoids, polyamines and phenylpropanoids in 2-year-old seedlings of the
13	Mediterranean shrub <i>Phillyrea latifolia</i> exposed to 110 ppb O ₃ for 90 days (Paolacci et
14	<u>al., 2007</u>). In 3-year-old European beech saplings exposed to O_3 for 20 months (with
15	monthly average twice ambient O ₃ concentrations ranging from 11 to 80 ppb),
16	O ₃ -induced changes in gene transcription were similar to those observed for herbaceous
17	species (Olbrich et al., 2009). Genes encoding proteins associated with plant stress
18	response, including ethylene biosynthesis, pathogenesis-related proteins and enzymes
19	detoxifying ROS, wereupregulated. Some genes associated with primary metabolism, cell
20	structure, cell division and cell growth were reduced (Olbrich et al., 2009). In a similar
21	study using adult European beech trees, it was determined that the magnitude of the
22	transcriptional changes described above was far greater in the saplings than in the adult
23	trees exposed to the same O_3 concentrations for the same time period (<u>Olbrich et al.</u> ,
24	<u>2010</u>).
25	The results from transcriptome studies described above have been substantiated by results
26	from proteome analysis in rice, poplar, European beech, wheat, and soybean. Exposure of
27	soybean to 120 ppb O_3 for 12 h/day for 3 days in growth chambers resulted in decreases
28	in the quantity of proteins associated with photosynthesis, while proteins involved with
29	antioxidant defense and carbon metabolism increased (Ahsan et al., 2010). Young poplar
30	plants exposed to 120 ppb O_3 in a growth chamber for 35 days also showed significant
31	changes in proteins involved in carbon metabolism (Bohler et al., 2007). Declines in
32	enzymes associated with carbon fixation, the Calvin cycle and photosystem II were
33	measured, while ascorbate peroxidase and enzymes associated with glucose catabolism
34	increased in abundance. In another study to determine the impacts of O_3 on both
35	developing and fully expanded poplar leaves, young poplars were exposed to 120 ppb O_3
36	for 13-h/day for up to 28 days (Bohler et al., 2010). Impacts on protein quantity only
37	occurred after the plants had been exposed to O_3 for 14 days, and at this point in time,

several Calvin cycle enzymes were reduced in quantity, while the effects on the light reactions appeared later, at 21 days after beginning treatment. Some of the antioxidant enzymes increased in abundance with O₃ treatment, while others (ascorbate peroxidase) did not. In relationship to leaf expansion, it was shown that O₃ did not affect protein quantity until leaves had reached full expansion, after about 7 days (Bohler et al., 2010).

6 Two-week-old rice seedlings exposed to varying levels of O_3 (4, 40, 80, 120 ppb) in a 7 growth chamber for 9 days showed reductions in quantities of proteins associated with 8 photosynthesis and energy metabolism, and increases in some antioxidant and defense 9 related proteins (Feng et al., 2008a). A subsequent study of O₃-treated rice seedlings 10 (exposed to 200 ppb O₃ for 24 hours) focusing on the integration of transcriptomics and 11 proteomics, supported and further enhanced these results (Cho et al., 2008). The authors 12 found that of the 22,000 genes analyzed from the rice genome, 1,535 were differentially 13 regulated by O_3 . Those differentially regulated genes were functionally categorized as 14 transcription factors, MAPK cascades, those encoding for enzymes involved in the 15 synthesis of jasmonic acid (JA), ethylene (ET), shikimate, tryptophan and lignin, and 16 those involved in glycolysis, the citric acid cycle, oxidative respiration and 17 photosynthesis. The authors determined that the proteome and metabolome (all small 18 molecule metabolites in a cell) analysis supported the results of the transcriptome 19 changes described above (Cho et al., 2008). This type of study, which ties together results 20 from changes in gene expression, protein quantity and activity, and metabolite levels, 21 provides the most complete picture of the molecular and biochemical changes occurring 22 in plants exposed to a stressor such as O_3 .

23 Sarkar et al. (2010) compared proteomes of two cultivars of wheat grown in OTCs at 24 several O_3 concentrations, including filtered air, ambient O_3 (mean concentration 47 ppb), 25 ambient + 10 ppb and ambient + 20 ppb for 5 h/day for 50 days. Declines in the rate of 26 photosynthesis and stomatal conductance were related to decreases in proteins involved 27 in carbon fixation and electron transport and increased proteolysis of photosynthetic 28 proteins such as the large subunit of ribulose-1,6-bisphosphate carboxylase/oxygenase 29 (Rubisco). Enzymes that take part in energy metabolism, such as ATP synthesis, were 30 also downregulated, while defense/stress related proteins were upregulated in O₃-treated 31 plants. In comparing the two wheat cultivars, Sarkar et al. (2010) found that while the 32 qualitative changes in protein expression between the two cultivars were similar, the 33 magnitude of these changes differed between the sensitive and tolerant wheat cultivars. 34 Greater foliar injury and a smaller decline in stomatal conductance was observed in the 35 sensitive cultivar as compared to the more tolerant cultivar, along with greater losses in 36 photosynthetic enzymes and higher quantities of antioxidant enzymes. Results from a 37 three-year exposure of European beech saplings to elevated O_3 (AOT40 value was 38 52.6 μ L/L•h for 2006 when trees were sampled) supported the results from the short-term

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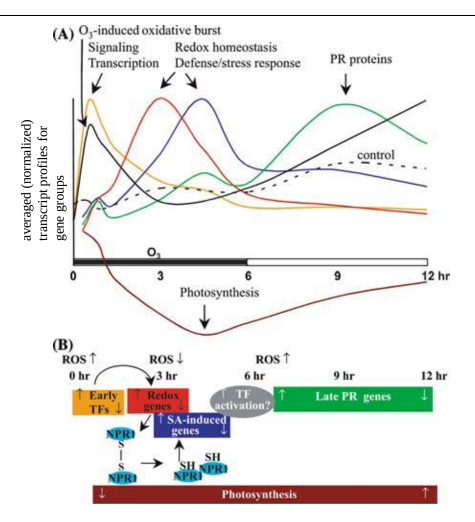
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1	exposure studies described above (Kerner et al., 2011). The O_3 treatment of the saplings
2	resulted in reductions in enzymes associated with the Calvin cycle, which could lead to
3	reduced carbon fixation. Enzymes associated with carbon metabolism/catabolism were
4	increased, and quantities of starch and sucrose were reduced in response to the O_3
5	treatment in these trees, indicating a potential impact of O ₃ on overall carbon metabolism
6	in long-term exposure conditions (Kerner et al., 2011).
7	Transcriptome and proteome studies have provided valuable information about O_3 effects
8	on plants. These studies allow for simultaneous analysis of changes in the expression
9	patterns of many different genes and proteins, and also provide information on how these
10	molecules might interact with one another as a result of plant exposure to oxidative
11	stress. Gene and protein expression patterns generally differ between O ₃ -sensitive and
12	tolerant plants, which could result from differential uptake or detoxification of O ₃ or from
13	differential regulation of the transcriptome and proteome.



Note: (A) Temporal profile of the oxidative stress response to ozone. The biphasic ozone-induced oxidative burst is represented in black, with the ROS control measurements shown as a broken line. Average transcript profiles are shown for early upregulated genes (yellow, peaks at 0.5-1 hours), and the 3 hours (blue), 4.5 hours (red) and 9-12 hours (green) late upregulated genes and for the downregulated genes coding for photosynthesis proteins (brown). (B) Diagrammatic representation of redox regulation of the oxidative stress response.

Source: Reprinted with permission of Springer (Mahalingam et al., 2005).

Figure 9-5 Composite diagram of major themes in the temporal evolution of the genetic response to ozone stress.

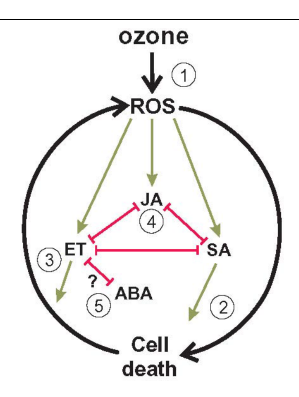
1	All of these studies describe common trends for changes in gene and protein expression
2	which occur in a variety of plant species exposed to O ₃ . While genes associated with
3	carbon assimilation and general metabolism are typically downregulated, genes
4	associated with signaling, catabolism, and defense are upregulated. The magnitude of
5	these changes in gene and protein expression appears to be related to plant species, age
6	and their sensitivity or tolerance to O_3 .

9.3.3.3 Role of Phytohormones in Plant Response to Ozone

 such as SA, ET and JA in determining plant response to 0₃. The 2006 0₃ AQCD documents the O₂-induced production of ET and its role in promoting the formation of leaf lesions. Transcriptome analysis and the use of a variety of mutants have allowed for further elucidation of the complex interactions between SA, ET, JA and the role of abscisic acid (ABA) in mediating plant response to 0₃ (Ludwikow and Sadowski, 2008). In addition to their roles in signaling pathways, phytohormones also appear to regulate, and be regulated by, the MAPK signaling cascades described previously. Most evidence suggests that while ET and SA are needed to develop O₂-induced leaf lesions, JA acts antagonistically to SA and ET to limit the lesions (Figure 9-6) (Kangasjarvi et al., 2005). The rapid production of ET in O₃ treated plants has been described in many plant species and has been further characterized through the use of a variety of mutants that either over-produce or are insensitive to ET. Production of stress ET in O₃-treated plants, which is thought to be part of a wounding response, was found to be correlated to the degree of injury development in leaves (U.S., EPA, 2006). More recent studies have supported these conclusions and have also focused on the interactions occurring between several oxidative-stress induced phytohormones, Yoshida et al. (2009) determined that ET likely amplifies the oxidative signal generated by ROS, thereby promoting lesion formation. By analyzing the O₂-induced transcriptome of several Arabidopsis mutants of the Col-0 exposure (200 ppb for 12 hours), the ET and JA signaling pathways were the main pathways used to activate plant defense responses, with a lesser role for SA. The authors also demonstrated that low level	1	Many studies of O ₃ effects on plants have analyzed the importance of plant hormones
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5further elucidation of the complex interactions between SA, ET, JA and the role of6abscisic acid (ABA) in mediating plant response to O3 (Ludwikow and Sadowski, 2008).7In addition to their roles in signaling cascades described previously. Most evidence8and be regulated by, the MAPK signaling cascades described previously. Most evidence9suggests that while ET and SA are needed to develop O ₂ -induced leaf lesions, JA acts10antagonistically to SA and ET to limit the lesions (Figure 9-6) (Kangasjarvi et al., 2005).11The rapid production of ET in O3 treated plants has been described in many plant species12and has been further characterized through the use of a variety of mutants that either13over-produce or are insensitive to ET. Production of stress ET in O ₂ -treated plants, which14is thought to be part of a wounding response, was found to be correlated to the degree of15injury development in leaves (U.S. EPA, 2006b). More recent studies have supported16these conclusions and have also focused on the interactions occurring between several17oxidative-stress induced phytohormones. Yoshida et al. (2009) determined that ET likely18amplifies the oxidative signal generated by ROS, thereby promoting lesion formation. By19analyzing the O ₂ -induced transcriptome of several Arabidopsis mutants of the Col-020ecotype, Tamaoki et al. (2003) determined that at 12 hours after initiating the O322pathways used to activate plant defense responses, with a lesser role for SA. The authors23also demonstrated that low levels of ET production could stimulate th	3	documents the O ₃ -induced production of ET and its role in promoting the formation of
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24defense genes, rather than promoting cell death which occurs when ET production is25high. Tosti et al. (2006) supported these findings by showing that plant exposure to O326not only results in activation of the biosynthetic pathways of ET, JA and SA, but also27increases the expression of genes related to the signal transduction pathways of these28phytohormones in O3-treated Arabidopsis plants (300 ppb O3 for 6 hours). Conversely, in29the O3 sensitive Ws ecotype, its sensitivity may, in part, be due to intrinsically high ET30levels leading to SA accumulation, and the high ET and SA may act to repress31JA-associated genes, which would serve to inhibit the spread of lesions (Mahalingam et32al., 2006). Ogawa et al. (2005) found that increases in SA in O3-treated plants leads to the33formation of leaf lesions in tobacco plants exposed to 200 ppb O3 for 6 hours.34Furthermore, in transgenic tobacco plants with reduced levels of ET production in35response to O3 exposure, several genes encoding for enzymes in the biosynthetic pathway	22	pathways used to activate plant defense responses, with a lesser role for SA. The authors
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34Furthermore, in transgenic tobacco plants with reduced levels of ET production in35response to O3 exposure, several genes encoding for enzymes in the biosynthetic pathway	32	al., 2006). Ogawa et al. (2005) found that increases in SA in O_3 -treated plants leads to the
35 response to O_3 exposure, several genes encoding for enzymes in the biosynthetic pathway	33	formation of leaf lesions in tobacco plants exposed to 200 ppb O_3 for 6 hours.
	34	Furthermore, in transgenic tobacco plants with reduced levels of ET production in
of SA were suppressed, suggesting that SA levels are in part, controlled by ET in the		
	36	of SA were suppressed, suggesting that SA levels are, in part, controlled by ET in the
37 presence of O_3 .	37	presence of O ₃ .

1	Exposure of the Arabidopsis mutant rcd1 to acute doses of O_3 (250 ppb O_3 for 8-h/day for
2	3 days) resulted in programmed cell death (PCD) and the formation of leaf lesions
3	(<u>Overmyer et al., 2000</u>). They determined that the observed induction of ET synthesis
4	promotes cell death, and that ET perception and signaling are required for the
5	accumulation of superoxide, which leads to cell death and propagation of lesions.
6	Jasmonic acid, conversely, contains the spread of leaf lesions (<u>Overmyer et al., 2000</u>).
7	Transcriptome analysis of several Arabidopsis mutants, which are insensitive to SA, ET
8	and JA, exposed to 12-h of 200 ppb O_3 showed that approximately 78 of the upregulated
9	genes measured in this study were controlled by ET and JA signaling pathways, while SA
10	signaling pathways were suggested to antagonize ET and JA pathways (<u>Tamaoki et al.</u> ,
10	
	<u>2003</u>). In a subsequent transcriptome study on the Col-0 ecotype exposed to 150 ppb O_3
12	for 48-h, JA and ET synthesis were downregulated, while SA was upregulated in O_3 -
13	treated plants. In cotton plants exposed to a range of O_3 concentrations (0-120 ppb) and
14	methyl jasmonate (MeJA), <u>Grantz et al. (2010b</u>) determined that exogenous applications
15	of MeJA did not protect plants from chronic O ₃ exposure.
16	Abscisic acid has been investigated for its role in regulating stomatal aperture and also
17	for its contribution to signaling pathways in the plant. The role of ABA and the
18	interaction between ABA and H ₂ O ₂ in O ₃ -induced stomatal closure was described in the
19	2006 O_3 AQCD. It was determined that the presence of H_2O_2 , which is formed from O_3
20	degradation, increases the sensitivity of guard cells to ABA and, therefore, more readily
21	results in stomatal closure. More recently, it was determined that synthesis of ABA was
22	induced in O_3 -treated Arabidopsis plants (250-350 ppb O_3 for 6 hours), with a more
23	pronounced induction in the O_3 sensitive rcd3 mutant as compared to the wildtype Col-0
24	(<u>Overmyer et al., 2008</u>). The rcd3 mutant also exhibited a lack of O_3 -induced stomatal
25	closure, and the RCD3 protein has been shown to be required for slow anion channels
26	(Overmyer et al., 2008). Ludwikow et al. (2009) used Arabidopsis ABI1td mutants, in
27	which a key negative regulator of ABA action (abscisic acid insensitive1 protein
28	phosphatase 2C) has been knocked out, to examine O_3 responsive genes in this mutant
29	compared to the Arabidopsis Col-0. Results of this study indicate a role for ABI1 in
30	negatively regulating the synthesis of both ABA and ET in O_3 -treated plants (350 ppb O_3
31	for 9 hours). Additionally, ABI1 may stimulate JA-related gene expression, providing
32	evidence for an antagonistic interaction between ABA and JA signaling pathways
33	(Ludwikow et al., 2009).
55	(<u>Ludwikow et al., 2009</u>).
34	Nitric oxide (NO) has also been shown to play a role in regulating gene expression in
35	plants in response to O_3 exposure. However, little is known to date about NO and its role
36	in the complex interactions of molecules in response to O_3 . Exposure of tobacco to O_3
37	(150 ppb for 5 hours) stimulated NO and NO-dependent ET production, while NO
38	production itself did not depend on the presence of ET (Ederli et al., 2006). Analysis of

1	O3-treated Arabidopsis indicated the possibility of a dual role for NO in the initiation of
2	cell death and later lesion containment (Ahlfors et al., 2009).
3	While much work remains to be done to better elucidate how plants detect O_3 , what
4	determines their sensitivity to the pollutant and how they might respond to it, it is clear
5	that the mechanism for O_3 detection and signal transduction is very complex. Many of the
6	phytohormones and other signaling molecules thought to be involved in these processes
7	are interactive and depend upon a variety of other factors, which could be either internal
8	or external to the plant. This results in a highly dynamic and complex system, capable of
9	resulting in a spectrum of plant sensitivity to oxidative stress and generating a variety of
10	plant responses to that stress.



Note: Ozone-derived radicals induce endogenous ROS production (1) which results in salicylic acid (SA) accumulation and programmed cell death; (2) Cell death triggers ethylene (ET) production, which is required for the continuing ROS production responsible for the propagation of cell death; (3) Jasmonates counteract the progression of the cycle by antagonizing the cell death promoting function of SA and ET; (4) Abscisic acid (ABA) antagonizes ET function in many situations and might also have this role in ozone-induced cell death; (5) Mutually antagonistic interactions between ET, SA and jasmonic acid (JA) are indicated with red bars.

Source: Reprinted with permission of Blackwell Publishing Ltd. (Kangasjarvi et al., 2005).

Figure 9-6 The oxidative cell death cycle.

9.3.4 **Detoxification**

9.3.4.1 Overview of Ozone-induced Defense Mechanisms

1 Plants are exposed to an oxidizing environment on a continual basis, and many reactions 2 that are part of the basic metabolic processes, such as photosynthesis and respiration, 3 generate ROS. As a result, there is an extensive and complex mechanism in place to 4 detoxify these oxidizing radicals, including both enzymes and metabolites, which are 5 located in several locations in the cell and also in the apoplast of the cell. As O₃ enters the 6 leaf through open stomata, the first point of contact of O_3 with the plant is likely in the 7 apoplast, where it breaks down to form oxidizing radicals such as H_2O_2 , O_2 , HO_2 and 8 HO₂. Another source of oxidizing radicals is an oxidative burst, generated by a 9 membrane-bound NADPH oxidase enzyme, which is recognized as an integral 10 component of the plant's defense system against pathogens (Schraudner et al., 1998). 11 Antioxidant metabolites and enzymes located in the apoplast are thought to form a first 12 line of defense by detoxifying O₃ and/or the ROS that are formed as breakdown products 13 of O_3 (Section 9.3.2). However, even with the presence of several antioxidants, including 14 ascorbate, the redox buffering capacity of the apoplast is far less than that of the 15 cytoplasm, as it lacks the regeneration systems necessary to retain a reduced pool of 16 antioxidants (Foyer and Noctor, 2005b). 17 Redox homeostasis is regulated by the presence of a pool of antioxidants, which are 18 typically found in a reduced state and detoxify ROS produced by oxidases or electron 19 transport components. As ROS increase due to environmental stress such as O₃, it is 20 unclear whether the antioxidant pool can maintain its reduced state (Foyer and Noctor, 21 2005b). As such, not only the quantity and types of antioxidant enzymes and metabolites 22 present, but also the cellular ability to regenerate those antioxidants are important 23 considerations in mechanisms of plant tolerance to oxidative stress (Dizengremel et al., 24 2008). Molecules such as glutathione (GSH), thioredoxins and NADPH play very 25 important roles in this regeneration process; additionally, it has been hypothesized that 26 alterations in carbon metabolism would be necessary to supply the needed reducing 27 power for antioxidant regeneration (Dizengremel et al., 2008).

9.3.4.2 **Role of Antioxidants in Plant Defense Responses**

28	Ascorbate has been the focus of many different studies as an antioxidant metabolite that
29	protects plants from exposure to O ₃ . It is found in several cellular locations, including the
30	chloroplast, the cytosol and the apoplast (Noctor and Foyer, 1998). Ascorbate is

1	synthesized in the cell and transported to the apoplast. Apoplastic ascorbate can be
2	oxidized to dehydroascorbate (DHA) with exposure to O_3 and is then transported back to
3	the cytoplasm. Here, DHA is reduced to ascorbate by the enzyme dehydroascorbate
4	reductase (DHAR) and reduced GSH, which is part of the ascorbate-glutathione cycle
5	(Noctor and Foyer, 1998). Many studies have focused on evaluating whether ascorbate is
6	the primary determining factor in differential sensitivity of plants to O ₃ . An evaluation of
7	several species of wildflowers in Great Smoky Mountains National Park showed a
8	correlation between higher quantities of reduced apoplastic ascorbate and lower levels of
9	foliar injury from O ₃ exposure in a field study on tall milkweed plants (Asclepsias
10	exaltata L.) (Burkey et al., 2006; Souza et al., 2006). Cheng et al. (2007) exposed two
11	soybean cultivars to elevated O_3 (77 ppb) and filtered air for 7-h/day for 6 days. The
12	differences in sensitivity between the two cultivars could not be explained by differential
13	O_3 uptake or by the fraction of reduced ascorbate present in the apoplast. However, total
14	antioxidant capacity of the apoplast was 2-fold higher in the tolerant Essex cultivar as
15	compared to the sensitive Forrest cultivar, indicating that there may be other compounds
16	in the leaf apoplast that scavenge ROS. D'Haese et al. (2005) exposed the NC-S
17	(sensitive) and NC-R (resistant) clones of white clover (<i>Trifolium repens</i>) to 60 ppb O_3
18	for 7-h/day for 5 days in environmental chambers. Surprisingly, the NC-S clone had a
19	higher constitutive concentration of apoplastic ascorbate with a higher redox status than
20	the NC-R clone. However, the redox status of symplastic GSH was higher in NC-R, even
21	though the concentration of GSH was not higher than in NC-S. In addition, total
22	symplastic antioxidative capacity was not a determining factor in differential sensitivity
23	between these two clones. Severino et al. (2007) also examined the role of antioxidants in
24	the differential sensitivity of the two white clover clones by growing them in the field for
25	a growing season and then exposing them to elevated O_3 (100 ppb for 8-h/day for
26	10 days) in OTC at the end of the field season. The NC-R clone had greater quantities of
27	total ascorbate and total antioxidants than the NC-S clone at the end of the experiment. In
28	snap bean, plants of the O ₃ tolerant Provider cultivar had greater total ascorbate and more
29	ascorbate in the apoplast than the sensitive S156 cultivar after exposure to 71 ppb O_3 for
30	10 days in OTC (Burkey et al., 2003). While most of the apoplastic ascorbate was in the
31	oxidized form, the ratio of reduced ascorbate to total ascorbate was higher in Provider
32	than S156, indicating that Provider is better able to maintain this ratio to maximize plant
33	protection from oxidative stress. Exposure of two wheat varieties to ambient (7-h average
34	44 ppb O_3) and elevated (7-h average 56 ppb O_3) O_3 for 60 days in open-air field
35	conditions showed higher concentrations of reduced ascorbate in the apoplast in the
36	tolerant Y16 variety than the more sensitive Y2 variety, however no varietal differences
37	were seen in the decrease in reduced ascorbate quantity in response to O_3 exposure (Feng
38	et al., 2010). There is much evidence that supports an important role for ascorbate,
39	particularly apoplastic ascorbate, in protecting plants from oxidative stressors such as O_3 ;

however, it is also clear that there is much variation in the importance of ascorbate for different plant species and differing exposure conditions. Additionally, the work of several authors suggests that there may be other compounds in the apoplast which have the capacity to act as antioxidants.

- 5 While the quantities of antioxidant metabolites such as ascorbate are an important 6 indicator of plant tolerance to O_3 , the ability of the plant to recycle oxidized ascorbate 7 efficiently also plays a large role in determining the plant's ability to effectively protect 8 itself from sustained exposure to oxidative stress. Tobacco plants over-expressing DHAR 9 were better protected from exposure to either chronic (100 ppb O_3 4-h/day for 30 days) or 10 acute (200 ppb O_3 for 2 hours) O_3 conditions than control plants and those with reduced 11 expression of DHAR (Chen and Gallie, 2005). The DHAR over-expressing plants 12 exhibited an increase in guard cell ascorbic acid, leading to a decrease in stomatal 13 responsiveness to O_3 and an increase in stomatal conductance and O_3 uptake. Despite 14 this, the presence of higher levels of ascorbic acid led to a lower oxidative load and a 15 higher level of photosynthetic activity in the DHAR over-expressing plants (Chen and 16 Gallie, 2005). A subsequent study with tobacco plants over-expressing DHAR confirmed 17 some of these results. Levels of ascorbic acid were higher in the transgenic tobacco 18 plants, and they exhibited greater tolerance to O_3 exposure (200 ppb O_3) as demonstrated 19 by higher photosynthetic rates in the transgenic plants as compared to the control plants 20 (Eltayeb et al., 2006). Over-expression of monodehydroascorbate reductase (MDAR) in 21 tobacco plants also showed enhanced stress tolerance in response to O_3 exposure 22 (200 ppb O₃), with higher rates of photosynthesis and higher levels of reduced ascorbic 23 acid as compared to controls (Eltayeb et al., 2007). Results of these studies demonstrate 24 the importance of ascorbic acid as a detoxification mechanism in some plant species, and 25 also emphasize that the recycling of oxidized ascorbate to maintain a reduced pool of 26 ascorbate is a factor in determining plant tolerance to oxidative stress.
- 27 The roles of other antioxidant metabolites and enzymes, including GSH, catalase (CAT), 28 peroxidase (POD) and superoxide dismutase (SOD), were comprehensively reviewed in 29 the 2006 O₃ AOCD. Based on the review of the literature, no conclusive and consistent 30 effects of O₃ on the quantity of GSH and CAT could be identified. Both apoplastic and 31 cvtosolic POD activity increased in response to O_3 exposure, while various isoforms of 32 SOD showed inconsistent changes in quantity in response to O_3 . Additional studies have 33 been conducted to further elucidate the roles of these antioxidant enzymes and 34 metabolites in protecting plants from oxidative stress. Superoxide dismutase and POD 35 activities were measured in both the tolerant Bel B and sensitive Bel W3 tobacco 36 cultivars exposed to ambient O_3 concentrations for 2 weeks 3 times throughout a growing 37 season (Borowiak et al., 2009). In this study, SOD and POD activity, including that of 38 several different isoforms, increased in both the sensitive and tolerant tobacco cultivars

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1	with exposure to O_3 , however the isoenzyme composition for POD differed between the
2	sensitive and tolerant tobacco cultivars (Borowiak et al., 2009) Tulip poplar
3	(Liriodendron tulipifera) trees exposed to increasing O ₃ concentrations (from 100 to
4	300 ppb O ₃ during a 2-week period) showed increases in activities of SOD, ascorbate
5	peroxidase (APX), glutathione reductase (GR), MDAR, DHAR, CAT and POD in the
6	2-week period, although individual enzyme activities increased at different times during
7	the 2-week period (<u>Ryang et al., 2009</u>).
8	Longer, chronic O ₃ exposures in trees revealed increases in SOD and APX activity in
9	Quercus mongolica after 45 days of plant exposure to 80 ppb O ₃ , which were followed by
10	declines in the activities and quantities of these enzymes after 75 days of exposure (Yan
11	et al., 2010). Similarly, activities of SOD, APX, DHAR, MDAR, and GR increased in
12	Gingko biloba trees during the first 50 days of exposure to 80 ppb O ₃ , followed by
13	decreases in activity below control values after 50 days of exposure (He et al., 2006).
14	Soybean plants exposed to 70 or 100 ppb O_3 for 4-h/day over the course of a growing
15	season showed elevated POD activity and a decrease in CAT activity at 40 and 60 days
16	after germination (Singh et al., 2010a).
17	Antioxidant enzymes and metabolites have been shown to play an important role in
18	determining plant tolerance to O ₃ and mediating plant responses to O ₃ . However, there is
19	also some evidence to suggest that the direct reaction of ascorbate with O ₃ could lead to
20	the formation of secondary toxicants, such as peroxy compounds, which may act upon
21	signal transduction pathways and modulate plant response to O_3 (Sandermann, 2008).
22	Therefore, the role of ascorbate and other antioxidants and their interaction with other
23	plant responses to O ₃ , such as the activation of signal transduction pathways, is likely far
24	more complex than is currently understood.

9.3.5 Effects on Primary and Secondary Metabolism

9.3.5.1 Light and Dark Reactions of Photosynthesis

25	Declines in the rate of photosynthesis and stomatal conductance in O ₃ -treated plants have
26	been documented for many different plant species (Booker et al., 2009; Wittig et al.,
27	<u>2007</u> ; U.S. EPA, 2006b). The 2006 O_3 AQCD described the mechanism by which plant
28	exposure to O ₃ reduces the quantity of Rubisco, and the more recent scientific literature
29	confirms these findings. While several measures of the light reactions of photosynthesis
30	are sensitive to exposure to O_3 (see below), photosynthetic carbon assimilation is
31	generally considered to be more affected by pollutant exposure, resulting in an overall
32	decline in photosynthesis (Guidi and Degl'Innocenti, 2008; Heath, 2008; Fiscus et al.,

1	2005). Loss of carbon assimilation capacity has been shown to result primarily from
2	declines in the quantity of Rubisco (Singh et al., 2009; Calatayud et al., 2007a).
3	Experimental evidence suggests that both decreases in Rubisco synthesis and enhanced
4	degradation of the protein contribute to the measured reduction in its quantity (U.S. EPA,
5	<u>2006b</u>). Reduced carbon assimilation has been linked to reductions in biomass and yield
6	(Wang et al., 2009b; He et al., 2007; Novak et al., 2007; Gregg et al., 2006; Keutgen et
7	<u>al., 2005</u>). Recent studies evaluating O_3 induced changes in the transcriptome and
8	proteome of several different species confirm these findings. Levels of mRNA for the
9	small subunit of Rubisco (rbcS) declined in European beech saplings exposed to 300 ppb
10	O_3 for 8-h/day for up to 26 days (Olbrich et al., 2005). Similar declines in rbcS mRNA
11	were also measured in the beech saplings in a free air exposure system over a course of
12	two growing seasons (Olbrich et al., 2009). Proteomics studies have also confirmed the
13	effects of O ₃ on proteins involved in carbon assimilation. Reductions in quantities of the
14	small and large subunit (rbcL) of Rubisco and Rubisco activase were measured in
15	soybean plants exposed to 120 ppb O_3 for 3 days in growth chambers (Ahsan et al.,
16	<u>2010</u>). Exposure of young poplar trees to 120 ppb O_3 for 35 days in exposure chambers
17	resulted in reductions of Rubisco, Rubisco activase, and up to 24 isoforms of Calvin
18	cycle enzymes, most of which play a role in regenerating the CO ₂ acceptor molecule,
19	ribulose-1.5-bisphosphate (Bohler et al., 2007). Reductions in protein quantity of both the
20	small and large subunit of Rubisco were seen in wheat plants exposed to ambient
21	(average concentration 47.3 ppb O_3) and elevated O_3 (ambient + 10 or 20 ppb O_3) in
22	open-top chambers for 5-h/day for 50 days (Sarkar et al., 2010). Lettuce plants exposed
23	to 100 ppb O_3 in growth chambers for 8-h/day for 3 weeks also showed reductions in
24	transcript and protein levels of the small and large subunits of Rubisco and Rubisco
25	activase (Goumenaki et al., 2010). The reductions in carbon assimilation have been
26	associated with declines in both the mRNA of the small and large subunits of Rubisco,
27	and with reductions in Rubisco activase mRNA and protein. Additionally, the reduction
28	in Rubisco quantity has also been associated with the O3-induced oxidative modification
29	of the enzyme, which is evidenced by the increases in carbonyl groups on the protein
30	after plant exposure to O_3 .
31	In addition to impacts on carbon assimilation, the deleterious effects of O_3 on the
32	photosynthetic light reactions have received more attention in recent years. Chlorophyll
33	fluorescence provides a useful measure of changes to the photosynthetic process from
34	exposure to oxidative stress. Decreases in the Fv/Fm ratio (a measure of the maximum
35	efficiency of Photosystem II) in dark adapted leaves indicate a decline in the efficiency of
36	the PSII photosystems and a concomitant increase in non-photochemical quenching
37	(Guidi and Degl'Innocenti, 2008; Scebba et al., 2006). Changes in these parameters have
38	been correlated to differential sensitivity of plants to the pollutant. In a study to evaluate

been correlated to differential sensitivity of plants to the pollutant. In a study to evaluate the response of 4 maple species to O_3 (exposed to an 8-h avg of 51 ppb for ambient and

39

- 1 79 ppb for elevated treatment in OTC), the 2 species which were most sensitive based on 2 visible injury and declines in CO₂ assimilation also showed the greatest decreases in 3 Fv/Fm in symptomatic leaves. In asymptomatic leaves, CO₂ assimilation decreased 4 significantly but there was no significant decline in Fv/Fm (Calatavud et al., 2007a). 5 Degl'Innocenti et al. (2007) measured significant decreases in Fv/Fm in young and 6 symptomatic leaves of a resistant tomato genotype (line 93.1033/1) in response to O_3 7 exposure (150 ppb O₃ for 3 hours in a growth chamber), but only minor decreases in 8 asymptomatic leaves with no associated changes in net photosynthetic rate. In the O₃ 9 sensitive tomato cultivar Cuor Di Bue, the Fv/Fm ratio did not change, while the 10 photosynthetic rate declined significantly in asymptomatic leaves (Degl'Innocenti et al., 2007). In two soybean cultivars, Fv/Fm also declined significantly with plant exposure to 11 12 O_3 (Singh et al., 2009). It appears that in asymptomatic leaves, photoinhibition, as 13 indicated by a decrease in Fv/Fm, is not the main reason for a decline in photosynthesis.
- 14 An evaluation of photosynthetic parameters of two white clover (Trifolium repens cv. 15 Regal) clones that differ in their O_3 sensitivity revealed that O_3 (40-110 ppb O_3 for 7-16 h/day for 5 days) increased the coefficient of non-photochemical quenching (q_{NP}) in both 17 the resistant (NC-R) and sensitive (NC-S) clones, however q_{NP} was significantly lower 18 for the sensitive clone (Crous et al., 2006). Sensitive Acer clones had a lower coefficient 19 of non-photochemical quenching, while exposure to O_3 increased q_{NP} in both sensitive 20 and tolerant clones (<u>Calatayud et al., 2007a</u>). While exposure to O_3 also increased q_{NP} in 21 tomato, there were no differences in the coefficient of photochemical quenching between 22 cultivars thought to be differentially sensitive to O_3 (Degl'Innocenti et al., 2007). Higher 23 q_{NP} as a result of exposure to O_3 indicates a reduction in the proportion of absorbed light 24 energy being used to drive photochemistry. A lower coefficient of non-photochemical 25 quenching in O₃ sensitive plants could indicate increased vulnerability to ROS generated 26 during exposure to oxidative stress (Crous et al., 2006).

27 Most of the research on O_3 effects on photosynthesis has focused on C3 (Calvin cycle) 28 plants because C4 (Hatch-Slack) plants have lower stomatal conductance and are, 29 therefore, thought to be less sensitive to O₃ stress. However, some studies have been 30 conducted to evaluate the effects of O_3 on C4 photosynthesis. In older maize leaves, 31 Leitao et al. (2007c); Leitao et al. (2007a) found that the activity, quantity and transcript 32 levels of both Rubisco and phosphoenolpyruvate carboxylase (PEPc) decreased as a 33 function of rising O_3 concentration. In younger maize leaves, the quantity, activity, and 34 transcript levels of the carboxylases were either increased or unaffected in plants exposed 35 to 40 ppb O₃ for 7- h/day for 28-33 days, but decreased at 80 ppb (Leitao et al., 2007b; 36 Leitao et al., 2007c). In another study, Grantz and Vu (2009) reported that O₃ exposures 37 (4, 58, and 114 ppb, 12-hour mean) decreased sugarcane biomass production by more than one third and allocation to roots by more than two thirds. 38

Respiration and Dark Respiration 9.3.5.2

1	While much research emphasis regarding O_3 effects on plants has focused on the negative
2	impacts on carbon assimilation, other studies have measured impacts on catabolic
3	pathways such as shoot respiration and photorespiration. Generally, shoot respiration has
4	been found to increase in plants exposed to O3. Bean plants exposed to ambient (average
5	12-h mean 43 ppb) and twice ambient (average 12-h mean 80 ppb) O_3 showed increases
6	in respiration. When mathematically partitioned, the maintenance coefficient of
7	respiration was significantly increased in O ₃ treated plants, while the growth coefficient
8	of respiration was not affected (Amthor, 1988). Loblolly pines were exposed to ambient
9	(12-h daily mean was 45 ppb) and twice ambient (12 hours daily mean was 86 ppb) O_3
10	for 12-h/day for approximately seven months per year for 3 and 4 years. While
11	photosynthetic activity declined with the age of the needles and increasing O ₃
12	concentration, enzymes associated with respiration showed higher levels of activity with
13	increasing O_3 concentration (<u>Dizengremel et al., 1994</u>). In their review on the role of
14	metabolic changes in plant redox status after O ₃ exposure, <u>Dizengremel et al. (2009</u>)
15	summarized multiple studies in which several different tree species were exposed to O ₃
16	concentrations ranging from ambient to 200 ppb O ₃ for at least several weeks. In all
17	cases, the activity of enzymes, including phosphofructokinase, pyruvate kinase and
18	fumarase, which are part of several catabolic pathways, were increased in O3 treated
19	plants.
19 20	•
20	Photorespiration is a light-stimulated process which consumes O_2 and releases CO_2 .
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9.3.5.3 Secondary Metabolism

1	Transcriptome analysis of Arabidopsis plants has revealed modulation of several genes
2	involved in plant secondary metabolism (Ludwikow and Sadowski, 2008). Phenylalanine
3	ammonia lyase (PAL) has been the focus of many studies involving plant exposure to O ₃
4	due to its importance in linking the phenylpropanoid pathway of plant secondary
5	metabolism to primary metabolism in the form of the shikimate pathway. Genes encoding
6	several enzymes of the phenylpropanoid pathway and lignin biosynthesis were
7	upregulated in transcriptome analysis of Arabidopsis plants (Col-0) exposed to 350 ppb
8	O ₃ for 6 hours, while 2 genes involved in flavonoid biosynthesis were downregulated
9	(Ludwikow et al., 2004). Exposure of Arabidopsis (Col-0) to lower O ₃ concentrations
10	(150 ppb for 8-h/day for 2 days) resulted in the induction of 11 transcripts involved in
11	flavonoid synthesis. In their exposure of 2-year-old Mediterranean shrub Phillyrea
12	latifolia to 110 ppb O ₃ for 90 days, Paolacci et al. (2007) identified four clones that were
13	upregulated and corresponded to genes involved in the synthesis of secondary
14	metabolites, such as isoprenoids, polyamines and phenylpropanoids. Upregulation of
15	genes involved in isoprene synthesis was also observed in Medicago trunculata exposed
16	to 300 ppb O_3 for 6 hours, while genes encoding enzymes of the flavonoid synthesis
17	pathway were either upregulated or downregulated (Puckette et al., 2008). Exposure of
18	red clover to $1.5 \times$ ambient O ₃ (average concentrations of 32.4 ppb) for up to 9 weeks in
19	an open field exposure system resulted in increases in leaf total phenolic content.
20	However, the types of phenolics that were increased in response to O ₃ exposure differed
21	depending upon the developmental stage of the plant. While almost all of the 31 different
22	phenolic compounds measured increased in quantity initially during the exposure, after
23	3 weeks the quantity of isoflavones decreased while other phenolics increased (Saviranta
24	et al., 2010). Exposure of beech saplings to ambient and $2 \times$ ambient O ₃ concentrations
25	over 2 growing seasons resulted in the induction of several enzymes which contribute to
26	lignin formation, while enzymes involved in flavonoid biosynthesis were downregulated
27	(<u>Olbrich et al., 2009</u>). Exposure of tobacco Bel W3 to 160 ppb O_3 for 5 hours showed
28	upregulation of almost all genes encoding for enzymes which are part of the
29	prechorismate pathway (Janzik et al., 2005). Isoprenoids can serve as antioxidant
30	compounds in plants exposed to oxidative stress (Paolacci et al., 2007).
31	The prechorismate pathway is the pathway leading to the formation of chorismate, a
32	precursor to the formation of the aromatic amino acids tryptophan, tyrosine and
33	phenylalanine. These amino acids are precursors for the formation of many secondary
34	aromatic compounds, and, therefore, the prechorismate pathway represents a branch-
35	point in the regulation of metabolites into either primary or secondary metabolism (Janzik
36	et al., 2005). Exposure of the O_3 sensitive Bel W3 tobacco cultivar at 160 ppb for 5 hours
37	showed an increase in transcript levels of most of the genes encoding enzymes of the

1	prechorismate pathway. However, shikimate kinase (SK) did not show any change in
2	transcript levels and only one of three isoforms of DAHPS (3-deoxy-D-arabino-
3	heptulosonat-7-phosphate synthase), the first enzyme in this pathway, was induced by O_3
4	exposure (Janzik et al., 2005). Differential induction of DAHPS isoforms was also
5	observed in European beech after 40 days of exposure to 150-190 ppb O ₃ . At this time
6	point in the beech experiment, transcript levels of shikimate pathway enzymes, including
7	SK, were generally strongly induced after an only weak initial induction after the first
8	40 days of exposure. Both soluble and cell-wall bound phenolic metabolites showed only
9	minimal increases in response to O_3 for the duration of the exposure period (Alonso et al.,
10	2007). Total leaf phenolics decreased with leaf age in <i>Populus nigra</i> exposed to 80 ppb
11	O ₃ for 12-h/day for 14 days. Ozone increased the concentration of total leaf phenolics in
12	newly expanded leaves, with the greatest increases occurring in compounds such as
13	quercitin glycoside, which has a high antioxidant capacity (Fares et al., 2010b). While
14	several phenylpropanoid pathway enzymes were induced in two poplar clones exposed to
15	60 ppb O_3 for 5-h/day for 15 days, the degree of induction differed between the two
16	clones. In the tolerant I-214 clone, PAL activity increased 9-fold in O ₃ -treated plants as
17	compared to controls, while there was no significant difference in PAL activity in the
18	sensitive Eridano clone (<u>DiBaccio et al., 2008</u>).
19	Polyamines such as putrescine, spermidine and spermine play a variety of roles in plants
20	and have been implicated in plant defense responses to both abiotic and biotic stresses.
21	They exist in both a free form and conjugated to hydroxycinnamic acids. Investigations
22	on the role of polyamines have found that levels of putrescine increase in response to
23	oxidative stress. This increase stems largely from the increase in the activity of arginine
24	decarboxylase (ADC), a key enzyme in the synthesis of putrescine (Groppa and
25	Benavides, 2008). Langebartels et al. (1991) described differences in putrescine
26	accumulation in O ₃ -treated tobacco plants exposed to several O ₃ concentrations, ranging
27	from 0-400 ppb for 5-7 hours. A large and rapid increase in putrescine occurred in the
28	tolerant Bel B cultivar and only a small increase in the sensitive Bel W3 cultivar, which
29	occurred only after the formation of necrotic leaf lesions. Van Buuren et al. (2002)
30	further examined the role of polyamines in these two tobacco cultivars during an acute
31	(130 ppb O_3 for 7-h in a growth chamber) exposure. They found that while free
32	putrescine accumulated in undamaged tissue of both cultivars, conjugated putrescine
33	predominantly accumulated in tissues undergoing cell death after plant exposure to O ₃
34	(Van Buuren et al., 2002). The authors suggest that while free putrescine may not play a
35	role in conferring tolerance in the Bel B cultivar, conjugated putrescine may play a role in
36	O ₃ -induced programmed cell death in Bel W3 plants.
37	Isoprene is emitted by some plant species and represents the predominant biogenic source
38	of hydrocarbon emissions in the atmosphere (Guenther et al., 2006). In the atmosphere,

1	the oxidation of isoprene by hydroxyl radicals can enhance O_3 formation in the presence
2	of NO_X , thereby impacting the O_3 concentration that plants are exposed to. While
3	isoprene emission varies widely between species, it has been proposed to stabilize
4	membranes and provide those plant species that produce it with a mechanism of
5	thermotolerance (Sharkey et al., 2008). It has also been suggested that isoprene may act
6	as an antioxidant compound to scavenge O ₃ (Loreto and Velikova, 2001). Recent studies
7	using a variety of plant species have shown conflicting results in trying to understand the
8	effects of O_3 on isoprene emission. Exposure to acute doses of O_3 (300 ppb for 3-h) in
9	detached leaves of Phragmites australis resulted in stimulation of isoprene emissions
10	(Velikova et al., 2005). Similar increases in isoprene emissions were measured in
11	<i>Populus nigra</i> after exposure to 100 ppb O_3 for 5 days continuously (Fares et al., 2008).
12	Isoprene emission in attached leaves of <i>Populus alba</i> , which were exposed to 150 ppb O_3
13	for 11-h/day for 30 days inside cuvettes, was inhibited, while isoprene emission and
14	transcript levels of isoprene synthase mRNA were increased in the leaves exposed to
15	ambient O_3 (40 ppb), which were located above the leaves enclosed in the exposure
16	cuvettes (Fares et al., 2006). Exposure of 2 genotypes of hybrid poplar to 120 ppb O ₃ for
17	6-h/day for 8 days resulted in a significant reduction in isoprene emission in the O_3 -
18	sensitive but not the tolerant genotype (<u>Ryan et al., 2009</u>). Similarly, O_3 treatment
19	(80 ppb 12-h/day for 14 days) of Populus nigra showed that isoprene emission was
20	reduced in the treated plants relative to the control plants (Fares et al., 2010b). Based on
21	results of this and other studies, Fares et al. (2010b) concluded that the isoprenoid
22	pathway may be induced in plants exposed to acute O ₃ doses, while at lower doses
23	isoprene emission may be inhibited. Vickers et al. (2009) developed transgenic tobacco
24	plants with the isoprene synthase gene from Populus alba and exposed them to 120 ppb
25	O3 for 6-h/day for 2 days. They determined that the wildtype plants showed significantly
26	more O ₃ damage, including the development of leaf lesions and a decline in
27	photosynthetic rates, than the transgenic, isoprene-emitting plants. Transgenic plants also
28	accumulated less H ₂ O ₂ and had lower levels of lipid peroxidation following exposure to
29	O_3 than the wildtype plants (Vickers et al., 2009). These results indicate that isoprene
30	may have a protective role for plants exposed to oxidative stress.

9.3.6 Summary

31	The results of recent studies on the effects of O ₃ stress on plants support and strengthen
32	those reported in the 2006 O ₃ AQCD. The most significant new body of evidence since
33	the 2006 O ₃ AQCD comes from research on molecular mechanisms of the biochemical
34	and physiological changes observed in many plant species in response to O_3 exposure.
35	Recent studies have employed new techniques, such as those used in evaluating

- 1 transcriptomes and proteomes to perform very comprehensive analyses of changes in 2 gene transcription and protein expression in plants exposed to O_3 . These newer molecular 3 studies not only provide very important information regarding the many mechanisms of 4 plant responses to O_3 , they also allow for the analysis of interactions between various 5 biochemical pathways which are induced in response to O_3 . However, many of these 6 studies have been conducted in artificial conditions with model plants, which are 7 typically exposed to very high, short doses of O_3 . Therefore, additional work remains to 8 elucidate whether these plant responses are transferable to other plant species exposed to 9 more realistic ambient conditions.
- 10 Ozone is taken up into leaves through open stomata. Once inside the substomatal cavity, 11 O_3 is thought to rapidly react with the aqueous layer surrounding the cell (apoplast) to 12 form breakdown products such as hydrogen peroxide (H_2O_2) , superoxide (O_2) , hydroxyl 13 radicals (HO) and peroxy radicals (HO₂). Plants could be detecting the presence of O_3 14 and/or its breakdown products in a variety different ways, depending upon the plant 15 species and the exposure parameters. Experimental evidence suggests that mitogen-16 activated protein kinases and calcium are important components of the signal 17 transduction pathways, which communicate signals to the nucleus and lead to changes in 18 gene expression in response to O_3 . It is probable that there are multiple detection 19 mechanisms and signal transduction pathways, and their activation may depend upon the 20 plant species, its developmental stage and/or O₃ exposure conditions. Initiation of signal 21 transduction pathways in O₃ treated plants has also been observed in stomatal guard cells. 22 Reductions in stomatal conductance have been described for many plant species exposed 23 to O_{3} , and new experimental evidence suggests that this reduction may be due not only to 24 a decrease in carboxylation efficiency, but also to a direct impact of O_3 on stomatal guard 25 cell function, leading to a changes in stomatal conductance.
- 26 Alterations in gene transcription that have been observed in O₃-treated plants are now 27 evaluated more comprehensively using DNA microarray studies, which measure changes 28 in the entire transcriptome rather than measuring the transcript levels of individual genes. 29 These studies have demonstrated very consistent trends, even though O_3 exposure 30 conditions (concentration, duration of exposure), plant species and sampling times vary 31 significantly. Genes involved in plant defense, signaling, and those associated with the 32 synthesis of plant hormones and secondary metabolism are generally upregulated in 33 plants exposed to O_3 , while those related to photosynthesis and general metabolism are 34 typically downregulated. Proteome studies support these results by demonstrating 35 concomitant increases or decreases in the proteins encoded by these genes. Transcriptome 36 analysis has also illuminated the complex interactions that exist between several different 37 phytohormones and how they modulate plant sensitivity and response to O_3 . 38 Experimental evidence suggests that while ethylene and salicylic acid are needed to

1develop O3-induced leaf lesions, jasmonic acid acts antagonistically to ethylene and2salicylic acid to limit the spread of the lesions. Abscisic acid, in addition to its role in3regulating stomatal aperture, may also act antagonistically to the jasmonic acid signaling4pathway. Changes in the quantity and activity of these phytohormones and the5interactions between them reveal some of the complexity of plant responses to an6oxidative stressor such as O3.

- 7 Another critical area of interest is to better understand and quantify the capacity of the 8 plant to detoxify oxygen radicals using antioxidant metabolites, such as ascorbate and 9 glutathione, and the enzymes that regenerate them. Ascorbate remains an important focus 10 of research, and, due to its location in the apoplast in addition to other cellular compartments, it is regarded as a first line of defense against oxygen radicals formed in 11 12 the apoplast. Most studies demonstrate that antioxidant metabolites and enzymes increase 13 in quantity and activity in plants exposed to O_3 , indicating that they play an important 14 role in protecting plants from oxidative stress. However, attempts to quantify the 15 detoxification capacity of plants have remained unsuccessful, as high quantities of 16 antioxidant metabolites and enzymes do not always translate into greater protection of the 17 plant. Considerable variation exists between plant species, different developmental 18 stages, and the environmental and O_3 exposure conditions which plants are exposed to.
- 19 As indicated earlier, the described alterations in transcript levels of genes correlate with 20 observed changes quantity and activity of the enzymes and metabolites involved in 21 primary and secondary metabolism. In addition to the generalized upregulation of the 22 antioxidant defense system, photosynthesis typically declines in O₃ treated plants. 23 Declines in C fixation due to reductions in quantity and activity of Rubisco were 24 extensively described in the 2006 O_3 AQCD. More recent studies support these results 25 and indicate that declines in Rubisco activity may also result from reductions in Rubisco 26 activase enzyme quantity. Other studies, which have focused on the light reactions of 27 photosynthesis, demonstrate that plant exposure to O_3 results in declines in electron 28 transport efficiency and a decreased capacity to quench oxidizing radicals. Therefore, the 29 overall declines in photosynthesis observed in O₃-treated plants likely result from 30 combined impacts on stomatal conductance, carbon fixation and the light reactions. 31 While photosynthesis generally declines in plants exposed to O_3 , catabolic pathways such 32 as respiration have been shown to increase. It has been hypothesized that increased 33 respiration may result from greater energy needs for defense and repair. Secondary 34 metabolism is generally upregulated in a variety of species exposed to O_3 as a part of a 35 generalized plant defense mechanism. Some secondary metabolites, such as flavonoids 36 and polyamines, are of particular interest as they are known to have antioxidant 37 properties. The combination of decreases in C assimilation and increases in catabolism

9.4 Nature of Effects on Vegetation and Ecosystems

9.4.1 Introduction

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2

3	Ambient O3 concentrations have long been known to cause visible symptoms, decreases
4	in photosynthetic rates, decreases in growth and yield of plants as well as many other
5	effects on ecosystems (U.S. EPA, 2006b, 1996c, 1986, 1978a). Numerous studies have
6	related O3 exposure to plant responses, with most effort focused on the yield of crops and
7	the growth of tree seedlings. Many experiments exposed individual plants grown in pots
8	or soil under controlled conditions to known concentrations of O ₃ for a segment of
9	daylight hours for some portion of the plant's life span. Information in this section also
10	goes beyond individual plant-scale responses to consider effects at the broader ecosystem
11	scale, including effects related to ecosystem services.

12This section will focus mainly on studies published since the release of the 2006 O313AQCD. However, because much O3 research was conducted prior to the 2006 O3 AQCD,14the present discussion of vegetation and ecosystem response to O3 exposure is largely15based on the conclusions of the 1978, 1986, 1996, and 2006 O3 AQCDs.

9.4.1.1 Ecosystem Scale, Function, and Structure

16	Information presented in this section was collected at multiple spatial scales or levels of
17	biological organization, ranging from the physiology of a given species to population,
18	community, and ecosystem investigations. An ecological population is a group of
19	individuals of the same species and a community is an assemblage of populations of
20	different species interacting with one another that inhabit an area. For this assessment,
21	"ecosystem" is defined as the interactive system formed from all living organisms and
22	their abiotic (physical and chemical) environment within a given area (IPCC, 2007a). The
23	boundaries of what could be called an ecosystem are somewhat arbitrary, depending on
24	the focus of interest or study. Thus, the extent of an ecosystem may range from very
25	small spatial scales or levels of biological organization to, ultimately, the entire Earth
26	(IPCC, 2007a). All ecosystems, regardless of size or complexity, have interactions and
27	physical exchanges between biota and abiotic factors, this includes both structural

(e.g., soil type and food web trophic levels) and functional (e.g., energy flow,
 decomposition, nitrification) attributes.

- 3 Ecosystems can be described, in part, by their structure, i.e., the number and type of 4 species present. Structure may refer to a variety of measurements including the species 5 richness, abundance, community composition and biodiversity as well as landscape 6 attributes. Competition among and within species and their tolerance to environmental 7 stressors are key elements of survivorship. When environmental conditions are shifted, 8 for example, by the presence of anthropogenic air pollution, these competitive 9 relationships may change and tolerance to stress may be exceeded. Ecosystems may also 10 be defined on a functional basis. "Function" refers to the suite of processes and 11 interactions among the ecosystem components and their environment that involve 12 nutrient and energy flow as well as other attributes including water dynamics and the flux 13 of trace gases. Plants, via such processes as photosynthesis, respiration, C allocation, 14 nutrient uptake and evaporation, affect energy flow, C, nutrient cycling and water 15 cycling. The energy accumulated and stored by vegetation (via photosynthetic C capture) 16 is available to other organisms. Energy moves from one organism to another through 17 food webs, until it is ultimately released as heat. Nutrients and water can be recycled. Air 18 pollution alters the function of ecosystems when elemental cycles or the energy flow are 19 altered. This alteration can also be manifested in changes in the biotic composition of 20 ecosystems.
- 21 There are at least three levels of ecosystem response to pollutants: (1) the individual 22 organism and its environment; (2) the population and its environment; and (3) the 23 biological community composed of many species and their environment (Billings, 1978). 24 Individual organisms within a population vary in their ability to withstand the stress of 25 environmental change. The response of individual organisms within a population is based 26 on their genetic constitution, stage of growth at time of exposure to stress, and the 27 microhabitat in which they are growing (Levine and Pinto, 1998). The stress range within 28 which organisms can exist and function determines the ability of the population to 29 survive.

9.4.1.2 Ecosystem Services

30Ecosystem structure and function may be translated into ecosystem services. Ecosystem31services are the benefits people obtain from ecosystems (UNEP, 2003). Ecosystems32provide many goods and services that are of vital importance for the functioning of the33biosphere and provide the basis for the delivery of tangible benefits to human society.

1	Hassan et al. (2005) define these benefits to include supporting, provisioning, regulating,
2	and cultural services:
3	 Supporting services are necessary for the production of all other ecosystem
4	services. Some examples include biomass production, production of
5	atmospheric O ₂ , soil formation and retention, nutrient cycling, water cycling,
6	and provisioning of habitat. Biodiversity is a supporting service that is
7	increasingly recognized to sustain many of the goods and services that humans
8	enjoy from ecosystems. These provide a basis for three higher-level categories
9	of services.
10	 Provisioning services, such as products (<u>Gitay et al., 2001</u>), i.e., food
11	(including game, roots, seeds, nuts and other fruit, spices, fodder), water, fiber
12	(including wood, textiles), and medicinal and cosmetic products (such as
13	aromatic plants, pigments).
14	 Regulating services that are of paramount importance for human society such
15	as (1) C sequestration, (2) climate and water regulation, (3) protection from
16	natural hazards such as floods, avalanches, or rock-fall, (4) water and air
17	purification, and (5) disease and pest regulation.
18	 Cultural services that satisfy human spiritual and aesthetic appreciation of
19	ecosystems and their components including recreational and other nonmaterial
20	benefits.
21	In the sections that follow, available information on individual, population and
22	community response to O_3 will be discussed. Effects of O_3 on productivity and
23	C sequestration, water cycling, below-ground processes, competition and biodiversity,
24	and insects and wildlife are considered below and in the context of ecosystem services
25	where appropriate.

9.4.2 Visible Foliar Injury and Biomonitoring

26 Visible foliar injury resulting from exposure to O₃ has been well characterized and 27 documented over several decades on many tree, shrub, herbaceous, and crop species 28 (U.S. EPA, 2006b, 1996b, 1984, 1978a). Visible foliar injury symptoms are considered 29 diagnostic as they have been verified experimentally in exposure-response studies, using 30 exposure methodologies such as CSTRs, OTCs, and free-air fumigation (see Section 9.2 31 for more detail on exposure methodologies). Several pictorial atlases and guides have 32 been published, providing details on diagnosis and identification of O₃-induced visible 33 foliar injury on many plant species throughout North America (Flagler, 1998; NAPAP, 34 1987) and Europe (Innes et al., 2001; Sánchez et al., 2001). Typical visible injury

1 2 3 4 5 6 7 8 9 10	symptoms on broad-leaved plants include: stippling, flecking, surface bleaching, bifacial necrosis, pigmentation (e.g., bronzing), chlorosis, and/or premature senescence. Typical visible injury symptoms for conifers include: chlorotic banding, tip burn, flecking, chlorotic mottling, and/or premature senescence of needles. Although common patterns of injury develop within a species, these foliar lesions can vary considerably between and within taxonomic groups. Furthermore, the degree and extent of visible foliar injury development varies from year to year and site to site (Orendovici-Best et al., 2008; Chappelka et al., 2007; Smith et al., 2003), even among co-members of a population exposed to similar O ₃ levels, due to the influence of co-occurring environmental and genetic factors (Souza et al., 2006; Chappelka et al., 2003; Somers et al., 1998).
11	Nevertheless, <u>Chappelka et al. (2007</u>) reported that the average incidence of O_3 -induced
12	foliar injury was 73% on milkweed observed in the Great Smoky Mountains National
13	Park in the years 1992-1996.
14	Although the majority of O ₃ -induced visible foliar injury occurrence has been observed
15	on seedlings and small plants, many studies have reported visible injury of mature
16	coniferous trees, primarily in the western U.S. (Arbaugh et al., 1998) and to mature
17	deciduous trees in eastern North America (Schaub et al., 2005; Vollenweider et al., 2003;
18	Chappelka et al., 1999a; Chappelka et al., 1999b; Somers et al., 1998; Hildebrand et al.,
19	<u>1996</u>).
20	It is important to note that visible foliar injury occurs only when sensitive plants are
21	
21	exposed to elevated O ₃ concentrations in a predisposing environment. A major modifying
21 22	exposed to elevated O_3 concentrations in a predisposing environment. A major modifying factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a
22	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a
22 23	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack
22 23 24 25 26	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits
22 23 24 25 26 27	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to
22 23 24 25 26 27 28	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (<u>Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1992; Temple et al., 2004; Context and Context an</u>
22 23 24 25 26 27 28 29	factor for O ₃ -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O ₃ entering the leaf that can cause injury (<u>Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988</u>). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O ₃ -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O ₃ ,
22 23 24 25 26 27 28 29 30	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O_3 -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O_3 , especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age
22 23 24 25 26 27 28 29 30 31	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O_3 -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O_3 , especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age influence the severity of symptom expression with older leaves showing greater injury
22 23 24 25 26 27 28 29 30	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O_3 -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O_3 , especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age
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22 23 24 25 26 27 28 29 30 31 32	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O_3 -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O_3 , especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age influence the severity of symptom expression with older leaves showing greater injury severity as a result of greater seasonal exposure (Zhang et al., 2010a).
22 23 24 25 26 27 28 29 30 31 32 33	 factor for O₃-induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O₃ entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O₃-induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O₃, especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age influence the severity of symptom expression with older leaves showing greater injury severity as a result of greater seasonal exposure (Zhang et al., 2010a). Although visible injury is a valuable indicator of the presence of phytotoxic
22 23 24 25 26 27 28 29 30 31 32 33 34	 factor for O₃-induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O₃ entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O₃-induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O₃, especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age influence the severity of greater seasonal exposure (Zhang et al., 2010a). Although visible injury is a valuable indicator of the presence of phytotoxic concentrations of O₃ in ambient air, it is not always a reliable indicator of other negative
22 23 24 25 26 27 28 29 30 31 32 33 34 35	factor for O_3 -induced visible foliar injury is the amount of soil moisture available to a plant during the year that the visible foliar injury is being assessed. This is because lack of soil moisture generally decreases stomatal conductance of plants and, therefore, limits the amount of O_3 entering the leaf that can cause injury (Matyssek et al., 2006; Panek, 2004; Grulke et al., 2003a; Panek and Goldstein, 2001; Temple et al., 1992; Temple et al., 1988). Consequently, many studies have shown that dry periods in local areas tend to decrease the incidence and severity of O_3 -induced visible foliar injury; therefore, the incidence of visible foliar injury is not always higher in years and areas with higher O_3 , especially with co-occurring drought (Smith et al., 2003). Other factors such as leaf age influence the severity of symptom expression with older leaves showing greater injury severity as a result of greater seasonal exposure (Zhang et al., 2010a). Although visible injury is a valuable indicator of the presence of phytotoxic concentrations of O_3 in ambient air, it is not always a reliable indicator of other negative effects on vegetation. The significance of O_3 injury at the leaf and whole plant levels

1	existing leaf area. Previous O ₃ AQCDs have noted the difficulty in relating visible foliar
2	injury symptoms to other vegetation effects such as individual plant growth, stand
3	growth, or ecosystem characteristics (U.S. EPA, 2006b, 1996b). As a result, it is not
4	presently possible to determine, with consistency across species and environments, what
5	degree of injury at the leaf level has significance to the vigor of the whole plant.
6	However, in some cases, visible foliar symptoms have been correlated with decreased
7	vegetative growth (Somers et al., 1998; Karnosky et al., 1996; Peterson et al., 1987;
8	Benoit et al., 1982) and with impaired reproductive function (Chappelka, 2002; Black et
9	al., 2000). Conversely, the lack of visible injury does not always indicate a lack of
10	phytotoxic concentrations of O_3 or a lack of non-visible O_3 effects (Gregg et al., 2006,
11	<u>2003</u>).

9.4.2.1 Biomonitoring

12 The use of biological indicators to detect phytotoxic levels of O_3 is a longstanding and 13 effective methodology (Chappelka and Samuelson, 1998; Manning and Krupa, 1992). A 14 plant bioindicator can be defined as a vascular or nonvascular plant exhibiting a typical 15 and verifiable response when exposed to a plant stress such as an air pollutant (Manning, 16 2003). To be considered a good indicator species, plants must (1) exhibit a distinct, 17 verified response; (2) have few or no confounding disease or pest problems; and (3) 18 exhibit genetic stability (U.S. EPA, 2006b). Such sensitive plants can be used to detect 19 the presence of a specific air pollutant such as O_3 in the ambient air at a specific location 20 or region and, as a result of the magnitude of their response, provide unique information 21 regarding specific ambient air quality. Bioindicators can be either introduced sentinels, 22 such as the widely used tobacco (Nicotiana tabacum) variety Bel W3 (Calatayud et al., 23 2007b; Laffray et al., 2007; Nali et al., 2007; Gombert et al., 2006; Kostka-Rick and 24 Hahn, 2005; Heggestad, 1991) or detectors, which are sensitive native plant species 25 (Chappelka et al., 2007; Souza et al., 2006). The approach is especially useful in areas 26 where O_3 monitors are not operated (Manning, 2003). For example, in remote wilderness 27 areas where instrument monitoring is generally not available, the use of bioindicator 28 surveys in conjunction with the use of passive samplers (Krupa et al., 2001) may be a 29 useful methodology (Manning, 2003). However, it requires expertise in recognizing those 30 signs and symptoms uniquely attributable to exposure to O_3 as well as in their 31 quantitative assessment.

32Since the 2006 O3 AQCD, new sensitive plant species have been identified from field33surveys and verified in controlled exposure studies (Kline et al., 2009; Kline et al., 2008).34Several multiple-year field surveys have also been conducted at National Wildlife

Refuges in Maine, Michigan, New Jersey, and South Carolina (<u>Davis, 2009</u>, <u>2007a</u>, <u>b</u>; <u>Davis and Orendovici, 2006</u>).

- 3 The USDA Forest Service through the Forest Health Monitoring Program (FHM) (1990 -4 2001) and currently the Forest Inventory and Analysis (FIA) Program has been collecting 5 data regarding the incidence and severity of visible foliar injury on a variety of O₃ 6 sensitive plant species throughout the U.S. (Coulston et al., 2003; Smith et al., 2003). The 7 plots where these data are taken are known as biosites. These biosites are located 8 throughout the country and analysis of visible foliar injury within these sites follows a set 9 of established protocols. For more details, see http://www.nrs.fs.fed.us/fia/topics/ozone/ 10 (USDA, 2011). The network has provided evidence of O_3 concentrations high enough to 11 induce visible symptoms on sensitive vegetation. From repeated observations and 12 measurements made over a number of years, specific patterns of areas experiencing 13 visible O₃ injury symptoms can be identified. (Coulston et al., 2003) used information 14 gathered over a 6-year period (1994-1999) from the network to identify several species 15 that were sensitive to O_3 over entire regions, including sweetgum (*Liquidambar*) 16 styraciflua), loblolly pine (Pinus taeda), and black cherry (P. serotina). In a study of the 17 west coast of the U.S, Campbell et al. (2007) reported O₃ injury in 25-37% of biosites in 18 California forested ecosystems from 2000-2005.
- 19 A study by Kohut (2007) assessed the estimated risk of O_3 -induced visible foliar injury 20 on bioindicator plants (NPS, 2006) in 244 national parks in support of the National Park 21 Service's Vital Signs Monitoring Network (NPS, 2007). The risk assessment was based 22 on a simple model relating response to the interaction of species, level of O_3 exposure, 23 and exposure environment. Kohut (2007) concluded that the estimated risk of visible 24 foliar injury was high in 65 parks (27%), moderate in 46 parks (19%), and low in 131 25 parks (54%). Some of the well-known parks with a high risk of O₃-induced visible foliar 26 injury include Gettysburg, Valley Forge, Delaware Water Gap, Cape Cod, Fire Island, 27 Antietam, Harpers Ferry, Manassas, Wolf Trap Farm Park, Mammoth Cave, Shiloh, 28 Sleeping Bear Dunes, Great Smoky Mountains, Joshua Tree, Sequoia and Kings Canyon, 29 and Yosemite.
- 30 Lichens have also long been used as biomonitors of air pollution effects on forest health 31 (Nash, 2008). It has been suspected, based on field surveys in the San Bernardino 32 Mountains surrounding the Los Angeles air basin, that declines in lichen diversity and 33 abundance were correlated with measured O₃ gradients (<u>Gül et al., 2011</u>). Several recent 34 studies in North America (Geiser and Neitlich, 2007; Gombert et al., 2006; Jovan and 35 McCune, 2006) and Europe (Nali et al., 2007; Gombert et al., 2006) have used lichens as 36 biomonitors of atmospheric deposition (e.g., N and S) and O₃ exposure. Nali et al. (2007) 37 found that epiphytic lichen biodiversity was not related to O₃ geographical distribution.

1

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1In addition, a recent study by Riddell et al. (2010) found that lichen species, Ramalina2menziesii, showed no decline in physiological response to low and moderate3concentrations of O3 and may not be a good indicator for O3 pollution. Mosses have also4been used as biomonitors of air pollution; however, there remains a knowledge gap in the5understanding of the effects of ozone on mosses as there has been very little information6available on this topic in recent years.

9.4.2.2 Summary

7	Visible foliar injury resulting from exposure to O ₃ has been well characterized and
8	documented over several decades of research on many tree, shrub, herbaceous, and crop
9	species (U.S. EPA, 2006b, 1996b, 1984, 1978a). Ozone-induced visible foliar injury
10	symptoms on certain bioindicator plant species are considered diagnostic as they have
11	been verified experimentally in exposure-response studies, using exposure methodologies
12	such as continuous stirred tank reactors (CSTRs), OTCs, and free-air fumigation.
13	Experimental evidence has clearly established a consistent association of visible injury
14	with O_3 exposure, with greater exposure often resulting in greater and more prevalent
15	injury. Since the 2006 O ₃ AQCD, results of several multi-year field surveys of
16	O ₃ -induced visible foliar injury at National Wildlife Refuges in Maine, Michigan, New
17	Jersey, and South Carolina have been published. New sensitive species showing visible
18	foliar injury continue to be identified from field surveys and verified in controlled
19	exposure studies.
20	The use of biological indicators in field surveys to detect phytotoxic levels of O_3 is a
21	longstanding and effective methodology. The USDA Forest Service through the Forest
22	Health Monitoring (FHM) Program (1990-2001) and currently the Forest Inventory and
23	Analysis (FIA) Program has been collecting data regarding the incidence and severity of
24	visible foliar injury on a variety of O ₃ sensitive plant species throughout the U.S. The
25	network has provided evidence that O ₃ concentrations were high enough to induce visible
26	symptoms on sensitive vegetation. From repeated observations and measurements made
27	over a number of years, specific patterns of areas experiencing visible O ₃ injury
28	symptoms can be identified. As noted in the preceding section, a study of 244 national
29	parks indicated that the estimated risk of visible foliar injury was high in 65 parks (27%),
30	moderate in 46 parks (19%), and low in 131 parks (54%).
31	Evidence is sufficient to conclude that there is a causal relationship between ambient
32	O_3 exposure and the occurrence of O_3 -induced visible foliar injury on sensitive
33	vegetation across the U.S.

9.4.3 Growth, Productivity and Carbon Storage in Natural Ecosystems

Ambient O3 concentrations have long been known to cause decreases in photosynthetic
rates, decreases in growth, and decreases in yield (U.S. EPA, 2006b, 1996c, 1986,
<u>1978a</u>). The O_3 -induced damages at the plant scale may translate to damages at the stand,
then ecosystem scales, and cause changes in productivity and C storage. This section
focuses on the responses of C cycling to seasonal or multi-year exposures to O_3 at levels
of organization ranging from individual plants to ecosystems. Quantitative responses
include changes in plant growth, plant biomass allocation, ecosystem production and
ecosystem C sequestration. Most information available on plant-scale responses was
obtained from studies that used a single species especially tree seedlings and crops, while
some used mixtures of herbaceous species. Ecosystem changes are difficult to evaluate in
natural settings, due to the complexity of interactions, the number of potential
confounders, and the large spatial and temporal scales. The discussion of ecosystem
effects focuses on new studies at the large-scale FACE experiments and on ecological
model simulations.

9.4.3.1 Plant Growth and Biomass Allocation

- 15The previous O3 AQCDs concluded that there is strong evidence that exposure to O316decreases photosynthesis and growth in numerous plant species (U.S. EPA, 2006b,171996b, 1984, 1978a). Studies published since the last review support those conclusions18and are summarized below.
- 19 In general, research conducted over several decades has indicated that exposure to O_3 20 alters stomatal conductance and reduces photosynthesis in a wide variety of plant species. 21 In a review of more than 55 studies, Wittig et al. (2007) reported that current O_3 22 concentrations in the northern hemisphere are decreasing stomatal conductance (13%) 23 and photosynthesis (11%) across tree species. It was also found that younger trees (<4 24 years) were affected less by O₃ than older trees. Further, the authors also found that 25 decreases in photosynthesis are consistent with the cumulative uptake of O_3 into the leaf. 26 In contrast, several studies reported that O₃ exposure may result in loss of stomatal 27 control, incomplete stomatal closure at night and a decoupling of photosynthesis and stomatal conductance, which may have implications for whole- plant water use 28 29 (Section 9.4.5).
- 30In a recently published meta-analysis, Wittig et al. (2009) quantitatively compiled peer31reviewed studies from the past 40 years on the effect of current and future O3 exposures32on the physiology and growth of forest species. They found that current ambient O3

1	concentrations as reported in those studies significantly decreased annual total biomass
2	growth (7%) across 263 studies. The authors calculated the ambient O_3 concentrations
3	across these studies to average 40 ppb. This average was calculated across the duration of
4	each study and there were therefore many hourly exposures well above 40 ppb. The
5	decreased growth effect was reported to be greater (11 to 17%) in elevated O_3 exposures
6	(97 ppb) (<u>Wittig et al., 2009</u>). This meta-analysis demonstrates the coherence of O_3
7	effects across numerous studies and species that used a variety of experimental
8	techniques, and these results support the conclusion of the previous AQCD that exposure
9	to O_3 decreases plant growth.
10	In two companion papers, McLaughlin et al. (2007a); (2007b) investigated the effects of
11	ambient O_3 on tree growth and hydrology at forest sites in the southern Appalachian
12	Mountains. The authors reported that the cumulative effects of ambient levels of O_3
13	decreased seasonal stem growth by 30-50% for most tree species in a high O_3 year in
14	comparison to a low O_3 year (McLaughlin et al., 2007a). The authors also reported that
15	high ambient O3 concentrations can disrupt whole-tree water use and in turn reduce late-
16	season streamflow (McLaughlin et al., 2007b); see Section 9.4.5 for more on water
17	cycling.
18	Since the 2006 O ₃ AQCD, several recent studies have reported results from the Aspen
19	FACE "free air" O ₃ and CO ₂ exposure experiment in Wisconsin (Darbah et al., 2008;
20	Riikonen et al., 2008; Darbah et al., 2007; Kubiske et al., 2007; Kubiske et al., 2006;
21	King et al., 2005). At the Aspen FACE site, single-species and two-species stands of trees
22	were grown in 12, 30-m diameter rings corresponding to three replications of a full
23	factorial arrangement of two levels each of CO ₂ and O ₃ exposure. Over the first
24	seven years of stand development, <u>Kubiske et al. (2006</u>) observed that elevated O_3
25	decreased tree heights, diameters, and main stem volumes in the aspen community by 11,
26	16, and 20%, respectively. In addition, <u>Kubiske et al. (2007</u>) reported that elevated O_3
27	may change intra- and inter-species competition. For example, O3 treatments increased
28	the rate of conversion from a mixed aspen-birch community to a birch dominated
29	community. In a comparison presented in Section <u>9.6.3</u> of this document, EPA found that
30	effects on biomass accumulation in aspen during the first seven years closely agreed with
31	the exposure-response function based on data from earlier OTC experiments.
32	Several studies at the Aspen FACE site also considered other growth-related effects of
33	elevated O_3 . Darbah et al. (2008); Darbah et al. (2007) reported that O_3 treatments
34	decreased paper birch seed weight and seed germination and that this would likely lead to
35	a negative impact of regeneration for that species. Riikonen et al. (2008) found that
36	elevated O_3 decreased the amount of starch in birch buds by 16%, and reduced aspen bud
37	size, which may have been related to the observed delay in spring leaf development. The

1results suggest that elevated O3 concentrations have the potential to alter C metabolism of2overwintering buds, which may have carry-over effects in the subsequent growing season3(Riikonen et al., 2008).

- 4 Effects on growth of understory vegetation were also investigated at Aspen FACE. 5 Bandeff et al. (2006) found that the effects of elevated CO_2 and O_3 on understory species 6 composition, total and individual species biomass, N content, and ¹⁵N recovery were a 7 result of overstory community responses to those treatments; however, the lack of 8 apparent direct O_3 treatment effects may have been due to high variability in the data. 9 Total understory biomass increased with increasing light and was greatest under the open 10 canopy of the aspen/maple community, as well as the more open canopy of the elevated 11 O₃ treatments (Bandeff et al., 2006). Similarly, data from a study by Awmack et al. 12 (2007) suggest that elevated CO₂ and O₃ may have indirect growth effects on red 13 (Trifolium pratense) and white (Trifolium repens) clover in the understory via overstory 14 community effects; however, no direct effects of elevated O₃ were observed.
- 15Overall, the studies at the Aspen FACE experiment are consistent with many of the OTC16studies that were evaluated in previous O3 AQCDs demonstrating that O3 exposure17decreases growth in numerous plant species. These results strengthen the understanding18of O3 effects on forests and demonstrate the relevance of the knowledge gained from19trees grown in open-top chamber studies.
- 20 For some annual species, particularly crops, the relevant measurement for an assessment 21 of the risk of O_3 exposure is yield or growth, e.g., production of grain or biomass. For 22 plants grown in mixtures such as hayfields, and natural or semi-natural grasslands 23 (including native nonagricultural species), affected factors other than production of 24 biomass may be important. Such endpoints include biodiversity or species composition, 25 and effects on those endpoints may be indirect, resulting, for example, from competitive 26 interactions among plants in mixed-species communities. Most of the available data on 27 non-crop herbaceous species are for grasslands, with many of the recent studies 28 conducted in Europe. See Section 9.4.7 for a review of the recent literature on O₃ effects 29 on competition and biodiversity in grasslands.

Root growth

30Although O3 does not penetrate soil, it could alter root development by decreasing31C assimilation via photosynthesis leading to less C allocation to the roots (Andersen,322003). The response of root development to O3 exposure depends on available33photosynthate within the plant and could vary over time. Many biotic and abiotic factors,34such as community dynamics and drought stress, have been found to alter root

1	development under elevated O ₃ . Generally, there is clear evidence that O ₃ reduces C
2	allocation to roots; however, results of a few recent individual studies have shown
3	negative (Jones et al., 2010), non-significant (Andersen et al., 2010; Phillips et al., 2009)
4	and positive effects (Pregitzer et al., 2008; Grebenc and Kraigher, 2007) on root biomass
5	and root: shoot ratio.
6	An earlier study at the Aspen FACE experiment found that elevated O ₃ reduced coarse
7	root and fine roots biomass in young stands of paper birch and trembling aspen (King et
8	al., 2001). However, this reduction disappeared several years later. Ozone significantly
9	increased fine-root (<1.0 mm) in the aspen community (Pregitzer et al., 2008). This
10	increase in fine root production was due to changes in community composition, such as
11	better survival of the O ₃ -tolerant aspen genotype, birch, and maple, rather than changes in
12	C allocation at the individual tree level (Pregitzer et al., 2008; Zak et al., 2007). In an
13	adult European beech/Norway spruce forest in Germany, drought was found to nullify the
14	O ₃ -driven stimulation of fine root growth. Ozone stimulated fine-root production of
15	beech during the humid year, but had no significant impact on fine root production in the
16	dry year (Matyssek et al., 2010; Nikolova et al., 2010).
17	Using a non-destructive method, Vollsnes et al. (2010) studied the in vivo root
18	development of subterranean clover (Trifolium subterraneum) before, during and after
19	short-term O ₃ exposure. It was found that O ₃ reduced root tip formation, root elongation,
20	the total root length, and the ratios between below- and above-ground growth within
21	one week after exposure. Those effects persisted for up to three weeks; however, biomass
22	and biomass ratios were not significantly altered at the harvest five weeks after exposure.
23	Several recent meta-analyses have generally indicated that O ₃ reduced C allocated to
24	roots. In one meta-analysis, <u>Grantz et al. (2006</u>) estimated the effect of O_3 on the
25	root:shoot allometric coefficient (k), the ratio between the relative growth rate of the root
26	and shoot. The results showed that O ₃ reduced the root:shoot allometric coefficient by
27	5.6%, and the largest decline of the root:shoot allometric coefficient was observed in
28	slow-growing plants. In another meta-analysis including 263 publications, Wittig et al.
29	(2009) found that current O_3 exposure had no significant impacts on root biomass and
30	root:shoot ratio when compared to pre-industrial O ₃ exposure. However, if O ₃
31	concentrations rose to 81-101 ppb (projected O_3 levels in 2100), both root biomass and
32	root:shoot ratio were found to significantly decrease. Gymnosperms and angiosperms
33	differed in their responses, with gymnosperms being less sensitive to elevated O ₃ . In two
34	other meta-analyses, Wang and Taub (2010) found elevated O_3 reduced biomass
35	allocation to roots by 8.3% at ambient CO_2 and 6.0% at elevated CO_2 , and Morgan et al.
36	(2003) found O_3 reduced root dry weight of soybean.

9.4.3.2 Summary

1	The previous O_3 AQCDs concluded that there is strong and consistent evidence that
2	ambient concentrations of O ₃ decrease photosynthesis and growth in numerous plant
3	species across the U.S. Studies published since the last review continue to support that
4	conclusion.
5	The meta-analyses by Wittig et al. (2009); Wittig et al. (2007) demonstrate the coherence
6	of O ₃ effects on plant photosynthesis and growth across numerous studies and species
7	using a variety of experimental techniques. Furthermore, recent meta-analyses have
8	generally indicated that O ₃ reduced C allocation to roots (Wittig et al., 2009; Grantz et
9	<u>al., 2006</u>). Since the 2006 O_3 AQCD, several studies were published based on the Aspen
10	FACE experiment using "free air," O ₃ , and CO ₂ exposures in a planted forest in
11	Wisconsin. Overall, the studies at the Aspen FACE experiment were consistent with
12	many of the open-top chamber (OTC) studies that were the foundation of previous O_3
13	NAAQS reviews. These results strengthen the understanding of O ₃ effects on forests and
14	demonstrate the relevance of the knowledge gained from trees grown in open-top
15	chamber studies.
16	Evidence is sufficient to conclude that there is a causal relationship between ambient
17	O_3 exposure and reduced growth of native woody and herbaceous vegetation.

Reproduction 9.4.3.3

18	Studies during recent decades have demonstrated O ₃ effects on various stages of plant
19	reproduction. The impacts of O_3 on reproductive development, as reviewed by <u>Black et</u>
20	al. (2000), can occur by influencing (1) age at which flowering occurs, particularly in
21	long-lived trees that often have long juvenile periods of early growth without flower and
22	seed production; (2) flower bud initiation and development; (3) pollen germination and
23	pollen tube growth; (4) seed, fruit, or cone yields; and (5) seed quality (Table 9-1) (U.S.
24	EPA, 2006b). Several recent studies since the 2006 O_3 AQCD further demonstrate the
25	effects of O ₃ on reproductive processes in herbaceous and woody plant species. Although
26	there have been documented effects of ozone on reproductive processes, a knowledge gap
27	still exists pertaining to the exact mechanism of these responses.
28	Ramo et al. (2007) exposed several meadow species to elevated O_3 (40-50 ppb) and CO_2
29	(+100 ppm), both individually and combined, over three growing seasons in ground-
30	planted mesocosms, using OTCs. Elevated O_3 delayed flowering of <i>Campanula</i>
31	<i>rotundifolia</i> and <i>Vicia cracca</i> . Ozone also reduced the overall number of produced
32	flowers and decreased fresh weight of individual Fragaria vesca berries.

1	Black et al. (2007) exposed Brassica campestris to 70 ppb for two days during late
2	vegetative growth or ten days during most of the vegetative phase. The two-day exposure
3	had no effect on growth or reproductive characteristics, while the 10 day exposure
4	reduced vegetative growth and reproductive site number on the terminal raceme,
5	emphasizing the importance of exposure duration and timing. Mature seed number and
6	weight per pod were unaffected due to reduced seed abortion, suggesting that, although
7	O_3 affected reproductive processes, indeterminate species such as <i>B. campestris</i> possess
8	enough compensatory flexibility to avoid reduced seed production <u>Black et al. (2007</u>).
9	In the determinate species, <i>Plantago major</i> , <u>Black et al. (2010</u>) found that O ₃ may have
10	direct effects on reproductive development in populations of differing sensitivity. Only
11	the first flowering spike was exposed to 120 ppb O_3 for 7 hours per day on 9 successive
12	days (corresponding to flower development) while the leaves and second spike were
13	exposed to charcoal-filtered air. Exposure of the first spike to O ₃ affected seed number
14	per capsule on both spikes even though spike two was not exposed. The combined seed
15	weight of spikes one and two was increased by 19% in the two resistant populations,
16	suggesting an overcompensation for injury; whereas, a decrease of 21% was observed in
17	the most sensitive population (Black et al., 2010). The question remains as to whether
18	these effects are true direct ozone-induced effects or compensatory responses.
19	Studies by Darbah et al. (2008); Darbah et al. (2007) of paper birch (Betula papyrifera)
20	trees at the Aspen FACE site in Rhinelander, WI investigated the effects of elevated O_3
21	and/or CO2 on reproductive fitness. Elevated O3 increased flowering, but decreased seed
22	weight and germination success rate of seeds from the exposed trees. These results
23	suggest that O_3 can dramatically affect flowering, seed production, and seed quality of
24	paper birch, ultimately affecting its reproductive fitness (Darbah et al., 2008; Darbah et
25	<u>al., 2007</u>).

Species	Condition Measures	References
Apocynum androsaemifolium (spreading dogbane)	Flowering time	Bergweiler and Manning (1999)
<i>Buddleia davidii</i> (butterfly bush)	Flowering time	Findley et al. (1997)
Rubus cuneifolius (sand blackberry)	Pollen germination	Chappelka (2002)
Plantago major (plantain)	Pollen tube elongation	<u>Stewart (1998)</u>
<i>Fragaria × ananassa</i> (cultivated strawberry)	Fruit yield	Drogoudi and Ashmore (2001); Drogoudi and Ashmore (2000)
<i>Plantago major</i> (plantain)	Seed yield	<u>Lyons and Barnes (1998); Pearson et al. (1996); Reiling and Davison (1992); Whitfield et al. (1997)</u>
Understory herbs	Seed yield	Harward and Treshow (1975)

Table 9-1	Ozone effects on plant reproductive processes
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Source: Derived from Table AX9-22 of the 2006 O₃ AQCD.

9.4.3.4 Ecosystem Productivity and Carbon Sequestration

1	During the previous NAAQS review, there were limited studies that investigated the
2	effect of O ₃ exposure on ecosystem productivity and C sequestration. Recent studies from
3	long-term FACE experiments provide more evidence of the association of O3 exposure
4	and changes in productivity at the ecosystem level of organization. In addition to
5	experimental studies, model studies also assessed the impact of O3 exposure on
6	productivity and C sequestration from stand to global scales.
7	In this section productivity of ecosystems is expressed in different ways depending on the
8	model or the measurements of a study. The most common metric of productivity is Gross
9	Primary Productivity. Gross Primary Productivity (GPP) is total carbon that enters the
10	ecosystem through photosynthesis by plants. Plants return a larger portion of this carbon
11	back to the atmosphere through respiration from roots and aboveground portions of plants
12	(R_{plant}) . Net primary production (NPP) is the difference between total carbon gain (GPP)
13	and carbon loss through R_{plant} . Net ecosystem productivity (NEP) is the difference
14	between NPP and carbon loss through heterotrophic respiration (R_{het}) (mostly
15	decomposition of dead organic matter) (Lambers et al., 1998). Similarly net ecosystem
16	exchange (NEE) is the net flux of carbon between the land and the atmosphere, typically
17	measured using eddy covariance techniques. Positive values of NEE usually refer to
18	carbon released to the atmosphere (i.e., a source), and negative values refer to carbon
19	uptake (i.e., a sink). Other studies have calculated net carbon exchange (NCE). NCE is
20	defined as NPP minus R_{het} , E_c (the carbon emission during the conversion of natural
21	ecosystems to agriculture) and E_p (the sum of carbon emission from the decomposition of

1	agricultural products). For natural vegetation, E_c and E_p are equal to 0, so NCE is equal
2	NEP (Felzer et al., 2005). In general, modeling studies take into account the effect of O_3
3	on C fixation of a system and there is generally not an effect on R_{plant} , R_{het} , E_c or E_p .
4	Therefore, decreases in GPP, NPP, NEP, NEE and NCE indicate a general decrease in
5	productivity of an ecosystem.
6	Two types of models are most often used to study the ecological consequences of O_3
7	exposure: (1) single plant growth models such as TREGRO and PnET-II (Hogsett et al.,
8	2008; Martin et al., 2001; Ollinger et al., 1997b), and (2) process-based ecosystem
9	models such as PnET-CN, Dynamic Land Ecosystem Model (DLEM), Terrestrial
10	Ecosystem Model (TEM), or Met Office Surface Exchange Scheme - Top-down
11	Representation of Interactive Foliage and Flora Including Dynamics (MOSES-TRIFFID)
12	(Felzer et al., 2009; Ren et al., 2007b; Sitch et al., 2007; Ollinger et al., 2002)
13	(Table 9-2). In these models, carbon uptake is simulated through photosynthesis
14	(TREGRO, PnET - II, PnET- CN, DLEM and MOSES-TRIFFID) or gross primary
15	production (TEM). Photosynthesis rate at leaf level is modeled by a function of stomatal
16	conductance and other parameters in TREGRO, PnET-II, PnET-CN, DLEM and
17	MOSES-TRIFFID. Photosynthesis at canopy level is calculated by summing either
18	photosynthesis of different leaf types (TREGRO, DLEM, and MOSES-TRIFFID) or
19	photosynthesis of different canopy layers (PnET -II, PnET- CN). The detrimental effect
20	of O_3 on plant growth is often simulated by multiplying photosynthesis rate by a
21	coefficient that is dependent on stomatal conductance and cumulative O ₃ uptake
22	(Table 9-2). Different plant functional groups (PFTs, such as deciduous trees, coniferous
23	trees or crops) show different responses to O3 exposure. PnET-iI, PnET-CN, TEM,
24	DLEM and MOSES-TRIFFID estimate this difference by modifying net photosynthesis
25	with coefficients that represent the O ₃ induced fractional reduction of photosynthesis for
26	each functional group. The coefficients used in PnET-iI, PnET-CN, TEM, DLEM are
27	derived from the functions of O ₃ exposure (AOT40) versus photosynthesis reduction
28	from <u>Reich (1987</u>) and <u>Tjoelker et al. (1995</u>). The coefficients used in MOSES-TRIFFID
29	are derived from the O_3 dose-photosynthesis response function from <u>Pleijel et al. (2004a</u>)

are derived from the O_3 dose-photosynthesis response function from <u>Pleijel et al. (2004a</u>) and <u>Karlsson et al. (2004</u>), where O_3 dose is estimated by a metric named CUOt (cumulative stomatal uptake of O_3). The O_3 threshold of CUOt is 1.6 nmol/m²/sec for woody PFT and 5 nmol/m²/sec for grass PFT, and is different from AOT40, which has an O_3 threshold level of 40 ppb for all PFTs. Experimental and model studies on ecosystem productivity and C sequestration at the forest stand scale as well as regional and global scales are reviewed in the following section.

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Model feature	Carbon uptake	Ozone effect	Reference
Hourly or daily step, single plant	Leaf: leaf photosynthesis is a function of stomatal conductance, mesophyll conductance and the gradient of CO_2 from atmosphere to the mesophyll cells	The effect of O_3 on photosynthesis is simulated by reducing mesophyll conductance, and increasing respiration. The degree of O_3 damage is determined	Hogsett et al. (2008); Weinstein et al.
model simulating vegetation growth process	Canopy: Leaf is divided into different ages. The canopy photosynthesis rate is the sum of the photosynthesis of all foliage groups	by ambient O_3 exposure, and the threshold O_3 concentration below which O_3 does not affect mesophyll conductance and respiration	<u>(2005</u>); <u>Tingey et</u> <u>al. (2004</u>)
PnET-il: Monthly time-step, single plant model PnET – CN: Monthly time-step, ecosystem model	Leaf: Maximum photosynthesis rate is determined by a function of foliar N concentration, and stomatal conductance is determined by a function of the actual rate of the photosynthesis. Canopy: canopy is divided into multiple, even- mass layers and photosynthesis is simulated by a multilayered canopy submodel	The effect of O_3 on photosynthesis is simulated by an equation of stomatal conductance and O_3 dose (AOT40). The model assumes that photosynthesis and stomatal conductance remain coupled under O_3 exposure, with a reduction in photosynthesis for a given month causing a proportion reduction in stomatal conductance.	Ollinger et al. (2002); Ollinger et al. (1997b); Pan et al. (2009)
Monthly time-step, ecosystemEcosystem: TEM is run at a 0.5*0.5 degree resolution. Each grid cell is classified by vegetation type and soil texture, and vegetation and detritus are assumed to distribute homogeneously within grid cells. Carbon flows into ecosystem via gross primary production, which is a function of maximum rate of assimilation, photosynthetically active radiation, the leaf area relative to the maximum annual leaf area, mean monthly air temperate, and nitrogen availability.		The direct O_3 reduction on GPP is simulated by multiplying GPP by $f(O_3)t$, where $f(O_3)t$ is determined by evapotranspiration, mean stomatal conductance, ambient AOT40, and empirically O_3 response coefficient derived from previous publications.	Felzer et al. (2005); (2004)
Daily time-step ecosystem model	Leaf: photosynthesis is a function of 6 parameters: photosynthetic photon flux density, stomatal conductance, daytime temperature, the atmospheric CO_2 concentration, the leaf N content and the length of daytime.	The detrimental effect of O_3 is simulated by multiplying the rate of photosynthesis by O_3 eff, where O_3 eff is a function of stomatal conductance, ambient AOT40, and O_3 sensitive coefficient. Ozone's	<u>Ren et al.</u> (2007b); (Ren et al., 2007a); Zhang et al.
	Canopy: Photosynthetic rates for sunlit leaf and shaded leaf scale up to the canopy level by multiplying the estimated leaf area index Ecosystem: GPP is the sum of gross C fixation of different plant function groups	indirect effect on stomatal conductance is also simulated, with a reduction in photosynthesis for a given month causing a reduction in stomatal conductance, and therefore canopy conductance.	<u>(2007a</u>)
30 minute time-step, dynamic global vegetation model	Leaf: photosynthesis is a function of environmental and leaf parameters and stomatal conductance; Stomatal conductance is a function of the concentration of CO_2 and H_2O in air at the leaf surface and the current rate of photosynthesis of the leaf	The effect of O_3 is simulated by multiplying the rate of photosynthesis by F, where F depends upon stomatal conductance, O_3 exposure, a critical threshold for O_3 damage, and O_3 sensitive coefficient (functional type dependent)	<u>Sitch et al.</u> (2007)
	Canopy: Photosynthetic rates scale up to the canopy level by multiplying a function of leaf area index and PAR extinction coefficient Ecosystem: GPP is the sum of gross C fixation of		
	feature Hourly or daily step, single plant model simulating vegetation growth process PnET-il: Monthly time-step, single plant model PnET – CN: Monthly time-step, ecosystem model Monthly time-step, ecosystem model	featureHourly or daily step, single plant modelLeaf: leaf photosynthesis is a function of stomatal conductance, mesophyll conductance and the gradient of CO2 from atmosphere to the mesophyll cells Canopy: Leaf is divided into different ages. The canopy photosynthesis rate is the sum of the photosynthesis of all foliage groupsPnET-il: Monthly time-step, single plant modelLeaf: Maximum photosynthesis rate is determined by a function of foliar N concentration, and stomatal conductance is determined by a function of the actual rate of the photosynthesis. Canopy: canopy is divided into multiple, even- mass layers and photosynthesis is simulated by a multilayered canopy submodelMonthly time-step, ecosystem modelEcosystem: TEM is run at a 0.5*0.5 degree resolution. Each grid cell is classified by vegetation type and soli texture, and vegetation and detritus are assumed to distribute homogeneously within grid cells. Carbon flows into ecosystem via gross primary production, which is a function of maximum rate of assimilation, photosynthetically active radiation, the leaf area relative to the maximum annual leaf area, mean monthly air temperature, the atmospheric CO2 concentration, the leaf N content and the length of daytime. Canopy: Photosynthetic rates for sunit leaf and shaded leaf scale up to the canopy level by multiplying the estimated leaf area index Ecosystem: GPP is the sum of gross C fixation of different plant function groups30 minute time-step, dynamic global vegetation modelLeaf: photosynthesis is a function of environmental and leaf parameters and stomatal conductance, Stomatal conductance is a function of the concentration of CO2 and H ₂ O in air at the leaf surface and the current rate	feature Houry or and the gradient of CO ₂ from atmosphere to the plant mesophylicells Leaf: leaf photosynthesis is a function of daily step, isopie The effect of O ₂ on photosynthesis is stomatal conductance, mesophylicenductance and the gradient of CO ₂ from atmosphere to the simulation; growth The effect of O ₂ on photosynthesis is stomatal conductance, and increasing respiration. The degree of O ₂ damage is determined by arbitron D ₃ concentration below which O ₃ does not affect mesophylic conductance and respiration PinET-I: Nonthly time-step, PinET-R Leaf: Maximum photosynthesis rate is determined by a function of toiar photosynthesis is simulated by a concentration, and stomatal conductance and O ₂ does (AOT40). The model assumes that photosynthesis is simulated by an equation of stomatal conductance and O ₂ does (AOT40). The model assumes that photosynthesis is simulated by an equation of stomatal conductance and O ₂ does (AOT40). The model assumes that photosynthesis is simulated by an equation of stomatal conductance. Monthly time-step, ecosystem Ecosystem: TEM is run at a 0.5*0.5 degree resolution. Each grid cell is Carbino flows into ecosystem via gross primary production, which is a function of maximum rate of assimilation, photosynthesic assumed to distribute formogeneously within grid cells. Carbon flows into ecosystem via gross primary production, which is a function of maximum rate of assimilation, photosynthesis is a function of 6 parameters: photosynthesis is a function of 6 parameters: photosynthesis is a function of stomatal conductance, damine the consystem: GPP is the sum of gross C fixation of the concentration of Go, and gross maximal conductance, ambient AOT40, and O ₃ sensitive coefficient. Czone's alos initaleconductance, ambient AOT40, and O ₃ sensinitic ceffi

Table 9-2Comparison of models used to simulate the ecological
consequences of ozone exposure.

Local scale

1	Both experimental and modeling studies have provided new information on effects of O ₃
2	exposure at the stand or site level, i.e., at the local scale. The above- and below-ground
3	biomass and net primary production (NPP) were measured at the Aspen FACE site after 7
4	years of O_3 exposure. Elevated O_3 caused 23, 13 and 14% reductions in total biomass
5	relative to the control in the aspen, aspen-birch and aspen-maple communities,
6	respectively (King et al., 2005). At the Kranzberg Forest FACE experiment in Germany,
7	O3 reduced annual volume growth by 9.5 m3/ha in a mixed mature stand of Norway
8	spruce and European beech (Pretzsch et al., 2010). At the grassland FACE experiment at
9	Alp Flix, Switzerland, O3 reduced the seasonal mean rates of ecosystem respiration and
10	GPP by 8%, but had no significant impacts on aboveground dry matter productivity or
11	growing season net ecosystem production (NEP) (Volk et al., 2011). Ozone also altered
12	C accumulation and turnover in soil, as discussed in Section <u>9.4.6</u> .
13	Changes in forest stand productivity under elevated O ₃ were assessed by several model
14	studies. TREGRO (Table 9-2) has been widely used to simulate the effects of O_3 on the
15	growth of several species in different regions in the U.S. Hogsett et al. (2008) used
16	TREGRO to evaluate the effectiveness of various forms and levels of air quality
17	standards for protecting tree growth in the San Bernardino Mountains of California. They
18	found that O ₃ exposures at the Crestline site resulted in a mean 20.9% biomass reduction
19	from 1980 to 1985 and 10.3% biomass reduction from 1995 to 2000, compared to the
20	"background" O3 concentrations (O3 concentration in Crook County, Oregon). The level
21	of vegetation protection projected was different depending on the air quality scenarios
22	under consideration. Specifically, when air quality was simulated to just meet the
23	California 8-h average maximum of 70 ppb and the maximum 3 months 12-h SUM06 of
24	25 ppm-h, annual growth reductions were limited to 1% or less, while air quality that just
25	met a previous NAAQS (the second highest 1-h max [125 ppb]) resulted in 6-7% annual
26	reduction in growth, resulting in the least protection relative to background O_3 (Hogsett et
27	<u>al., 2008</u>).
28	ZELIG is a forest succession gap model, and has been used to evaluate the dynamics of
29	natural stand succession. Combining TREGRO with ZELIG, Weinstein et al. (2005)
30	simulated the effects of different O_3 levels (0.5, 1.5, 1.75, and 2 times [×] ambient) on the
31	growth and competitive interactions of white fir and ponderosa pine at three sites in
32	California: Lassen National Park, Yosemite National Park, and Crestline. Their results
33	suggested that O_3 had little impact on white fir, but greatly reduced the growth of
34	ponderosa pine. If current O_3 concentrations continue over the next century, ambient O_3
35	exposure (SUM06 of 110 ppm-h) at Crestline was predicted to decrease individual tree
36	C budget by 10% and decrease ponderosa pine abundance by 16%. Effects at Lassen

National Park and Yosemite National Park sites were found to be smaller because of lower O_3 exposure levels (<u>Weinstein et al., 2005</u>).

3 To evaluate the influence of interspecies competition on O_3 effects, the linked TREGRO 4 and ZELIG modeling system was used to predict the effects of O_3 over 100 years on the 5 basal area of species in a Liriodendron tulipifera-dominated forest in the Great Smoky 6 Mountains National Park (Weinstein et al., 2001). Ambient O_3 was predicted to decrease 7 individual tree C budget by 28% and reduce the basal area of L. tulipifera by 10%, 8 whereas a 1.5×-ambient exposure was predicted to cause a 42% decrease in the individual 9 tree C budget and a 30% reduction in basal area. Individual tree C balance for Acer 10 rubrum decreased 14% and 23% under ambient and 1.5×-ambient exposure, respectively. Prunus serotina was predicted to have less than a 2% decrease in tree C balance in all 11 12 scenarios, but its basal area was greatly altered by the O_3 effects on the other tree species. 13 Basal area of A. rubrum and P. serotina was predicted to increase for some years, but 14 then decrease by up to 30%, depending on the scenario. The effects of O_3 on stand 15 productivity and dynamics were also studied by other tree growth or stand models, such 16 as ECOPHYS, INTRASTAND and LINKAGES. ECOPHYS is a functional-structural 17 tree growth model. The model used the linear relationship between the maximum 18 capacity of carboxylation and O₃ dose to predict the relative effect of O₃ on leaf 19 photosynthesis (Martin et al., 2001). Simulations with ECOPHYS found that O₃ 20 decreased stem dry matter production, stem diameter and leaf dry matter production, 21 induced earlier leaf abscission, and inhibited root growth (Martin et al., 2001). 22 INTRASTAND is an hourly time step model for forest stand carbon and water budgets. 23 LINKAGES is a monthly time step model simulating forest growth and community 24 dynamics. Linking INTRASTAND with LINKAGES, Hanson et al. (2005) found that a 25 simulated increase in O_3 concentration in 2100 (a mean 20-ppb increase over the current 26 O₃ concentration) yields a 35% loss of net ecosystem C exchange (NEE) with respect to the current conditions (174 g $C/m^2/year$). 27

Regional and global scales

- Since the publication of the 2006 O₃ AQCD, there is additional evidence suggesting that
 O₃ exposure alters ecosystem productivity and biogeochemical cycling at the regional
 scale, i.e., at scales ranging from watershed to subcontinental divisions, and at continental
 and global scales. Most of those studies were conducted by using process-based
 ecosystem models (Table 9-2) and are briefly reviewed in the following sections.
 Ollinger et al. (1997a) simulated the effect of O₃ on hardwood forest productivity of 64
 hardwood sites in the northeastern U.S. with PnET-iI (Table 9-2). Their simulations
 - indicated that O_3 caused a 3-16% reduction in NPP from 1987 to 1992 (<u>Table 9-3</u>). The

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1	interactive effects of O ₃ , N deposition, elevated CO ₂ and land use history on C dynamics
2	were estimated by PnET-CN (Table 9-2) (Ollinger et al., 2002). The results indicated that
3	O_3 offset the increase in net C exchange caused by elevated CO_2 and N deposition by
4	13% (25.0 g C/m ² /year) under agriculture site history, and 23% (33.6 g C/m ² /year) under
5	timber harvest site history. PnET-CN was also used to assess changes in C sequestration
6	of U.S. Mid-Atlantic temperate forest. Pan et al. (2009) designed a factorial modeling
7	experiment to separate the effects of changes in atmospheric composition, historical
8	climatic variability and land-disturbances on the C cycle. They found that O_3 acted as a
9	negative factor, partially offsetting the growth stimulation caused by elevated CO_2 and
10	N deposition in U.S. Mid-Atlantic temperate forest. Ozone decreased NPP of most forest
11	types by 7-8%. Among all the forest types, spruce-fir forest was most resistant to O_3
12	damage, and NPP decreased by only 1% (Pan et al., 2009).

- 13 Felzer et al. (2004) developed TEM 4.3 (Table 9-2) to simulate the effects of O_3 on plant 14 growth and estimated effects of O₃ on NPP and C sequestration of deciduous trees, 15 conifers and crops in the conterminous U.S. The results indicated that O_3 reduced NPP 16 and C sequestration in the U.S. (Table 9-3) with the largest decreases (over 13% in some 17 locations) in NPP occurring in the Midwest agricultural lands during the mid-summer. 18 TEM was also used to evaluate the magnitude of O_3 damage at the global scale 19 (Table 9-3) (Felzer et al., 2005). Simulations for the period 1860 to 1995 show that the 20 largest reductions in NPP and net C exchange occurred in the mid western U.S., eastern 21 Europe, and eastern China (Felzer et al., 2005). DLEM (Table 9-2) was developed to 22 simulate the detrimental effect of O_3 on ecosystems, and has been used to examine the O_3 23 damage on NPP and C sequestration in Great Smoky Mountains National Park (Zhang et 24 al., 2007a), grassland ecosystems and terrestrial ecosystems in China (Ren et al., 2007b; 25 Ren et al., 2007a). Results of those simulations are listed in Table 9-3.
- 26 Instead of using AOT40 as their O₃ exposure metric as PnET, TEM and DLEM did, Sitch 27 et al. (2007) incorporated a different O_3 metric named CUOt (cumulative stomatal uptake 28 of O₃), derived from Pleijel et al. (2004a), into the MOSES-TRIFFID coupled model 29 (Table 9-2). In the CUOt metric, the fractional reduction of plant production is dependent 30 on O_3 uptake by stomata over a critical threshold for damage with this threshold level 31 varying by plant functional type. Consistent with previous studies, their model simulation 32 indicated that O₃ reduced global gross primary production (GPP), C-exchange rate and 33 C sequestration (Table 9-3). The largest reductions in GPP and land-C storage were 34 projected over North America, Europe, China and India. In the model, reduced ecosystem 35 C uptake due to O_3 damage results in additional CO_2 accumulation in the atmosphere and 36 an indirect radiative forcing of climate change. Their simulations indicated that the indirect radiative forcing caused by O_3 (0.62-1.09 W/m²) could have even greater impact 37 on global warming than the direct radiative forcing of O_3 (0.89 W/m²) (Sitch et al., 2007). 38

1	Results from the various model studies presented in <u>Table 9-3</u> are difficult to compare
2	because of the various spatial and temporal scales used. However, all the studies showed
3	that O_3 exposure decreased ecosystem productivity and C sequestration. These results are
4	consistent and coherent with experimental results obtained from studies at the leaf, plant
5	and ecosystem scales (Sitch et al., 2007; Felzer et al., 2005). Many of the models use the
6	same underlying function to simulate the effect of O_3 exposure to C uptake. For example
7	the functions of O ₃ exposure (AOT40) versus photosynthesis reduction for PnET-iI,
8	PnET-CN, TEM, DLEM were all from <u>Reich (1987</u>) and <u>Tjoelker et al. (1995</u>).
9	Therefore, it is not surprising that the results are similar. While these models can be
10	improved and more evaluation with experimental data can be done, these models
11	represent the state of the science for estimating the effect of O ₃ exposure on productivity
12	and C sequestration.

9.4.3.5 Summary

13	During the previous NAAQS reviews, there were very few studies that investigated the
14	effect of O ₃ exposure on ecosystem productivity and C sequestration. Recent studies from
15	long-term FACE experiments, such as Aspen FACE, SoyFACE and the Kranzberg Forest
16	(Germany), provide evidence of the association of O ₃ exposure and reduced productivity
17	at the ecosystem level of organization. Studies at the leaf and plant scales show that O_3
18	decreased photosynthesis and plant growth, which provides coherence and biological
19	plausibility for the decrease in ecosystem productivity. Results across different ecosystem
20	models, such as TREGRO, PnET, TEM and DLEM, are consistent with the FACE
21	experimental evidence, which show that O3 reduced productivity of various ecosystems.
22	Productivity is measured by various metrics such as GPP, NPP, NEP, NCE, NEE and
23	individual tree biomass gain. All these metrics indicate a decrease in CO ₂ fixation by the
24	systems that were studied.
25	Although O ₃ generally causes negative effects on plant growth, the magnitude of the
26	response varies among plant communities. For example, O ₃ had little impact on white fir,
27	but greatly reduced growth of ponderosa pine in southern California (Weinstein et al.,
28	2005). Ozone decreased net primary production (NPP) of most forest types in the Mid-
29	Atlantic region, but had small impacts on spruce-fir forest (Pan et al., 2009).
30	In addition to plant growth, other indicators that are typically estimated by model studies
31	include net ecosystem CO ₂ exchange (NEE), C sequestration, and crop yield. Model
32	simulations consistently found that O ₃ exposure caused negative impacts on these
33	indicators, but the severity of these impacts was influenced by multiple interactions of
34	biological and environmental factors. The suppression of ecosystem C sinks results in

1	more CO ₂ accumulation in the atmosphere. Globally, the indirect radiative forcing caused
2	by O ₃ exposure through lowering the ecosystem C sink could have an even greater impact
3	on global warming than the direct radiative forcing of O_3 (Sitch et al., 2007). Ozone
4	could also affect regional C budgets through interacting with multiple factors, such as
5	N deposition, elevated CO_2 and land use history. Model simulations suggested that O_3
6	partially offset the growth stimulation caused by elevated CO ₂ and N deposition in both
7	Northeast- and Mid-Atlantic-region forest ecosystems of the U.S. (Pan et al., 2009;
8	<u>Ollinger et al., 2002</u>).
9	The evidence is sufficient to infer that there is a causal relationship between O_3
10	exposure and reduced productivity, ${ m and}$ a likely causal relationship between O $_3$
11	exposure and reduced carbon sequestration in terrestrial ecosystems.

	Scale	Model	Index	O ₃ Impacts	Reference
GPP	Global	MOSES- TRIFFID	CUOt ^a	Decreased by 14-23% over the period 1901-2100	<u>Sitch et al.</u> (2007)
NPP	Global	TEM	AOT40	Decreased by 0.8% without agricultural management and a decrease of 2.9% with optimal agricultural management	<u>Felzer et al.</u> (2005)
	U.S.	TEM	AOT40	Reduced by 2.3% without optimal N fertilization and 7.2% with optimal N fertilization from 1983-1993	<u>Felzer et al.</u> (2005)
	U.S.	TEM	AOT40	Reduced by 2.6–6.8% during the late 1980s to early 1990s.	<u>Felzer et al.</u> (2004)
	Northeastern U.S.	PnET	AOT40	A reduction of 3-16% from 1987-1992	<u>Ollinger et al</u> (1997a)
	U.S. Mid- Atlantic	PnET	AOT40	Decreased NPP of most forest types by 7-8%	<u>Pan et al.</u> (2009)
	China	DLEM	AOT40	Reduced NPP of grassland in China by 8.5 Tg ^b C from 1960s to 1990s	<u>Ren et al.</u> (2007a)
C exchange	Global	TEM	AOT40	Reduced net C exchange (1950–1995) by 0.1 Pg C/yr without agricultural management and 0.3 Pg C/yr with optimal agricultural management	<u>Felzer et al.</u> (2005)
	Global	MOSES- TRIFFID	CUOt	Decreased global mean land–atmosphere C fluxes by 1.3 Pg C/yr and 1.7 Pg C/yr for the 'high' and 'low' plant O_3 sensitivity models, respectively	<u>Sitch et al.</u> (2007)
C sequestration	Global	MOSES- TRIFFID	CUOt	Reduced land-C storage accumulation by between 143 Pg C/yr and 263 Pg C/yr from 1900–2100	<u>Sitch et al.</u> (2007)
	U.S.	TEM	AOT40	Reduced C sequestration by 18–38 Tg C/yr from 1950 to 1995	<u>Felzer et al.</u> (2004)
	GSM National Park	DLEM	AOT40	Decreased the ecosystem C storage of deciduous forests by 2.5% and pine forest by 1.4% from 1971 to 2001	<u>Zhang et al.</u> (2007a)
	China	DLEM	AOT40	Reduced total C storage by 0.06% in 1960s and 1.6% in 1990s in China's terrestrial ecosystems	<u>Ren et al.</u> (2007b)
	China	DLEM	AOT40	O_3 exposure reduced the net C sink of China's terrestrial ecosystem by 7% from 1961 to 2005	<u>Tian et al.</u> (2011)
	China	DLEM	AOT40	Ozone induced net carbon exchange reduction ranged from 0.4-43.1%, depending on different forest type	<u>Ren et al.</u> (2011)

Table 9-3Modeled effects of ozone on primary production, C exchange,
and C sequestration.

^aCUOt is defined as the cumulative stomatal uptake of O₃, using a constant O₃-uptake rate threshold of t nmol/m²/sec. ^bPg equals 1×10^{15} grams.

9.4.4 Crop Yield and Quality in Agricultural Systems

1	The detrimental effect of O_3 on crop production has been recognized since the 1960s and
2	a large body of research has stemmed from that recognition. Previous O_3 AQCDs have
3	extensively reviewed this body of literature. <u>Table 9-4</u> summarizes recent experimental
4	studies of O_3 effects on agricultural crops, exclusive of growth and yield. Growth and
5	yield results are summarized in <u>Table 9-17</u> .

1 The actual concentration and duration threshold	for O ₃ damage varies from species to
2 species and sometimes even among genotypes of	of the same species (Guidi et al., 2009;
3 Sawada and Kohno, 2009; Biswas et al., 2008; A	Ariyaphanphitak et al., 2005; Dalstein and
4 <u>Vas, 2005; Keutgen et al., 2005</u>). A number of c	comprehensive reviews and meta-analyses
5 have recently been published discussing both th	e current understanding of the
6 quantitative effects of O ₃ concentration on a var	iety of crop species and the potential
7 focus areas for biotechnological improvement to	o a future growing environment that will
8 include higher O ₃ concentrations (Bender and W	Veigel, 2011; Booker et al., 2009;
9 <u>VanDingenen et al., 2009; Ainsworth, 2008; Fer</u>	ng et al., 2008; <u>Hayes et al., 2007; Mills</u>
10 <u>et al., 2007; Grantz et al., 2006; Morgan et al., 2</u>	2003). Since the 2006 O ₃ AQCD (<u>U.S.</u>
11 <u>EPA, 2006b</u>), exposure-response indices for a v	ariety of crops have been suggested
12 (<u>Mills et al., 2007a</u>) and many reports have inve	estigated the effects of O ₃ concentration
13 on seed or fruit quality to extend the knowledge	base beyond yield quantity. This section
14 will outline the key findings from these papers a	as well as highlight some of the recent
15 research addressing the endpoints such as yields	s and crop quality.

16This section will also highlight recent literature that focuses on O_3 damage to crops as17influenced by other environmental factors. Genetic variability is not the only factor that18determines crop response to O_3 damage. Ozone concentration throughout a growing-19season is not homogeneous and other environmental conditions such as elevated CO_2 20concentrations, drought, cold or nutrient availability may alleviate or exacerbate the21oxidative stress response to a given O_3 concentration.

9.4.4.1 Yield

22	It is well known that yield is negatively impacted in many crop species in response to
23	high O ₃ concentration. However, the concentrations at which damage is observed vary
24	from species to species. Numerous analyses of experiments conducted in OTCs and with
25	naturally occurring gradients demonstrate that the effects of O_3 exposure also vary
26	depending on the growth stage of the plant; plants grown for seed or grain are often most
27	sensitive to exposure during the seed or grain-filling period (Soja et al., 2000; Pleijel et
28	al., 1998; Younglove et al., 1994; Lee et al., 1988a). AX9.5.4.1 of the 2006 O ₃ AQCD
29	summarized many previous studies on crop yield.

Field studies and meta-analyses

30The effect of O3 exposure on U.S. crops remains an important area of research and31several studies have been published on this topic since the 2006 O3 AQCD (U.S. EPA,322006b) (Table 9-4 and Table 9-17). For example, one study with cotton in a crop-weed

- 1 interaction study (Grantz and Shrestha, 2006) utilizing OTCs suggests that 12-hour 2 average O_3 concentrations of 79.9 ppb decreased cotton biomass by 25% and 12-hour 3 average O_3 concentration of 122.7 ppb decreased cotton biomass by 75% compared to 4 charcoal filtered control (12-h avg: 12.8 ppb). Further, this study suggests that the weed, 5 vellow nutsedge, was less sensitive to increasing O_3 concentration, which would increase 6 weed competition (Grantz and Shrestha, 2006). In a study of peanuts in North Carolina, 7 near ambient and elevated exposures of O_3 reduced photosynthesis and yield compared to 8 very low O₃ conditions (Booker et al., 2007; Burkey et al., 2007). In another study, 9 Grantz and Vu (2009) reported that sugarcane biomass growth significantly declined 10 under O_3 exposure.
- 11 The average yield loss reported across a number of meta-analytic studies have been 12 published recently for soybean (Morgan et al., 2003), wheat (Feng et al., 2008b), rice 13 (Ainsworth, 2008), semi-natural vegetation (Hayes et al., 2007), potato, bean and barley 14 (Feng and Kobayashi, 2009). Meta-analysis allows for the objective development of a 15 quantitative consensus of the effects of a treatment across a wide body of literature. 16 Further, this technique allows for a compilation of data across a range of O_3 fumigation 17 techniques, durations and concentrations in order to assemble the existing literature in a 18 meaningful manner.
- 19 Morgan et al. (2003) reported an average seed yield loss for soybean of 24% compared to 20 charcoal filtered air across all O₃ concentrations used in the 53 compiled studies. The 21 decrease in seed yield appeared to be the product of nearly equal decreases (7-12%) in 22 seed weight, seed number and pod number. As would be expected, the lowest O₃ 23 concentration (30-59 ppb) resulted in the smallest yield losses, approximately 8%, while 24 the highest O₃ concentration (80-120 ppb) resulted in the largest yield losses, 25 approximately 35% (Morgan et al., 2003). Further, the oil/protein ratio within the 26 soybean seed was altered due to growth at elevated O_3 concentrations, with a decrease in 27 oil content. The studies included in this meta-analysis all used enclosed fumigation 28 systems or growth chambers which may have altered the coupling of the atmosphere to 29 the lower plant canopy (McLeod and Long, 1999), although the results of Morgan et al. 30 (2006), Betzelberger et al. (2010), and the comparisons presented in Section 9.6.3 31 strongly suggest that decreases in yield between ambient and elevated exposures are not 32 affected by exposure method. Utilizing the Soybean Free Air gas Concentration 33 Enrichment Facility (SoyFACE; www.soyface.illinois.edu). Morgan et al. (2006) 34 reported a 20% seed yield loss due to a 23% increase in average daytime O_3 35 concentration (56-69 ppb) within a single soybean cultivar across two growing seasons in 36 Illinois, which lies within the range predicted by the meta-analysis. A further breakdown 37 of the effects of current O_3 concentrations (AOT40 of 4.7 ppm-h) on bean seed quality 38 (*Phaseolus vulgaris*) has identified that growth at current O₃ concentrations compared to

1	charcoal-filtered air raised total lipids, total crude protein and dietary fiber content (Iriti et
2	al., 2009). An increase in total phenolics was also observed, however the individual
3	phenolic compounds responded differently, with significant decreases in anthocyanin
4	content. The seeds from ambient O ₃ exposed plants also displayed increased total
5	antioxidant capacity compared to charcoal-filtered air controls (Iriti et al., 2009).
6	Betzelberger et al. (2010) has recently utilized the SoyFACE facility to compare the
7	impact of elevated O ₃ concentrations across 10 soybean cultivars to investigate
8	intraspecific variability of the O ₃ response to find physiological or biochemical markers
9	for eventual O_3 tolerance breeding efforts (<u>Betzelberger et al., 2010</u>). They report an
10	average 17% decrease in yield across all 10 cultivars across two growing seasons due to a
11	doubling of ambient O ₃ concentrations, with the individual cultivar responses ranging
12	from -7% to -36%. The exposure-response functions derived for these 10 current
13	cultivars were similar to the response functions derived from the NCLAN studies
14	conducted in the 1980s (Heagle, 1989), suggesting there has not been any selection for
15	increased tolerance to O ₃ in more recent cultivars. More complete comparisons between
16	yield predictions based on data from cultivars used in NCLAN studies, and yield data for
17	modern cultivars from SoyFACE are reported in Section 9.6.3 of this document. They
18	confirm that the response of soybean yield to O ₃ exposure has not changed in current
19	cultivars.
20	A meta-analysis has also been performed on studies investigating the effects of O ₃

- 21 concentrations on wheat (Feng et al., 2008b). Across 23 studies included, elevated O₃ 22 concentrations (ranging from a 7-h daily average of 31-200 ppb) decreased grain yield by 23 29%. Winter wheat and spring wheat did not differ in their responses; however the 24 response in both varieties to increasing O_3 concentrations resulted in successively larger 25 decreases in yield, from a 20% decrease in 42 ppb to 60% in 153 ppb O_3 . These yield 26 losses were mainly caused by a combination of decreases in individual grain weight 27 (-18%), ear number per plant (-16%), and grain number per ear (-11%). Further, the grain 28 starch concentration decreased by 8% and the grain protein yield decreased by 18% due 29 to growth at elevated O₃ concentrations as well. However, increases in grain calcium and 30 potassium levels were reported (Feng et al., 2008b).
- 31A recent meta-analysis found that growth at elevated O3 concentrations negatively32impacts nearly every aspect of rice performance as well (Ainsworth, 2008). While rice is33not a major crop in the U.S., it provides a staple food for over half of the global34population (IRRI, 2002) and the effects of rising O3 concentrations on rice yields merit35consideration. On average, rice yields decreased 14% in 62 ppb O3 compared to charcoal-36filtered air. This yield loss was largely driven by a 20% decrease in grain number37(Ainsworth, 2008).

1	Feng and Kobayashi (2009) have recently compiled yield data for six major crop species,
2	potato, barley, wheat, rice, bean and soybean and grouped the O ₃ treatments used in those
3	studies into three categories: baseline O_3 concentrations (<26 ppb), current ambient 7- or
4	12-h daily O_3 concentrations (31-50 ppb), and future ambient 7- or 12-h daily O_3
5	concentrations (51-75 ppb). Using these categories, they have effectively characterized
6	the effects of current O_3 concentrations and the effects of future O_3 concentrations
7	compared to baseline O ₃ concentrations. At current O ₃ concentrations, which ranged from
8	41-49 ppb in the studies included, soybean (-7.7%), bean (-19.0%), barley (-8.9%), wheat
9	(-9.7%), rice (-17.5%) and potato (-5.3%) all had yield losses compared to the baseline
10	O_3 concentrations (<26 ppb). At future O_3 concentrations, averaging 63 ppb, soybean
11	(-21.6%), bean (-41.4%), barley (-14%), wheat (-28%), rice (-17.5%) and potato (-11.9%)
12	all had significantly larger yield losses compared to the losses at current O_3
13	concentrations (<26 ppb) (Feng and Kobayashi, 2009).
14	A review of OTC studies has determined the AOT40 critical level that causes a 5% yield
15	reduction across a variety of agricultural and horticultural species (Mills et al., 2007a).
16	The authors classify the species studied into three groups: sensitive, moderate and
17	tolerant. The sensitive crops, including watermelon, beans, cotton, wheat, turnip, onion,
18	soybean, lettuce, and tomato, respond with a 5% reduction in yield under a 3-month
19	AOT40 of 6 ppm-h. Watermelon was the most sensitive with a critical level of
20	1.6 ppm-h. The moderately sensitive crops, including sugar beet, oilseed rape, potato,
21	tobacco, rice, maize, grape and broccoli, responded with a 5% reduction in yield between
22	8.6 and 20 ppm-h. The crops classified as tolerant, including strawberry, plum and barley,
23	responded with a 5% yield reduction between 62-83.3 ppm-h (Mills et al., 2007a).
24	Feng and Kobayashi (2009) compared their exposure-response results to those published
25	by Mills et al. (2007a) and found the ranges of yield loss to be similar for soybean, rice
26	and bean. However, Feng and Kobayashi (2009) reported smaller yield losses for potato
27	and wheat and larger yield losses for barley compared to the dose-response functions
28	published by Mills et al. (2007a), which they attributed to their more lenient criteria for
29	literature inclusion.
30	While the studies investigating the impact of various O ₃ concentrations on yield are
31	important and aid in determining the vulnerability of various crops to a variety of O_3
32	concentrations, there is still uncertainty as to how these crops respond under field
33	conditions with interacting environmental factors such as temperature, soil moisture, CO_2
34	concentration, and soil fertility (Booker et al., 2009). Further, there appears to be a
35	distinct developmental and genotype dependent influence on plant sensitivity to O ₃ that
36	has yet to be fully investigated across O ₃ concentrations in a field setting. The potentially
37	mitigating effect of breeding selection for O ₃ resistance has received very little attention

in the published scientific literature. Anecdotal reports suggest that such selection may
 have occurred in recent decades for some crops in areas of the country with high ambient
 exposures. However, the only published literature available is on soybean and these
 studies indicate that sensitivity has not changed in cultivars of soybean between the
 1980s and the 2000s (Betzelberger et al., 2010). This conclusion for soybeans is
 confirmed by comparisons presented in Section 9.6.3 of this document.

Yield loss at regional and global scales

- 7 Because O_3 is heterogeneous in both time and space and O_3 monitoring stations are 8 predominantly near urban areas, the impacts of O₃ on current crop yields at large 9 geographical scales are difficult to estimate. Fishman et al. (2010) have used satellite 10 observations to estimate O₃ concentrations in the contiguous tri-state area of Iowa, 11 Illinois and Indiana and have combined that information with other measured 12 environmental variables to model the historical impact of O₃ concentrations on soybean 13 vield across the 2002-2006 growing seasons. When soybean yield across Iowa, Indiana 14 and Illinois was modeled as a function of seasonal temperature, soil moisture and O_3 15 concentrations, O₃ had the largest contribution to the variability in yield for the southern-16 most latitudes included in the dataset. Fishman et al. (2010) determined that O_3 17 concentrations significantly reduced soybean yield by 0.38 to 1.63% for every 18 additional ppb of exposure across the 5 years. This value is consistent with previous 19 chamber studies (Heagle, 1989) and results from SoyFACE (Morgan et al., 2006). 20 Satellite estimates of tropospheric O_3 concentrations exist globally (Fishman et al., 2008), 21 therefore utilizing this historical modeling approach is feasible across a wider 22 geographical area, longer time-span and perhaps for more crop species.
- 23 The detrimental effects of O_3 on crop production at regional or global scales were also 24 assessed by several model studies. Two large scale field studies were conducted in the 25 U.S. (NCLAN) and in Europe (European Open Top Chamber Programme, EOTCP) to 26 assess the impact of O_3 on crop production. Ozone exposure-response regression models 27 derived from the two programs have been widely used to estimate crop yield loss 28 (Avnery et al., 2011a, b; VanDingenen et al., 2009; Tong and Mauzerall, 2008; Wang and 29 Mauzerall, 2004). Those studies found that O₃ generally reduced crop yield and that 30 different crops showed different sensitivity to O₃ pollution (
- 31Table 9-5). Ozone was calculated to induce a possible 45-82 million metric tons loss for32wheat globally. Production losses for rice, maize and soybean were on the order of3317-23 million metric tons globally (VanDingenen et al., 2009). The largest yield losses34occur in high-production areas exposed to high O3 concentrations, such the Midwest and

the Mississippi Valley regions in the U.S., Europe, China and India (<u>VanDingenen et al.</u>, <u>2009</u>; <u>Tong et al.</u>, <u>2007</u>).

9.4.4.2 Crop Quality

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2

3 In general, it appears that increasing O₃ concentrations above current ambient 4 concentrations can cause species-dependent biomass losses, decreases in root biomass 5 and nutritive quality, accelerated senescence and shifts in biodiversity. A study conducted 6 with highbush blackberry has demonstrated decreased nutritive quality with increasing O_3 7 concentration despite no change in biomass between charcoal-filtered control, ambient O₃ 8 and $2 \times$ ambient O₃ exposures (Ditchkoff et al., 2009). A study conducted with sedge 9 using control (30 ppb), low (55 ppb), medium (80 ppb) and high (105 ppb) O_3 treatments 10 has demonstrated decreased root biomass and accelerated senescence in the medium and 11 high O₃ treatments (Jones et al., 2010). Alfalfa showed no biomass changes across 12 two years of double ambient O₃ concentrations (AOT40 of 13.9 ppm-h) using FACE 13 fumigation (Maggio et al., 2009). However a modeling study has demonstrated that 84% 14 of the variability in the relative feed value in high-yielding alfalfa was due to the 15 variability in mean O₃ concentration from 1998-2002 (Lin et al., 2007). Further, in a 16 managed grassland FACE system, the reduction in total biomass harvest over five years 17 decreased twice as fast in the elevated treatment (AOT40 of 13-59 ppm-h) compared to 18 ambient (AOT40 of 1-20.7 ppm-h). Compared with the ambient control, loss in annual 19 dry matter yield was 23% after 5 year. Further, functional groups were differentially 20 affected, with legumes showing the strongest negative response (Volk et al., 2006). 21 However, a later study by Stampfli and Fuhrer (2010) at the same site suggested that 22 Volk et al. (2006) likely overestimated the effects of O_3 on yield reduction because the 23 overlapping effects of species dynamics caused by heterogeneous initial conditions and a 24 change in management were not considered by these authors. An OTC study conducted 25 with *Trifolium subterraneum* exposed to filtered (<15 ppb), ambient, and 40 ppb above 26 ambient O_3 demonstrated decreases in biomass in the highest O_3 treatment as well as 10-27 20% decreased nutritive quality which was mainly attributed to accelerated senescence 28 (Sanz et al., 2005). A study conducted with Eastern gamagrass and big bluestem in OTCs 29 suggested that big bluestem was not sensitive to O_3 , but gamagrass displayed decreased 30 nutritive quality in the $2 \times$ ambient O₃ treatment, due to higher lignin content and 31 decreased N (Lewis et al., 2006).

9.4.4.3 Summary

1	The detrimental effect of O_3 on crop production has been recognized since the 1960's and
2	a large body of research has subsequently stemmed from those initial findings. Previous
3	O_3 AQCDs have extensively reviewed this body of literature (<u>U.S. EPA, 2006b</u>). Current
4	O_3 concentrations across the U.S. are high enough to cause yield loss for a variety of
5	agricultural crops including, but not limited to, soybean, wheat, potato, watermelon,
6	beans, turnip, onion, lettuce, and tomato. Continued increases in O_3 concentration may
7	further decrease yield in these sensitive crops. Despite the well-documented yield losses
8	due to increasing O_3 concentration, there is still a knowledge gap pertaining to the exact
9	mechanisms of O_3 -induced yield loss. Research has linked increasing O_3 concentration to
10	decreased photosynthetic rates and accelerated senescence, which are related to yield.
11	New research is beginning to consider the mechanism of damage caused by prolonged,
12	lower O_3 concentration (so-called chronic exposure) compared to short, very high O_3
13	concentration (so-called acute exposure). Both types of O ₃ exposure cause damage to
14	agricultural crops, but through very different mechanisms. Historically, most research on
15	the mechanism of O ₃ damage used acute exposure studies. During the last decade, it has
16	become clear that the cellular and biochemical processes involved in the response to
17	acute O3 exposure are not involved in response to chronic O3 exposure, even though both
18	cause yield loss in agriculturally important crops.
19	In addition, recent research has highlighted the effects of O ₃ on crop quality. Increasing
20	O3 concentration decreases nutritive quality of grasses, decreases macro- and micro-
21	nutrient concentrations in fruits and vegetable crops, and decreases cotton fiber quality.
22	These areas of research require further investigation to determine mechanisms and
23	exposure-response relationships.
24	During the previous NAAQS reviews, there were very few studies that estimated O_3
25	impacts on crop yields at large geographical scales. Recent modeling studies found that
26	O3 generally reduced crop yield, but the impacts varied across regions and crop species.
27	For example, the largest O3-induced crop yield losses occurred in high-production areas
28	exposed to high O_3 concentrations, such the Midwest and the Mississippi Valley regions
29	of the U.S. (VanDingenen et al., 2009). Among crop species, the estimated yield loss for
30	wheat and soybean were higher than for rice and maize (VanDingenen et al., 2009).
31	Using satellite air-column observations with direct air-sampling O ₃ data, Fishman et al.
32	(2010) modeled the yield-loss due to O_3 over the continuous tri-state area of Illinois,
33	Iowa and Wisconsin. They determined that O ₃ concentrations significantly reduced
34	soybean yield, which further reinforces previous results from FACE-type experiments
35	and OTC experiments. Evidence is sufficient to conclude that there is a causal

1relationship between O3 exposure and reduced yield and quality of agricultural2crops.

Table 9-4Summary of recent studies of ozone effects on crops (exclusive of
growth and yield).

	•				
Species Facility Location	Exposure Duration	Ozone Exposure ^a (Additional treatment)	Variable(s) measured	Percent (%) change from CF ^b (% change from ambient)	Reference
Alfalfa (<i>Medicago</i> <i>sativa</i> cv. Beaver) Growth chambers	1, 2 or 4 days	3 or 5 h/day 85 ppb (Exposure duration)	Relative feed value	n.s. *high variability among treatment groups (N/A)	<u>Muntifering et</u> <u>al. (2006b</u>)
Bean (Phaseolus vulgaris I. cv Borlotto) OTC, ground- planted Curno, Italy	4 months	Seasonal AOT40: CF = 0.5 ppm-h; Ambient = 4.6 ppm-h (N/A)	Seed lipid, Protein content Fiber content	+28.5 (N/A) +7.88 (N/A) +14.54 (N/A)	<u>Iriti et al.</u> (2009)
Big Blue Stem (<i>Andropogon</i> <i>gerardii</i>) OTC Alabama, U.S.	4 months	12-h avg: CF = 14 ppb; Ambient = 29 ppb; Elevated = 71 ppb (N/A)	Relative feed value	n.s. (n.s.)	<u>Lewis et al.</u> (2006)
<i>Brassica napus</i> Growth chambers Belgium	4 days	CF & 176 ppb for 4 h/day (N/A)	Glucosinolates	-41 (N/A)	<u>Gielen et al.</u> (2006)
Brassica napus cv. Westar Growth chambers Finland	17-26 days	8-h avg: CF & 100 ppb (Bt/non-Bt; herbivory)	VOC emissions	−30.7 (N/A); −34 (N/A)	<u>Himanen et</u> <u>al. (2009b</u>)
Eastern Gamagrass (<i>Tripsacum</i> <i>dactyloides</i>) OTC Alabama, U.S.	4 months	12-h avg: CF = 14 ppb; Ambient = 29 ppb; Elevated = 71 ppb (N/A)	Relative feed value	-17 (-12)	<u>Lewis et al.</u> (2006)
Lettuce (<i>Lactuca sativa</i>) OTC Carcaixent Experimental Station, Spain	30 days	12-h mean: CF = 10.2 ppb; NF = 30.1 ppb; $NF+O_3 = 62.7 \text{ ppb}$ (4 cultivars)	Lipid peroxidation; Root length	+77 (+38) -22 (-14)	<u>Calatayud et</u> <u>al. (2002</u>)
Peanut (<i>Arachis hypogaea</i>) OTC Raleigh, NC; U.S.	З уг	12-h avg: CF = 22 ppb; Ambient = 46 ppb; Elevated = 75 ppb (CO ₂ : 375 ppm; 548 ppm; 730 ppm)	Harvest biomass	-40 (-10)	Booker et al. (2007)
Poa pratensis OTC Braunschweig, Germany	3 yr; 4-5 weeks in the spring	8-h avg: CF+25 = 21.7 ppb; NF+50 = 73.1 ppb (Competition)	Relative feed value	N/A (n.s.; -8)	<u>Bender et al.</u> (2006)

Species Facility Location	Exposure Duration	Ozone Exposure ^a (Additional treatment)	Variable(s) measured	Percent (%) change from CF ^b (% change from ambient)	Reference
Potato (Solanum tuberosum cv. Bintje) OTC Sweden & Finland	2 yr	CF = 10 ppb; Ambient = 25 ppb); Ambient(+) = (36 ppb); Ambient(++) = (47 ppb) (N/A)	[K], [Ca], [Mg], [P], [N] per dry weight of tubers *dose-response regression, report significant positive or negative slope with increasing [O ₃]	[N] [P] [Ca] n.s.; [K] & [Mg] sig + (N/A)	<u>Piikki et al.</u> (2007)
Potato (Solanum tuberosum cv. Indira) Climate chambers Germany	8 weeks	CF = 10 ppb; Ambient = 50 ppb; 2× Ambient = 100 ppb (CO ₂ : 400 ppm & 700 ppm)	Pathogen infestation using percent necrosis	+52 (n.s.)	Plessl et al. (2007)
Soybean OTC Italy	3 уг	AOT40: CF = 0 ppm-h; Ambient = 3.4 ppm-h; Elevated = 9.0 ppm-h (Well-watered & water-stressed)	Daily evapotranspiration	-28 (-14)	<u>Bou Jaoudé</u> <u>et al. (2008a</u>)
Soybean (<i>Glycine max</i> cv. 93B15) SoyFACE Urbana, IL; U.S.	3 yr May-Oct	AOT40: Ambient = 5-22 ppm-h; Elevated = 20-43 ppm-h (CO ₂ : 550 ppm; environmental variability)	Photosynthesis in new leaves,	N/A (n.s.)	<u>Bernacchi et</u> <u>al. (2006</u>)
Soybean (<i>Glycine max</i> cv. 93B15) SoyFACE Urbana, IL; U.S.	4 months	8-h avg: Ambient = 38.5 ppb; Elevated = 52 ppb (Herbivory)	Herbivory defense-related genes	N/A (N/A)	<u>Casteel et al.</u> (2008)
Soybean (<i>Glycine max</i> cv. Essex) OTC, ground- planted Raleigh, NC; U.S.	2 yr	12-h avg: CF = 21 ppb; 1.5x Ambient = 74 ppb (CO ₂ : 370 ppm & 714 ppm)	Post-harvest residue	N/A (-15.46)	Booker et al. (2005)
Soybean (<i>Glycine max</i> cv. Essex) OTCs, 21 L pots Raleigh, NC; U.S.	3 months	12-h avg: CF = 18 ppb); Elevated = 72 ppb) (CO ₂ : 367 & 718)	Water-use efficiency	n.s. (N/A)	Booker et al. (2004b)
Soybean (<i>Glycine max</i>) 10 cultivars) SoyFACE Urbana, IL; U.S.	2 yr	8-h avg (ppb): Ambient = 46.3 & 37.9; Elevated = 82.5 & 61.3 (Cultivar comparisons)	Total antioxidant capacity	N/A (+19)	Betzelberger et al. (2010)
Spring Wheat (<i>Triticum</i> aestivum	7 yr	Seasonal AOT40s ranged from:	Seed protein content;	N/A (Significant negative correlation)	<u>Piikki et al.</u> (2008b)
cv. Minaret; Satu; Drabant; Dragon) OTCs Belgium, Finland, & Sweden		0 to16 ppm-h (N/A)	1,000-seed weight regressed across all experiments	N/A (Significant negative correlation)	

Species Facility Location	Exposure Duration	Ozone Exposure ^a (Additional treatment)	Variable(s) measured	Percent (%) change from CF ^b (% change from ambient)	Reference
Strawberry (<i>Fragaria x</i> <i>ananassa</i> Duch. Cv. Korona & Elsanta) Growth chambers Bonn, Germany	2 months	8-h avg: CF = 0 ppb; Elevated = 78 ppb (N/A)	Total leaf area	−16 (N/A)	<u>Keutgen et al.</u> (2005)
Sweet Potato Growth Chambers Bonn, Germany	4 weeks	8-h avg: CF = 0 ppb; Ambient <40 ppb; Elevated = 255 ppb (N/A)	Tuber weight	-14 (-11.5)	<u>Keutgen et al.</u> (2008)
Tomato (<i>Lycopersicon</i> <i>esculentum</i>) OTC Valencia, Spain	133 days	8- mean: CF = 16.3 ppb; NF = 30.1 ppb; NF(+) = 83.2 ppb (Various cultivars; early & late harvest)	Brix degree	-7.2 (-3.6)	Dalstein and Vas (2005)
Trifolium repens & Trifolium pretense Aspen FACE Rhinelander, WI; U.S.	3 months	3-mo daylight avg: Ambient = 34.8 ppb; 1.2x Ambient = 42.23 ppb (CO ₂ ; 560 ppm)	Lignin; Dry-matter digestibility	N/A (+19.3) N/A (-4.2)	<u>Muntifering et</u> <u>al. (2006a</u>)

^aOzone exposure in ppb unless otherwise noted.

^bCF = Carbon-filtered air.

NF = Non-filtered air.

Table 9-5Modeled effects of ozone on crop yield loss at regional and global
scales

Scale	Index	O ₃ Impacts	Reference
Global	M7a; M12b; AOT40	Reduced by 7.3% to 12.3% for wheat, 5.4% to 15.6% for soybean, 2.8% to 3.7% for rice, and 2.4% to 4.1% for maize in year 2000.	Van Dingenen et al. (2009)
Global	M12b; AOT40	O_3 -induced global yield reductions ranged from 8.5-14% for soybean, 3.9-15% for wheat, and 2.2-5.5% for maize in year 2000. Global crop production losses totaled 79-121 million metric tons, worth \$11-18 billion annually (in U.S. Dollars; 2000).	<u>Avnery et al. (2011a)</u>
U.S.	M7; M12; AOT40	Reduced by 4.1% to 4.4% for wheat, 7.1% to 17.7% for soybean, 2.6% to 3.2% for rice, and 2.2% to 3.6% for maize in year 2000.	Van Dingenen et al. (2009)
U.S.	SUM06	Caused a loss of 53.8 million to 438 million bushels in soybean production, which account for 1.7–14.2% of total U.S. soybean production in 2005	<u>Tong et al. (2007)</u>
East Asia	M7; M12	Reduced the yield of wheat, rice and corn by 1–9% and soybean by 23– 27% in China, Japan and South Korea in 1990	Wang and Mauzerall (2004)

^aM7 is defined as 7-h mean O_3 concentration (ppb).

 $^{\text{b}}\text{M12}$ is defined as 12-h mean O_3 concentration (ppb).

9.4.5 Water Cycling

1	Ozone can affect water use in plants and ecosystems through several mechanisms
2	including damage to stomatal functioning and loss of leaf area. Figure 9-7 provides a
3	simple illustration of potential effects of O_3 exposure on water cycling. Section <u>9.3.2</u>
4	reviewed possible mechanisms for effects of O ₃ exposure on stomatal functioning. This
5	section on water cycling discusses how this alteration of stomatal functioning may affect
6	water use in leaves, whole plants, a planted forest and watersheds

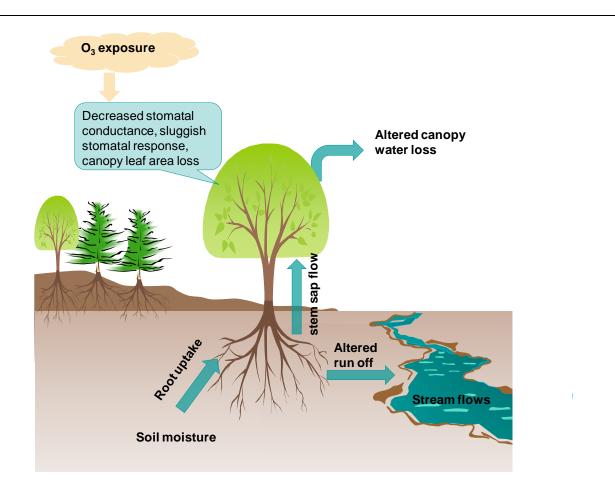


Figure 9-7 The potential effects of ozone exposure on water cycling.

1	In the literature, there is not a clear consensus on the nature of leaf-level stomatal
2	conductance response to O_3 exposure. At the leaf level, O_3 exposure is known to result in
3	stomatal patchiness (Paoletti and Grulke, 2005; Omasa et al., 1987; Ellenson and
4	Amundson, 1982), i.e., the heterogeneous aperture widths of stomata on the leaf surface,
5	and, as a result, the collective response of groups of stomata on leaves and canopies
6	determines larger-scale responses to O ₃ . When measured at steady-state high light
7	conditions, leaf-level stomatal conductance is often found to be reduced when exposed to
8	O_3 . For example, a meta-analysis of 55 studies found that O_3 reduced stomatal
9	conductance by 11% (Wittig et al., 2007). However, these steady-state measurements
10	were generally taken at saturating light conditions and steady-state vapor pressure deficit
11	(VPD). Saturating light and steady-state VPD conditions are not common in the field
12	since many parts of the plant canopy are shaded throughout the day. When studied under
13	varying environmental conditions, many studies have reported incomplete stomatal
14	closure with elevated O ₃ exposure during the day (Mills et al., 2009; Grulke et al., 2007b;
15	Matyssek et al., 1995; Wieser and Havranek, 1995) or at night (Grulke et al., 2004). This

1 may be due to sluggish stomatal response. Sluggish stomatal response, defined as a delay 2 in stomatal response to changing environmental factors relative to controls (Paoletti and 3 Grulke, 2010) has also been documented by several researchers (Grulke et al., 2007c; 4 Matyssek et al., 1995; Pearson and Mansfield, 1993; Wallin and Skärby, 1992; Lee et al., 5 1990; Skarby et al., 1987; Keller and Häsler, 1984; Reich and Lassoie, 1984). Sluggish 6 stomatal response associated with O_3 exposure suggests an uncoupling of the normally 7 tight relationship between carbon assimilation and stomatal conductance as measured 8 under steady-state conditions (Gregg et al., 2006; Paoletti and Grulke, 2005). Several tree 9 and ecosystem models, such as TREGRO, PnET and DLEM, rely on this tight 10 relationship to simulate water and carbon dynamics. The O₃-induced impairment of 11 stomatal control may be more pronounced for plants growing under water stress 12 (Wilkinson and Davies, 2010; Grulke et al., 2007a; Paoletti and Grulke, 2005; Bonn et 13 al., 2004; Kellomaki and Wang, 1997; Tjoelker et al., 1995; Reich and Lassoie, 1984). 14 Since leaf-level stomatal regulation is usually assessed in a steady state rather than as a 15 dynamic response to changing environmental conditions, steady state measurements 16 cannot detect sluggish stomatal response. Because of sluggish stomatal responses, water 17 loss from plants could be greater or reduced under dynamic environmental conditions 18 over days and months. In situations where stomata fail to close under low light or water 19 stressed conditions, water loss may be greater over time. In other situations, it is possible 20 that slugglish stomata may fail to completely open in response to environmental stimuli 21 and result in decreased water loss.

22 In addition to the impacts on stomatal performance, O_3 -induced physiological changes, 23 such as reduced leaf area index and accelerated leaf senescence could alter water use 24 efficiency. It is well established from chamber and field studies that O_3 exposure is 25 correlated with lower foliar retention (Karnosky et al., 2003; Topa et al., 2001; Pell et al., 26 1999; Grulke and Lee, 1997; Karnosky et al., 1996; Miller et al., 1972; Miller et al., 27 1963). However, Lee et al. (2009a) did not find changes in needle area of ponderosa pine 28 and reported that greater canopy conductance followed by water stress under elevated O_3 29 may have been caused by stomatal dysfunction. At the Aspen FACE experiment, stand-30 level water use, as indicated by sap flux per unit ground area, was not significantly 31 affected by elevated O_3 despite a 22% decrease in leaf area index and 20% decrease in 32 basal area (Uddling et al., 2008). The lack of negative effect of elevated O_3 on stand 33 water use may be due to the substantially increased leaf area-specific hydraulic 34 conductance (Uddling et al., 2009). The increased leaf area-specific hydraulic 35 conductance may be caused by the sluggish stomatal response. For example, in the pure 36 aspen stands, the stomatal closure response to increasing vapor pressure deficit was less 37 sensitive and mid-day leaf water potential was more negative under elevated O_3 38 compared to controls. This suggests that O₃ impaired stomatal control over transpiration 39 (Uddling et al., 2009). Another potential factor contributing to the unchanged stand-level

1 water use included the higher proportion of sun leaves in trees under elevated O₃ 2 compared with control trees (Uddling et al., 2008). 3 Elevated O₃ could also affect evapotranspiration by altering tree crown interception of 4 precipitation. Ozone was shown to change branch architectural parameters, and the 5 effects were species-dependent at the Aspen FACE experiment (Rhea et al., 2010). The 6 authors found that there was a significant correlation between canopy architecture 7 parameters and stemflow (the flow of intercepted water down the stem of a tree) for birch 8 but not aspen. 9 It is difficult to scale up physiology measurements from leaves to ecosystems. Thus, the 10 current understanding of how stomatal response at the leaf scale is integrated at the scale 11 of whole forest canopies, and therefore how it influences tree and forest stand water use 12 is limited. Field studies by (McLaughlin et al., 2007a; 2007b) provided valuable insight 13 into the possible consequences of stomatal sluggishness for ecosystem water cycling. 14 McLaughlin et al. (2007a); (2007b) indicated that O₃ increased water use in a mixed 15 deciduous forest in eastern Tennessee. McLaughlin et al. (2007a); (2007b) found that O₃, 16 with daily maximum levels ranging from 69.2 to 82.9 ppb, reduced stem growth by 30-17 50% in the high- O_3 year 2002. The decrease in growth rate was caused in part by 18 amplification of diurnal cycles of water loss and recovery. Peak hourly O₃ exposure 19 increased the rate of water loss through transpiration as indicated by the increased stem 20 sap flow. The authors suggested that a potential mechanism for the increased sap flow 21 could be altered stomatal regulation from O_3 exposure, but this was inferred through sap 22 flow measurements and was not directly measured. The increased canopy water loss 23 resulted in higher water uptake by the trees as reflected in the reduced soil moisture in the 24 rooting zone. The change in tree water use led to further impacts on the hydrological 25 cycle at the landscape level. Increased water use under high O_3 exposure was reported to 26 reduce late-season modeled streamflow in three forested watersheds in eastern Tennessee 27 (McLaughlin et al., 2007b). 28 Felzer et al. (2009) used TEM-Hydro to assess the interactions of O_3 , climate, elevated CO₂ and N limitation on the hydrological cycle in the eastern U.S. They found that 29 30 elevated CO_2 decreased evapotranspiration by 2-4% and increased runoff by 3-7%, as 31 compared to the effects of climate alone. When O₃ damage and N limitation were 32 included, evapotranspiration was reduced by an additional 4-7% and runoff was increased 33 by an additional 6-11% (Felzer et al., 2009). Based upon simulation with INTRAST and 34 LINKAGES, Hanson et al. (2005) found that increasing O₃ concentration by 20 ppb 35 above the current ambient level yields a modest 3% reduction in water use. Those 36 ecological models were generally built on the assumption that O_3 induces stomatal

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closure and have not incorporated possible stomatal sluggishness due to O₃ exposure.

Because of this assumption, results of those models normally found that O_3 reduced water use.

9.4.5.1 Summary

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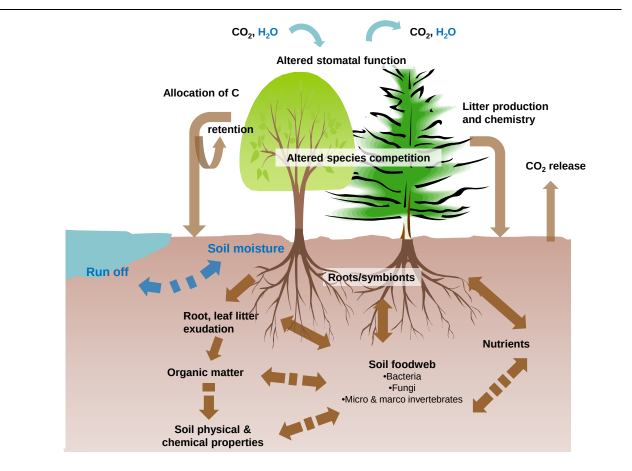
3	Although the evidence was from a limited number of field and modeling studies, findings
4	showed an association between O ₃ exposure and alteration of water use and cycling in
5	vegetation, and at the watershed level. There is not a clear consensus on the nature of
6	leaf-level stomatal conductance response to O ₃ exposure. When measured under steady-
7	state high light conditions, leaf-level stomatal conductance is often found to be reduced
8	when plants are exposed to O ₃ . However, measurements of stomatal conductance under
9	dynamic light and VPD conditions indicate sluggish responses under elevated O ₃
10	exposure, which could potentially lead to increased water loss from vegetation in some
11	situations. Field studies conducted by McLaughlin et al. (2007a); (2007b) suggested that
12	peak hourly O ₃ exposure increased the rate of water loss from several tree species, and
13	led to a reduction in the late-season modeled stream flow in three forested watersheds in
14	eastern Tennessee. Sluggish stomatal responses during O3 exposure was suggested as a
15	possible mechanism for increased water loss during peak O ₃ exposure. Currently, the
16	O ₃ -induced reduction in stomatal aperture is the biological assumption for most process-
17	based models. Because of this assumption, results of those models normally found that
18	O_3 reduced water loss. For example, <u>Felzer et al. (2009</u>) found that O_3 damage and
19	N limitation together reduced evapotranspiration and increased runoff.
20	Although the direction of the response differed among studies, the evidence is sufficient

Although the direction of the response differed among studies, the evidence is sufficient to conclude that there is likely to be a causal relationship between O_3 exposure and the alteration of ecosystem water cycling.

9.4.6 Below-Ground Processes

23	Above-ground and below-ground processes are tightly interconnected. Because roots and
24	soil organisms are not exposed directly to O_3 , below-ground processes are affected by O_3
25	through alterations in the quality and quantity of C supply from photosynthates and
26	litterfall (Andersen, 2003). Ozone can decrease leaf C uptake by reducing photosynthesis
27	(Section 9.3). Ozone can also increase metabolic costs by stimulating the production of
28	chemical compounds for defense and repair processes, and by increasing the synthesis of
29	antioxidants to neutralize free radicals (see Section 9.3), both of which increase the
30	allocation of carbon for above-ground processes. Therefore, O ₃ could significantly reduce

1	the amount of C available for allocation to below-ground by decreasing C uptake while
2	increasing C consumption of above-ground processes (Andersen, 2003).
3	Since the 2006 O_3 AQCD, there is additional evidence for O_3 effects on below-ground
4	processes. Ozone has been found to alter root growth, soil food web structure,
5	decomposer activities, C turnover, water cycling and nutrient flow (Figure 9-8). Ozone
6	effects on root development and root biomass production and soil food web structure are
7	reviewed in Section $9.4.3.1$ and Section $9.4.9.2$, respectively. The focus in this section is
8	on the response of litter input, decomposer activities, soil respiration, soil C formation
9	and nutrient cycling.



Note: Arrows denote C flux pathways that are affected by ozone. Dashed lines indicate where the impact of ozone is suspected but unknown.

Source: Modified from Andersen (2003).

Figure 9-8 Conceptual diagram showing where ozone alters C, water and nutrient flow in a tree-soil system, including transfer between biotic and abiotic components below ground that influence soil physical and chemical properties.

9.4.6.1 Litter Carbon Chemistry, Litter Nutrient and Their Ecosystem Budgets

Consistent with previous findings, recent studies show that, although the responses are often species-dependent, O_3 tends to alter litter chemistry (U.S. EPA, 2006b). Alterations in chemical parameters, such as changes in C chemistry and nutrient concentrations, were observed in both leaf and root litter (Table 9-6).

5 At the Aspen FACE site, several studies investigated litter chemistry changes (Parsons et 6 al., 2008; Johnson and Pregitzer, 2007; Chapman et al., 2005; Liu et al., 2005). In both 7 aspen and birch leaf litter, elevated O_3 increased the concentrations of soluble sugars, 8 soluble phenolics and condensed tannins (Parsons et al., 2008; Liu et al., 2005). 9 Compared to other treatments, aspen litter under elevated O₃ had the highest fiber 10 concentration, with the lowest concentration associated with the birch litter under the 11 same conditions (Parsons et al., 2008). Chapman et al. (2005) measured chemical 12 changes in fine root litter and found that elevated O₃ decreased lignin concentration. 13 O_3 -induced chemistry changes were also reported from other experimental sites. Results 14 from an OTC study in Finland suggested that elevated O₃ increased the concentration of 15 acid-soluble lignin, but had no significant impact on other chemicals such as total sugars, 16 hemicelluloses, cellulose or total lignin in the litter of silver birch (Kasurinen et al., 17 2006). Results from the free air canopy O_3 exposure experiment at Kranzberg Forest 18 showed that O₃ increased starch concentrations but had no impact on cellulose and lignin 19 in beech and spruce leaf litter (Aneja et al., 2007). The effect of O_3 on three antioxidants 20 (ascorbate, glutathione and α -tocopherol) in fine roots of beech was also assessed at 21 Kranzberg Forest. The results indicated that O_3 had no significant effect on α -tocopherol 22 and ascorbate concentrations, but decreased glutathione concentrations in fine roots 23 (Haberer et al., 2008). In addition to changing C chemistry, O₃ also altered nutrient 24 concentrations in green leaves and litter (Table 9-6).

25 The combined effects of O₃ on biomass productivity and chemistry changes may alter 26 C chemicals and nutrient contents at the canopy or stand level. For example, although O_3 27 had different impacts on their concentrations, annual fluxes of C chemicals (soluble 28 sugar, soluble phenolics, condensed tannins, lipid and hemicelluloses), macro nutrients 29 (N, P, K and S) and micro nutrients (Mg, B, Cu and Zn) to soil were all reduced due to 30 lower litter biomass productivity at Aspen FACE (Liu et al., 2007; Liu et al., 2005).. In a 31 2-year growth chamber experiment in Germany, N content of a spruce canopy in a mixed 32 culture and Ca content of a beech canopy in a monoculture was increased due to elevated 33 O_3 , although leaf production was not significantly altered by O_3 (Rodenkirchen et al., 34 2009).

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Study Site	Species	O ₃ Concentration	Response	Reference
Suonenjoki Research Station, Finland	Silver birch	Ambient: 10-60 ppb Elevated: 2x ambient	Decreased the concentration of P, Mn, Zn and B in leaf litter	<u>Kasurinen et al.</u> (2006)
Aspen FACE	Aspen and birch	Ambient: 50-60 ppb Elevated: 1.5x ambient	Decreased the concentrations of P, S, Ca and Zn, but had no impact on the concentrations of N, K, Mg, Mn, B and Cu in leaf litter.	<u>Liu et al. (2007)a</u>)
Aspen FACE	Birch	Ambient: 50-60 ppb Elevated: 1.5× ambient	Increase N concentration in birch litter	<u>Parsons et al.</u> (2008)
Kranzberg Forest, Germany	Beech and spruce	Ambient: 9-41 ppb Elevated: 2x ambient	Increased N concentration in beach leaf, but not in spruce needle	<u>Kozovits et al.</u> (2005)
Kranzberg Forest, Germany	Beech and spruce	Ambient: 9-41 ppb Elevated: 2× ambient	 (1) Had no significant effects on spruce needle chemistry; (2) increased Ca concentration in beech leaves in monoculture, but had no impacts on other nutrients 	
Salerno, Italy	Holm oak	Non-filtered OTC: 29 ppb Filtered OTC: 17ppb	O_3 had no significant impacts on litter C, N, lignin and cellulose concentrations	<u>Baldantoni et al.</u> (2011)
Kuopio University Research Garden, Finland	Red Clover	Ambient: 25.7 ppb Elevated: 1.5x ambient	Increased the total phenolic content of leaves and had minor effects on the concentrations of individual phenolic compounds	<u>Saviranta et al.</u> (2010)

Table 9-6	The effect of elevated ozone on leaf/litter nutrient concentrations.

9.4.6.2 Decomposer Metabolism and Litter Decomposition

1	The above- and below-ground physiological changes caused by O_3 exposure cascade
2	through the ecosystem and affect soil food webs. In the 2006 O_3 AQCD, there were very
3	few studies on the effect of O_3 on the structure and function of soil food webs, except two
4	studies conducted by Larson et al. (2002) and Phillips et al. (2002). Since the last O_3
5	AQCD, new studies have provided more information on how O ₃ affects the metabolism
6	of soil microbes and soil fauna.
7	Chung et al. (2006) found that the activity of the cellulose-degrading enzyme
8	1,4- β -glucosidase was reduced by 25% under elevated O ₃ at Aspen FACE. The decrease
9	in cellulose-degrading enzymatic activity was associated with the lower cellulose
10	availability under elevated O_3 (Chung et al., 2006). However, a later study at the same
11	site, which was conducted in the 10th year of the experiment, found that O ₃ had no
12	impact on cellulolytic activity in soil (Edwards and Zak, 2011). In a lysimeter study of
13	beech trees (Fagus sylvatica) in Germany, soil enzyme activity was found to be
14	suppressed by O ₃ exposure (Esperschutz et al., 2009; Pritsch et al., 2009). Except for

xylosidase, enzyme activities involved in plant cell wall degradation (cellobiohydrolase,
beta-glucosidase and glucuronidase) were decreased in rhizosphere soil samples under
elevated O_3 (2 × ambient level) (<u>Pritsch et al., 2009</u>). Similarly, <u>Chen et al. (2009</u>) found
O ₃ exposure, with a 3-month AOT40 of 21-44 ppm-h, decreased the microbial metabolic
capability in the rhizosphere and bulk soil of wheat, although the observed reduction in
bulk soil was not significant.

7 Ozone-induced change in soil organisms' activities could affect litter decomposition 8 rates. Results of recent studies indicated that O₃ slightly reduced or had no impacts on 9 litter decomposition (Liu et al., 2009b; Parsons et al., 2008; Kasurinen et al., 2006) 10 (Baldantoni et al., 2011). The responses varied among species, sites and exposure length. 11 Parsons et al. (2008) collected litter from aspen and birch seedlings at Aspen FACE site, 12 and conducted a 23-month field litter incubation starting in 1999. They found that 13 elevated O_3 had different impacts on the decomposition of aspen and birch litter. Elevated 14 O₃ was found to reduce aspen litter decomposition. However, O₃ accelerated birch litter 15 decomposition under ambient CO_2 , but reduced it under elevated CO_2 (Parsons et al., 16 2008). Liu et al. (2009b) conducted another litter decomposition study at Aspen FACE 17 from 2003 to 2006, when stand leaf area index (LAI) reached its maximum. During the 18 935-day field incubation, elevated O_3 was shown to reduce litter mass loss in the first 19 year, but not in the second year. They suggested that higher initial tannin and phenolic 20 concentrations under elevated O₃ reduced microbial activity in the first year (Liu et al., 21 2009b). In an OTC experiment, Kasurinen et al. (2006) collected silver birch leaf litter 22 from three consecutive growing seasons and conducted three separate litter-bag 23 incubation experiments. Litter decomposition was not affected by O_3 exposure in the first 24 two incubations, but a slower decomposition rate was found in the third incubation. Their 25 principle component analysis indicated that the litter chemistry changes caused by O_3 26 (decreased Mn, P, B and increased C:N) might be partially responsible for the decreased 27 mass loss of their third incubation. In another OTC experiment, Baldantoni et al. (2011) 28 found that O_3 significantly reduced leaf litter decomposition of *Quercus ilex* L, although 29 litter C, N, lignin and cellulose concentrations were not altered by O₃ exposure.

9.4.6.3 Soil Respiration and Carbon Formation

30	Ozone could reduce the availability of photosynthates for export to roots, and thus,
31	indirectly increase root mortality and turnover rates. Ozone has also been shown to
32	reduce above-ground litter productivity and alter litter chemistry, which would affect the
33	quality and quantity of the C supply to soil organisms (Section 9.4.6.1). The complex
34	interactions among those changes make it difficult to predict the response of soil
35	C cycling under elevated O_3 . The 2006 O_3 AQCD concluded that O_3 had no consistent

1	impact on soil respiration (U.S. EPA, 2006b). Ozone could increase or decrease soil
2	respiration, depending on the approach and timing of the measurements. Ozone may also
3	alter soil C formation. However, very few experiments directly measured changes in soil
4	organic matter content under O_3 fumigation (U.S. EPA, 2006b). Recent studies on soil
5	respiration and soil C content also found mixed responses. Most importantly, recent
6	results from long-term fumigation experiments, such as the Aspen FACE experiment,
7	suggest that ecosystem response to O_3 exposure can change over time. Observations
8	made during the late exposure years can be inconsistent with those during the early years,
9	highlighting the need for caution when assessing O ₃ effects based on short-term studies
10	(<u>Table 9-7</u>).

Table 9-7The temporal variation of ecosystem responses to ozone exposure
at Aspen FACE site

Endpoint	Period of Measurement	Response	Reference
Litter decomposition	1999-2001	O_3 reduced aspen litter decomposition. However, O_3 accelerated birch litter decomposition under ambient CO_2 , but reduced it under elevated CO_2	<u>Parsons et al. (2008</u>)
	2003-2006	O_3 reduced litter mass loss in the first year, but not in the second year.	<u>Liu et al. (2009b</u>)
Fine root production	1999	O_3 had no significant impact on fine root biomass	King et al. (2001)
	2002, 2005	O_3 increased fine root biomass	Pregitzer et al. (2008)
Soil respiration	1998-1999	Soil respiration under $+CO_2+O_3$ treatment was lower than that under $+CO_2$ treatment	<u>King et al. (2001)</u>
	2003-2007	Soil respiration under $+CO_2+O_3$ treatment was 5-25% higher than under elevated CO_2 treatment.	Pregitzer et al. (2008); Pregitzer et al. (2006)
Soil C formation	1998-2001	O_3 reduced the formation rates of total soil C by 51% and acid-insoluble soil C by 48%	<u>Loya et al. (2003</u>)
	2004-2008	No significant effect of O_3 on the new C formed under elevated CO_2	<u>Talhelm et al. (2009</u>)

Soil Respiration

11	Ozone has shown inconsistent impacts on soil respiration. A sun-lit
12	controlled-environment chamber study found that O_3 had no significant effects on soil
13	respiration, fine root biomass or any of the soil organisms in a reconstructed ponderosa
14	pine/soil-litter system (Tingey et al., 2006). In an adult European beech/Norway spruce
15	forest at Kranzberg Forest, the free air O ₃ fumigation (AOT40 of 10.2-117 ppm-h)
16	increased soil respiration under both beech and spruce during a humid year (Nikolova et
17	al., 2010). The increased soil respiration under beech has been accompanied by the

1	increase in fine root biomass and ectomycorrhizal fungi diversity and turnover (Grebenc
2	and Kraigher, 2007). The stimulating effect on soil respiration disappeared under spruce
3	in a dry year, which was associated with a decrease in fine root production in spruce
4	under drought. This finding suggested that drought was a more dominant stress than O_3
5	for spruce (<u>Nikolova et al., 2010</u>). <u>Andersen et al. (2010</u>) labeled the canopies of
6	European beech and Norway spruce with CO_2 depleted in ¹³ C at the same site. They did
7	not observe any significant changes in soil respiration for either species.
8	The nearly 10 year long studies at Aspen FACE indicated that the response of soil
9	respiration to O_3 interacted with CO_2 exposure and varied temporally (<u>Table 9-7</u>)
10	(Pregitzer et al., 2008; Pregitzer et al., 2006; King et al., 2001). Ozone treatment alone
11	generally had the lowest mean soil respiration rates, although those differences between
12	control and elevated O ₃ were usually not significant. However, soil respiration rates were
13	different with O ₃ alone and when acting in combination with elevated CO ₂ . In the first
14	five years (1998-2002), soil respiration under $+CO_2+O_3$ treatment was similar to that
15	under control and lower than that under +CO ₂ treatment (Pregitzer et al., 2006; King et
16	<u>al., 2001</u>). Since 2003, $+CO_2+O_3$ treatment started to show the greatest impact on soil
17	respiration. Compared to elevated CO_2 , soil respiration rate under $+CO_2+O_3$ treatment
18	was 15-25% higher from 2003-2004, and 5-10% higher from 2005-2007 (Pregitzer et al.,
19	2008; Pregitzer et al., 2006). Soil respiration was highly correlated with the biomass of
20	roots with diameters of <2 mm and <1 mm, across plant community and atmospheric
21	treatments. The authors suggested that the increase in soil respiration rate may be due to
22	+CO ₂ +O ₃ increased fine root (<1.0 mm) biomass production (<u>Pregitzer et al., 2008</u>).
23	Changes in leaf chemistry and productivity due to O_3 exposure have been shown to affect
24	herbivore growth and abundance (See Section <u>9.4.9.1</u>). Canopy insects could affect soil
25	carbon and nutrient cycling through frass deposition, or altering chemistry and quantity
26	of litter input to the forest floor. A study at the Aspen FACE found that although elevated
27	O ₃ affected the chemistry of frass and greenfall, these changes had small impact on
28	microbial respiration and no effect on nitrogen leaching (Hillstrom et al., 2010a).
29	However, respiratory carbon loss and nitrate immobilization were nearly double in
30	microcosms receiving herbivore inputs than those receiving no herbivore inputs
31	(Hillstrom et al., 2010a).

Soil Carbon Formation

32	Ozone-induced reductions in plant growth can result in reduced C input to soil and
33	therefore soil C content (Andersen, 2003). The simulations of most ecosystem models
34	support this prediction (Ren et al., 2007b; Zhang et al., 2007a; Felzer et al., 2004).
35	However, very few studies have directly measured soil C dynamics under elevated O ₃ .

1	After the first four years of fumigation (from 1998 to 2001) at the Aspen FACE site,
2	Loya et al. (2003) found that forest stands exposed to both elevated O_3 and CO_2
3	accumulated 51% less total soil C, and 48% less acid-insoluble soil C compared to stands
4	exposed only to elevated CO ₂ . Soil organic carbon (SOC) was continuously monitored at
5	the Aspen FACE site, and the later data showed that the initial reduction in new
6	C formation (soil C derived from plant litter since the start of the experiment) by O_3
7	under elevated CO ₂ is only a temporary effect (<u>Table 9-7</u>) (<u>Talhelm et al., 2009</u>). The
8	amount of new soil C in the elevated CO_2 and the combined elevated CO_2 and O_3
9	treatments has converged since 2002. There was no significant effect of O_3 on the new C
10	formed under elevated CO_2 over the last four years of the study (2004-2008). <u>Talhelm et</u>
11	al. (2009) suggested the observed reduction in the early years of the experiment might be
12	driven by a suppression of C allocated to fine root biomass. During the early exposure
13	years, O_3 had no significant impact on fine root production (King et al., 2001). However,
14	the effect of O_3 on fine root biomass was observed later in the experiment. Ozone
15	increased fine root production and the highest fine root biomass was observed under the
16	combined elevated CO_2 and O_3 treatment in the late exposure years (<u>Table 9-7</u>) (<u>Pregitzer</u>)
17	et al., 2006). This increase in fine root production was due to changes in community
18	composition, such as better survival of an O ₃ -tolerant aspen genotype, birch and maple,
19	rather than changes in C allocation at the individual tree level (Pregitzer et al., 2008; Zak
20	<u>et al., 2007</u>).

9.4.6.4 Nutrient Cycling

21	Ozone can affect nutrient cycling by changing nutrient release from litter, nutrient uptake
22	by plants, and soil microbial activity. Nitrogen is the limiting nutrient for most temperate
23	ecosystems, and several studies examined N dynamics under elevated O ₃ . Nutrient
24	mineralization from decomposing organic matter is important for sustaining ecosystem
25	production. Holmes et al. (2006) found that elevated O3 decreased gross N mineralization
26	at the Aspen FACE site, indicating that O_3 may reduce N availability. Other N cycling
27	processes, such as NH_4^+ immobilization, gross nitrification, microbial biomass N and soil
28	organic N, were not affected by elevated O ₃ (Holmes et al., 2006). Similarly, Kanerva et
29	al. (2006) found total N, NO ₃ -, microbial biomass N, potential nitrification and
30	denitrification in their meadow mesocosms were not affected by elevated O_3 (40-50 ppb).
31	Ozone was found to decrease soil mineral N content at SoyFACE, which was likely
32	caused by a reduction in plant material input and increased denitrification (Pujol Pereira
33	et al., 2011). Ozone also showed small impact on other micro and macro nutrients. Liu et
34	al. (2007)a) assessed N, P, K, S, Ca, Mg, Mn, B, Zn and Cu release dynamics at Aspen
35	FACE, and they found that O_3 had no effects on most nutrients, except to decrease N and

1	Ca release from litter. These studies reviewed above suggest that soil N cycling processes
2	are not affected or slightly reduced by O ₃ exposure. However, in a lysimeter study with
3	young beech trees, Stoelken et al. (2010) found that elevated O_3 stimulated N release
4	from litter which was largely attributed to an enhanced mobilization of inert nitrogen
5	fraction.
6	Using the SImple NItrogen Cycle model (SINIC), Hong et al. (2006) evaluated the
7	impacts of O ₃ exposure on soil N dynamics and streamflow nitrate flux. The detrimental
8	effect of O_3 on plant growth was found to reduce plant uptake of N and therefore increase
9	nitrate leaching. Their model simulation indicated that ambient O ₃ exposure increased the
10	mean annual stream flow nitrate export by 12% (0.042 g N/m ² /year) at the Hubbard
11	Brook Experimental Watershed from 1964-1994 (Hong et al., 2006).

9.4.6.5 Dissolved Organic Carbon and Biogenic Trace Gases Emission

- 12The O3-induced changes in plant growth, C and N fluxes to soil and microbial13metabolism can alter other biogeochemical cycling processes, such as soil dissolved14organic carbon (DOC) turnover and trace gases emission.
- 15 Jones et al. (2009) collected fen cores from two peatlands in North Wales, UK and 16 exposed them to one of four levels of O₃ (AOT40 of 0, 3.69, 5.87 and 13.80 ppm-h for 17 41 days). They found the concentration of porewater DOC in fen cores was significantly 18 decreased by increased O₃ exposure. A reduction of the low molecular weight fraction of 19 DOC was concurrent with the observed decrease in DOC concentration. Their results 20 suggested that O₃ damage to overlying vegetation may decrease utilizable C flux to soil. 21 Microbes, therefore, have to use labile C in the soil to maintain their metabolism, which, 22 the authors hypothesized, leads to a decreased DOC concentration with a shift of the 23 DOC composition to more aromatic, higher molecular weight organic compounds.
- 24 Several studies since the 2006 O_3 AQCD have examined the impacts of O_3 on nitrous 25 oxide (N_2O) and methane (CH_4) emission. Kanerva et al. (2007) measured the fluxes of 26 N_2O and CH_4 in meadow mesocosms, which were exposed to elevated CO_2 and O_3 in 27 OTCs in south-western Finland. They found that the daily N₂O fluxes were decreased in 28 the NF+O₃ (non-filtered air + elevated O_3 , 40-50 ppb) after three seasons of exposure. 29 Elevated O_3 alone or combined with CO_2 did not have any significant effect on the daily 30 fluxes of CH_4 (Kanerva et al., 2007). In another study conducted in central Finland, the 31 4 year open air O_3 fumigation (AOT40 of 20.8-35.5 ppm-h for growing season) slightly 32 increased potential CH_4 oxidation by 15% in the peatland microcosms, but did not affect 33 the rate of potential CH_4 production or net CH_4 emissions, which is the net result of the

1	potential CH ₄ production and oxidation (Morsky et al., 2008). However, several studies
2	found that O_3 could significantly reduce CH_4 emission. Toet et al. (2011) exposed
3	peatland mesocosms to O_3 in OTCs for two years, and found that CH_4 emissions were
4	significantly reduced by about 25% during midsummer periods of both years. In an OTC
5	study of rice paddy, Zheng et al. (2011) found that the daily mean CH_4 emissions were
6	significantly lower under elevated O_3 treatments than those in charcoal-filtered air and
7	nonfiltered air treatments. They found that the seasonal mean CH ₄ emissions were
8	negatively related with AOT40, but positively related to the relative rice yield,
9	aboveground biomass and underground biomass.

9.4.6.6 Summary

10Since the 2006 O3 AQCD, more evidence has shown that although the responses are11often site specific, O3 altered the quality and quantity of litter input to soil, microbial12community composition, and C and nutrient cycling. Biogeochemical cycling of below-13ground processes is fueled by C input from plants. Studies at the leaf and plant level have14provided biologically plausible mechanisms, such as reduced photosynthetic rates,15increased metabolic cost, and reduced root C allocation for the association of O3 exposure16and the alteration of below-ground processes.

17 Results from Aspen FACE and other experimental studies consistently found that O₃ 18 reduced litter production and altered C chemistry, such as soluble sugars, soluble 19 phenolics, condensed tannins, lignin, and macro/micro nutrient concentration in litter 20 (Parsons et al., 2008; Kasurinen et al., 2006; Liu et al., 2005). Under elevated O₃, the 21 changes in substrate quality and quantity could alter microbial metabolism and therefore 22 soil C and nutrient cycling. Several studies indicated that O₃ suppressed soil enzyme 23 activities (Pritsch et al., 2009; Chung et al., 2006). However, the impact of O₃ on litter 24 decomposition was inconsistent and varied among species, sites and exposure length. 25 Similarly, O₃ had inconsistent impacts on dynamics of micro and macro nutrients.

- 26Studies from the Aspen FACE experiment suggested that the response of below-ground27C cycle to O3 exposure, such as litter decomposition, soil respiration and soil C content,28changed over time. For example, in the early part of the experiment (1998-2003), O3 had29no impact on soil respiration but reduced the formation rates of total soil C under30elevated CO2. However, after 10-11 years of exposure, O3 was found to increase soil31respiration but have no significant impact on soil C formation under elevated CO2.
- 32The evidence is sufficient to infer that there is a causal relationship between O333exposure and the alteration of below-ground biogeochemical cycles.

9.4.7 Community Composition

The effects of O_3 on species competition (AX9.3.3.4) and community composition
(AX9.6.4) were summarized in the 2006 O ₃ AQCD. Plant species differ in their
sensitivity to O_3 . Further, different genotypes of a given species also vary in their
sensitivity. This differential sensitivity could change the competitive interactions that
lead to loss in O ₃ sensitive species or genotypes. In addition, O ₃ exposure has been found
to alter reproductive processes in plants (See Section $9.4.3.3$). Changes in reproductive
success could lead to changes in species composition. However, since ecosystem-level
responses result from the interaction of organisms with one another and with their
physical environment, it takes longer for a change to develop to a level of prominence at
which it can be identified and measured. A shift in community composition in forest and
grassland ecosystems noted in the 2006 O_3 AQCD has continued to be observed from
experimental and gradient studies. Additionally, research since the last review has shown
that O ₃ can alter community composition and diversity of soil microbial communities.

9.4.7.1 Forest

14	In the San Bernardino Mountains in southern California, O ₃ pollution caused a significant
15	decline in ponderosa pine (Pinus ponderosa) and Jeffrey pine (Pinus jeffreyi) (U.S. EPA,
16	2006b). Pine trees in the young mature age class group exhibited higher mortality rates
17	compared with mature trees at a site with severe O ₃ visible foliar injury. The vulnerability
18	of young mature pines was most likely caused by the fact that trees in this age class were
19	emerging into the canopy, where higher O_3 concentrations were encountered (McBride
20	and Laven, 1999). Because of the loss of O_3 -sensitive pines, mixed forests of ponderosa
21	pine, Jeffery Pine and white fir (Abies concolor) shifted to predominantly white fir
22	(Miller, 1973). Ozone may have indirectly caused the decline in understory diversity in
23	coniferous forests in the San Bernardino Mountains through an increase in pine litterfall.
24	This increase in litterfall from O_3 exposure results in an understory layer that may
25	prohibit the establishment of native herbs, but not the exotic annual Galium aparine
26	(<u>Allen et al., 2007</u>).
27	Ozone damage to conifer forests has also been observed in several other regions. In the
28	Valley of Mexico, a widespread mortality of sacred fir (Abies religiosa) was observed in
29	the heavily polluted area of the Desierto de los Leones National Park in the early 1980s
30	(de Lourdes de Bauer and Hernandez-Tejeda, 2007; Fenn et al., 2002). Ozone damage
31	was widely believed to be an important causal factor in the dramatic decline of sacred fir.
32	In alpine regions of southern France and the Carpathians Mountains, O3 was also
33	considered as the major cause of the observed decline in cembran pine (Pinus cembra)

(Wieser et al., 2006). However, many environmental factors such as light, temperature, nutrient and soil moisture, and climate extremes such as unusual dry and wet periods could interact with O_3 and alter the response of forest to O_3 exposure. For those pollution gradient studies, several confounding factors, such as drought, insect outbreak and forest management, may also contribute to or even be the dominant factors causing the mortality of trees (de Lourdes de Bauer and Hernandez-Tejeda, 2007; Wieser et al., 2006).

8 Recent evidence from long-term free O₃ fumigation experiments provided additional 9 support for the potential impacts of O₃ on species competition and community 10 composition changes in forest ecosystems. At the Aspen FACE site, community 11 composition at both the genetic and species levels was altered after seven years of 12 fumigation with O_3 (Kubiske et al., 2007). In the pure aspen community, O_3 fumigation 13 reduced growth and increased mortality of sensitive clone 259, while the O_3 tolerant 14 clone 8L emerged as the dominant clone. Growth of clone 8L was even greater under 15 elevated O_3 compared to controls, probably due to O_3 alleviated competitive pressure on 16 clone 8L by reducing growth of other clones. In the mixed aspen-birch and aspen-maple 17 communities, O₃ reduced the competitive capacity of aspen compared to birch and maple 18 (Kubiske et al., 2007). In a phytotron study, O₃ fumigation reduced growth of beech but 19 not spruce in mixed culture, suggesting a higher susceptibility of beech to O₃ under 20 interspecific competition (Kozovits et al., 2005).

9.4.7.2 Grassland and Agricultural Land

21	The response of managed pasture, often cultivated as a mixture of grasses and clover, to
22	O ₃ pollution has been studied for many years. The tendency for O ₃ -exposure to shift the
23	biomass of grass-legume mixtures in favor of grass species, reported in the previous O_3
24	AQCD has been generally confirmed by recent studies. In a mesocosm study, Trifolium
25	repens and Lolium perenne mixtures were exposed to an episodic rural O3 regime within
26	solardomes for 12 weeks. T. repens showed significant changes in biomass but not L.
27	perenne, and the proportion of T. repens decreased in O3-exposed mixtures compared to
28	the control (Hayes et al., 2009). The changes in community composition of grass-legume-
29	forb mixtures were also observed at the Le Mouret FACE experiment, Switzerland.
30	During the 5-year O ₃ fumigation (AOT40 of 13.3-59.5 ppm-h), the dominance of
31	legumes in fumigated plots declined more quickly than those in the control plots (Volk et
32	al., 2006). However, Stampfli and Fuhrer (2010) reanalyzed the species and soil data and
33	suggested that Volk et al. (2006) overestimated the O ₃ effect. Stampfli and Fuhrer (2010)
34	found that the difference in the species dynamics between control and O_3 treatment was
35	more caused by heterogeneous initial conditions than O3 exposure. Several studies also

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1	suggested that mature/species-rich ecosystems were more resilient to O_3 exposure. At
2	another FACE experiment, located at Alp Flix, Switzerland, O3 fumigation (AOT40 of
3	15.2-64.9 ppm-h) showed no significant impact on community composition of this
4	species-rich pasture (Bassin et al., 2007b). Although most studies demonstrated an
5	increase in grass: forb ratio with O ₃ exposure (Hayes et al., 2009; U.S. EPA, 2006b), a
6	study on a simulated upland grassland community showed that O3 reduced the grass:forb
7	ratio (Hayes et al., 2010) which may be due to the grass species in this community. The
8	grass species studied by Hayes et al. (2010), Anthoxanthum odoratum, was more
9	sensitive to O_3 than other grass species such as <i>L. perenne</i> (<u>Hayes et al., 2009</u>). <u>Pfleeger</u>
10	et al. (2010) collected seed bank soil from an agricultural field and examined how the
11	plant community responded over several generations to elevated O ₃ exposures. Sixty
12	plant species from 22 families emerged in the chambers over their four year study.
13	Overall, they found that O_3 appeared to have small impacts on seed germination and only
14	a minor effect on species richness of pioneer plant communities.
15	Several review papers have discussed the physiological and ecological characteristics of
16	O ₃ -sensitive herbaceous plants. <u>Hayes et al. (2007</u>) assessed species traits associated with
17	O_3 sensitivity by the changes in biomass caused by O_3 exposure. Plants of the therophyte
18	(e.g., annual) life form were particularly sensitive to O ₃ . Species with higher mature leaf
19	N concentration tended to be more sensitive than those with lower leaf N concentration.
20	Plants growing under high oxidative stress environments, such as high light or high
21	saline, were more sensitive to O_3 . Using the same dataset from <u>Hayes et al. (2007</u>), <u>Mills</u>
22	et al. $(2007b)$ identified the O ₃ sensitive communities. They found that the largest number
23	of these O ₃ sensitive communities were associated with grassland ecosystems. Among
24	grassland ecosystems, alpine grassland, sub-alpine grassland, woodland fringe, and dry
25	grassland were identified as the most sensitive communities.

9.4.7.3 Microbes

26	Several methods have been used to study microbial composition changes associated with
27	elevated O ₃ . Phospholipid fatty acid (PLFA) analysis is widely used to determine whether
28	O3 elicits an overall effect on microbial community composition. However, since PLFA
29	markers cover a broad range of different fungi, resolution of this method may be not fine
30	enough to detect small changes in the composition of fungal communities. Methods, such
31	as microscopic analyses and polymerase chain reaction-denaturing gradient gel
32	electrophoresis (PCR-DGGE), have better resolution to specifically analyze the fungal
33	community composition. The resolution differences among those methods needs to be
34	considered when assessing the O_3 impact on microbial community composition.

1	Kanerva et al. (2008) found that elevated O_3 (40-50 ppb) decreased total, bacterial,
2	actinobacterial and fungal PLFA biomass values as well as fungal:bacterial PLFA
3	biomass ratio in their meadow mesocosms in south-western Finland. The relative
4	proportions of individual PLFAs between the control and elevated O ₃ treatments were
5	significantly different, suggesting that O_3 modified the structure of the microbial
6	community. Morsky et al. (2008) exposed boreal peatland microcosms to elevated O_3 ,
7	with growing season AOT40 of 20.8-35.3 ppm-h, in an open-air O ₃ exposure field in
8	Central Finland. They also found that microbial composition was altered after three
9	growing seasons with O ₃ fumigation, as measured by PLFA. Ozone tended to increase
10	the presence of Gram-positive bacteria and the biomass of fungi in the peatland
11	microcosms. Ozone also resulted in higher microbial biomass, which co-occurred with
12	the increases in concentrations of organic acids and leaf density of sedges (Morsky et al.,
13	2008). In a lysimeter experiment in Germany, O ₃ was found to alter the PLFA profiles in
14	the upper 0-20 cm rhizosphere soil of European beech. Elevated O ₃ reduced bacterial
15	abundance but had no detectable effect on fungal abundance (Pritsch et al., 2009). Using
16	microscopic analyses, Kasurinen et al. (2005) found that elevated O ₃ , with 5 or 6 months
17	of AOT40 of 20.6-30.9 ppm-h, decreased the proportions of black and liver-brown
18	mycorrhizas and increased that of light brown/orange mycorrhizas. In an herbaceous
19	plant study, SSCP (single-strand conformation polymorphism) profiles indicated that O_3
20	stress (about 75 ppb) had a very small effect on the structural diversity of the bacterial
21	community in rhizospheres (Dohrmann and Tebbe, 2005). At the Aspen FACE site, O_3
22	had no significant effect on fungal relative abundance, as indicated by PLFA profile.
23	However, elevated O_3 altered fungal community composition, according to the
24	identification of 39 fungal taxonomic units from soil using polymerase chain reaction-
25	denaturing gradient gel electrophoresis (PCR-DGGE) (Chung et al., 2006). In another
26	study at Aspen FACE, phylogenetic analysis suggested that O ₃ exposure altered the
27	agaricomycete community. The ectomycorrhizal communities developing under elevated
28	O ₃ had higher proportions of <i>Cortinarius</i> and <i>Inocybe</i> species, and lower proportions of
29	Laccaria and Tomentella (Edwards and Zak, 2011). Ozone was found to change
30	microbial community composition in an agricultural system. Chen et al. (2010b) found
31	elevated O ₃ (100-150 ppb) had significant effects on soil microbial composition
32	expressed as PLFA percentage in a rice paddy in China.

9.4.7.4 Summary

In the 2006 O₃ AQCD, the impact of O₃ exposure on species competition and community
 composition was assessed. Ozone was found to cause a significant decline in ponderosa
 and Jeffrey pine in the San Bernardino Mountains in southern California. Ozone exposure

1	also tended to shift the grass-legume mixtures in favor of grass species (U.S. EPA,
2	<u>2006b</u>). Since the 2006 O_3 AQCD, more evidence has shown that O_3 exposure changed
3	the competitive interactions and could lead to loss of O ₃ sensitive species or genotypes.
4	Studies at plant level found that the severity of O ₃ damage on growth, reproduction and
5	foliar injury varied among species, which provided the biological plausibility for the
6	alteration of community composition. Additionally, research since the last review has
7	shown that O ₃ can alter community composition and diversity of soil microbial
8	communities.
9	The decline of conifer forests under O ₃ exposure was continually observed in several
10	regions. Ozone damage was believed to be an important causal factor in the dramatic
11	decline of sacred fir in the valley of Mexico (de Lourdes de Bauer and Hernandez-
12	Tejeda, 2007), as well as cembran pine in southern France and the Carpathian Mountains
13	(Wieser et al., 2006). Results from the Aspen FACE site indicated that O_3 could alter
14	community composition of broadleaf forests as well. At the Aspen FACE site, O_3
15	reduced growth and increased mortality of a sensitive aspen clone, while the O ₃ tolerant
16	clone emerged as the dominant clone in the pure aspen community. In the mixed aspen-
17	birch and aspen-maple communities, O ₃ reduced the competitive capacity of aspen
18	compared to birch and maple (<u>Kubiske et al., 2007</u>).
19	The tendency for O_3 -exposure to shift the biomass of grass-legume mixtures in favor of
20	grass species, was reported in the 2006 O ₃ AQCD and has been generally confirmed by
21	recent studies. However, in a high elevation mature/species-rich grass-legume pasture, O ₃
22	fumigation showed no significant impact on community composition (Bassin et al.,
23	<u>2007b</u>).
24	Ozone exposure not only altered community composition of plant species, but also
25	microorganisms. The shift in community composition of bacteria and fungi has been
26	observed in both natural and agricultural ecosystems, although no general patterns could
27	be identified (Kanerva et al., 2008; Morsky et al., 2008; Kasurinen et al., 2005).
28	The evidence is sufficient to conclude that there is likely to be a causal relationship
29	between O_3 exposure and the alteration of community composition of some
30	ecosystems.

9.4.8 Factors that Modify Functional and Growth Response

31	Many biotic and abiotic factors, including insects, pathogens, root microbes and fungi,
32	temperature, water and nutrient availability, and other air pollutants, as well as elevated
33	CO ₂ , influence or alter plant response to O ₃ . These modifying factors were

1	comprehensively reviewed in AX9.3 of the 2006 O3 AQCD and thus, this section serves
2	mainly as a brief summary of the previous findings. A limited number of new studies
3	published since the 2006 O_3 AQCD add to the understanding of the role of these
4	interactions in modifying O3-induced plant responses. Many of these modifying factors
5	and interactions are integrated into discussions elsewhere in this chapter and the reader is
6	directed to those sections.

9.4.8.1 Genetics

7	It is well known that species vary greatly in their responsiveness to O_3 . Even within a
8	given species, individual genotypes or populations can also vary significantly with
9	respect to O_3 sensitivity (U.S. EPA, 2006b). Therefore, caution should be taken when
10	considering a species' degree of sensitivity to O_3 . Plant response to O_3 is determined by
11	genes that are directly related to oxidant stress and to an unknown number of genes that
12	are not specifically related to oxidants, but instead control leaf and cell wall thickness,
13	stomatal conductance, and the internal architecture of the air spaces. It is rarely the case
14	that single genes are responsible for O_3 tolerance. Studies using molecular biological
15	tools and transgenic plants have positively verified the role of various genes and gene
16	products in O ₃ tolerance and are continuing to increase the understanding of O ₃ toxicity
17	and differences in O_3 sensitivity. See Section <u>9.3.3.2</u> of this document for a discussion of
18	recent studies related to gene expression changes in response to O_3 .

9.4.8.2 Environmental Biological Factors

19	As stated in the 2006 O_3 AQCD, the biological factors within the plant's environment
20	that may influence its response to O ₃ encompass insects and other animal pests, diseases,
21	weeds, and other competing plant species. Ozone may influence the severity of a disease
22	or infestation by a pest or weed, either by direct effects on the causal species, or
23	indirectly by affecting the host, or both. In addition, the interaction between O ₃ , a plant,
24	and a pest, pathogen, or weed may influence the response of the target host species to O_3
25	(U.S. EPA, 2006b). Several recent studies on the effects of O_3 on insects via their
26	interactions with plants are discussed in Section <u>9.4.9.1</u> . In addition, O_3 has also been
27	shown to alter soil fauna communities (Section 9.4.9.2).
28	In contrast to detrimental biological interactions, there are mutually beneficial
29	relationships or symbioses involving higher plants and bacteria or fungi. These include
30	(1) the nitrogen-fixing species <i>Rhizobium</i> and <i>Frankia</i> that nodulate the roots of legumes
31	and alder and (2) the mycorrhizae that infect the roots of many crop and tree species, all

1 2	of which may be affected by exposure of the host plants to O_3 . Some discussion of mycorrhizae can be found in Section 9.4.6.
2	$\frac{9.4.0}{2}$
3	In addition to the interactions involving animal pests, O_3 also has indirect effects on
4	higher herbivorous animals, e.g., livestock, due to O_3 -induced changes in feed quality.
5	Recent studies on the effects of O ₃ on nutritive quality of plants are discussed in
6	Section <u>9.4.4.2</u> .
7	Intra- and interspecific competition are also important factors in determining vegetation
8	response to O ₃ . Plant competition involves the ability of individual plants to acquire the
9	environmental resources needed for growth and development: light, water, nutrients, and
10	space. Intraspecific competition involves individuals of the same species, typically in
11	monoculture crop situations, while interspecific competition refers to the interference
12	exerted by individuals of different species on each other when they are in a mixed
13	culture. This topic was previously reviewed in AX9.3.3.4 of the 2006 O ₃ AQCD. Recent
14	studies on competition and its implications for community composition are discussed in
15	Section <u>9.4.7</u> .

9.4.8.3 Physical Factors

16	Physical or abiotic factors play a large role in modifying plant response to O_3 , and have
17	been extensively discussed in previous O ₃ AQCDs. This section summarizes those
18	findings as well as recent studies published since the last review.
19	Although some studies have indicated that O_3 impact significantly increases with
20	increased ambient temperature (Ball et al., 2000; Mills et al., 2000), other studies have
21	indicated that temperature has little effect (Balls et al., 1996; Fredericksen et al., 1996). A
22	recent study by Riikonen et al. Riikonen et al. (2009) at the Ruohoniemi open air
23	exposure field in Kuopio, Finland found that the effects of temperature and O_3 on total
24	leaf area and photosynthesis of Betula pendula were counteractive. Elevated O3 reduced
25	the saplings' ability to utilize the warmer growth environment by increasing the stomatal
26	limitation for photosynthesis and by reducing the redox state of ascorbate in the apoplast
27	in the combination treatment as compared to temperature alone (Riikonen et al., 2009).
28	Temperature affects the rates of all physiological processes based on enzyme catalysis
29	and diffusion; each process and overall growth (the integral of all processes) has a
30	distinct optimal temperature range. It is important to note that a plant's response to
31	changes in temperature will depend on whether it is growing near its optimum
32	temperature for growth or near its maximum temperature (Rowland-Bamford, 2000).
33	However, temperature is very likely an important variable affecting plant O ₃ response in

1	the presence of the elevated CO_2 levels contributing to global climate change. In contrast,
2	some evidence suggests that O ₃ exposure sensitizes plants to low temperature stress
3	(<u>Colls and Unsworth, 1992</u>) and, also, that O_3 decreases below-ground carbohydrate
4	reserves, which may lead to responses in perennial species ranging from rapid demise to
5	impaired growth in subsequent seasons (i.e., carry-over effects) (Andersen et al., 1997).
6	Light, a component of the plant's physical environment, is an essential "resource" of
7	energy content that drives photosynthesis and C assimilation. It has been suggested that
8	increased light intensity may increase the O3 sensitivity of light-tolerant species while
9	decreasing that of shade-tolerant species, but this appears to be an oversimplification with
10	many exceptions. Several studies suggest that the interaction between O_3 sensitivity and
11	light environment is complicated by the developmental stage as well as the light
12	environment of individual leaves in the canopy (Kitao et al., 2009; Topa et al., 2001;
13	Chappelka and Samuelson, 1998).
14	Although the relative humidity of the ambient air has generally been found to increase the
15	effects of O ₃ by increasing stomatal conductance (thereby increasing O ₃ flux into the
16	leaves), abundant evidence also indicates that the ready availability of soil moisture
17	results in greater O_3 sensitivity (Mills, 2002). The partial "protection" against the effects
18	of O_3 afforded by drought has been observed in field experiments (Low et al., 2006) and
19	modeled in computer simulations (Broadmeadow and Jackson, 2000). Conversely,
20	drought may exacerbate the effects of O_3 on plants (<u>Pollastrini et al., 2010</u> ; <u>Grulke et al.</u> ,
21	<u>2003b</u>). There is also some evidence that O_3 can predispose plants to drought stress
22	(Maier-Maercker, 1998). Hence, the nature of the response is largely species-specific and
23	will depend to some extent upon the sequence in which the stressors occur.

9.4.8.4 Interactions with other Pollutants

Ozone-nitrogen interactions

24	Elevated O_3 exposure and N deposition often co-occur. However, the interactions of O_3
25	exposure and N deposition on vegetation are complex and less well understood compared
26	to their independent effects. Consistent with the conclusion of the 2006 O_3 AQCD, the
27	limited number of studies published since the last review indicated that the interactive
28	effects of N and O_3 varied among species and ecosystems (<u>Table 9-8</u>). To better
29	understand these interactions in ecosystems across the U.S., more information is needed
30	considering combined O ₃ exposure and N deposition related effects.
31	Nitrogen deposition could stimulate relative growth rate (RGR), and lead to increased
32	stomatal conductance. Therefore, plants might become more susceptible to O ₃ exposure.

1	Alternatively, N deposition may increase the availability of photosynthates for use in
2	detoxification and plants could become more tolerant to O ₃ (Bassin et al., 2007a). Only a
3	few recent studies have investigated the interactive effects of O ₃ and N in the U.S. Grulke
4	et al. (2005) measured stomatal conductance of California black oak (Quercus kelloggii)
5	at a long-term N-enrichment site located in the San Bernardino Mountains, which is
6	accompanied by high O ₃ exposure (80 ppb, 24-h avg. over a six month growing season).
7	The authors found that N amendment led to poor stomatal control in full sun in
8	midsummer of the average precipitation years, but enhanced stomatal control in shade
9	leaves of California black oak. In an OTC study, <u>Handley and Grulke (2008</u>) found that
10	O_3 lowered photosynthetic ability and water-use efficiency, and increased leaf chlorosis
11	and necrosis of California black oak. Nitrogen fertilization tended to reduce plant
12	sensitivity to O ₃ exposure; however, the interaction was not statistically significant. In
13	another study, Grulke et al. (2008) reported that various lines of phenomenological and
14	experimental evidence indicate that N deposition and O ₃ pollution contribute to the
15	susceptibility of forests to wildfire in the San Bernadino Mountains by increasing stress
16	due to drought, weakening trees, and predisposing them to bark beetle infestation
17	(<u>U.S. EPA, 2008; NOx/SOx ISA</u>).
18	Studies and here is to be U.S. and the summaries the Table 0.S. Concerlles the
	Studies conducted outside the U.S. are also summarized in <u>Table 9-8</u> . Generally, the
19	responses were species specific. The O ₃ -induced reduction in photosynthetic rate and
20	biomass loss were greater in the relatively high N treatment for watermelon (Citrillus
21	lanants) (Calatayud et al., 2006) and Japanese beech (Fagus crenata) seedlings
22	(<u>Yamaguchi et al., 2007</u>). However, there was no significant interactive effect of O_3 and
23	N on biomass production for Quercus serrata seedlings (Watanabe et al., 2007), young
24	Norway spruce (Picea abies) trees (Thomas et al., 2005), and young European beech
25	(Fagus sylvatica) trees Thomas et al. (2006).

Table 9-8	Response of plants to the interactive effects of elevated ozone
	exposure and nitrogen enrichment.

Site	Species	Ozone exposure	N addition	Responses	References
San Bernardino Mountains, U.S.	California black oak (<i>Quercus</i> <i>kelloggii</i>)	80 ppb	0, and 50 kg N/ ha/yr	N-amended trees had lower late summer C gain and greater foliar chlorosis in the drought year, and poor stomatal control and lower leaf water use efficiency and in midsummer of the average precipitation year.	<u>Grulke et al. (2005</u>)
San Bernardino Mountains, U.S.	California black oak (Quercus kelloggii)	0, 75, and 150 ppb	0, and 50 kg N/ ha/yr	N fertilization tended to reduce plant sensitivity to O_3 exposure; however the interaction was not statistically significant.	<u>Handley and Grulke</u> (2008)

Site	Species	Ozone exposure	N addition	Responses	References
Switzerland	Spruce trees (<i>Picea abies</i>)	Filtered (19.4- 28.1 ppb); ambient (37.6-47.4 ppb)	0, 20, 40 and 80 kg N/ ha/yr	Higher N levels alleviated the negative impact of O_3 on root starch concentrations	<u>Thomas et al.</u> (2005)
Switzerland	Beech trees (<i>Fagus</i> <i>sylvatica</i>)	Filtered (19.4- 28.1 ppb); ambient (37.6-47.4 ppb)	0, 20, 40 and 80 kg N/ ha/yr	N addition amplified the negative effects of O_3 on leaf area and shoot elongation.	<u>Thomas et al.</u> (2006)
Switzerland	Alpine pasture	Ambient (AOT40 of 11.1-12.6 ppm-h); 1.2 ambient (AOT40 of 15.2-29.5 ppm-h) and 1.6 ambient (28.4- 64.9 ppm-h)	0, 5, 10, 25, 50 kg N/ ha/yr	The positive effects of N addition on canopy greenness were counteracted by accelerated leaf senescence in the highest O ₃ treatment.	Bassin et al. (2007b)
Switzerland	Alpine pasture	Ambient (AOT40 of 11.1-12.6 ppm-h); 1.2 ambient (AOT40 of 15.2-29.5 ppm-h) and 1.6 ambient (28.4- 64.9 ppm-h)	0, 5, 10, 25, 50 kg N/ha/yr	Only a small number of species showed significant O_3 and N interactive effects on leaf chlorophyll concentration, leaf weight and change in ¹⁸ O, and the patterns were not consistent.	<u>Bassin et al. (2009</u>)
Switzerland	Alpine pasture	Ambient (AOT40 of 11.1-12.6 ppm-h); 1.2 ambient (AOT40 of 15.2-29.5 ppm-h) and 1.6 ambient (28.4- 64.9 ppm-h)	0, 5, 10, 25, 50 kg N/ ha/yr	The positive effects of N addition on canopy greenness were counteracted by accelerated leaf senescence in the highest O ₃ treatment.	Bassin et al. (2007b)
Switzerland	Alpine pasture	Ambient (AOT40 of 11.1-12.6 ppm-h); 1.2 ambient (AOT40 of 15.2-29.5 ppm-h) and 1.6 ambient (28.4- 64.9 ppm-h)	0, 5, 10' 25, 50 kg N/ ha/yr	Highest N addition resulted in carbon loss, but there was no interaction between O₃ and N treatments.	<u>Volk et al. (2011</u>)
Spain	Watermelon (<i>Citrillus</i> <i>lanants</i>)	O_3 free (AOT40 of 0 ppm-h), ambient (AOT40 of 5.1- 6.3 ppm-h) and elevated O_3 (AOT40 of 32.5-35.6 ppm-h)	140, 280, and 436 kg N/ ha/yr	High N concentration enhanced the detrimental effects of O_3 on Chlorophyll a fluorescence parameters, lipid peroxidation, and the total yield.	<u>Calatayud et al.</u> (2006)
Spain	Trifolium striatum	Filtered (24-h avg. of 8- 22 ppb); ambient (29- 34 ppb), elevated O ₃ (35-56 ppb)	10, 30, and 60 kg N/ ha/yr	O_3 reduced total aerial biomass. N fertilization counterbalanced O_3 -induced effects only when plants were exposed to moderate O_3 levels (ambient) but not under elevated O_3 concentrations.	<u>Sanz et al. (2007</u>)
Japan	Japanese beech seedlings (<i>Fagus</i> <i>crenata</i>)	Filtered (24-h avg. of 10.3-13.2 ppb); ambient (42.0- 43.3 ppb), 1.5 ambient (62.6-63.9 ppb) and 2.0 ambient (82.7- 84.7 ppb)	0, 20 and 50 kg N/ ha/yr	The O_3 -induced reduction in net photosynthesis and whole-plant dry mass were greater in the relatively high N treatment than that in the low N treatment.	Yamaguchi et al. (2007)
Japan	Quercus serrata seedlings	Filtered (24-h avg. of 10.3-13.2 ppb); ambient (42.0- 43.3 ppb), 1.5 ambient (62.6-63.9 ppb) and 2.0 ambient (82.7- 84.7 ppb)	0, 20 and 50 kg N/ ha/yr	No significant interactive effects of O_3 and N load on the growth and net photosynthetic rate were detected.	<u>Watanabe et al.</u> (2007)

Ozone-carbon dioxide interactions

1	Several decades of research has shown that exposure to elevated CO_2 increases
2	photosynthetic rates (Bernacchi et al., 2006; Bernacchi et al., 2005; Tissue et al., 1999;
3	Tissue et al., 1997; Will and Ceulemans, 1997), decreases stomatal conductance
4	(Ainsworth and Rogers, 2007; Paoletti et al., 2007; Bernacchi et al., 2006; Leakey et al.,
5	2006; Medlyn et al., 2001) and generally increases the growth of plants (McCarthy et al.,
6	2009; Norby et al., 2005). This is in contrast to the decrease in photosynthesis and growth
7	in many plants that are exposed to elevated O ₃ . The interactive effects on vegetation have
8	been the subject of research in the past two decades due to the implications on
9	productivity and water use of ecosystems. This area of research was discussed in detail in
10	AX9.3.8.1 of the 2006 O_3 AQCD and the conclusions made then are still relevant (U.S.
11	<u>EPA, 2006b</u>).
12	The bulk of the available evidence shows that, under the various experimental conditions
13	used (which almost exclusively employed abrupt or "step" increases in CO_2
14	concentration, as discussed below), increased CO_2 levels (ambient + 200 to 400 ppm)
15	may protect plants from the negative effects of O_3 on growth. This protection may be
16	afforded in part by CO_2 acting together with O_3 in inducing stomatal closure, thereby
17	reducing O_3 uptake, and in part by CO_2 reducing the negative effects of O_3 on Rubisco
18	and its activity in CO_2 -fixation. Although both CO_2 -induced and O_3 -induced decreases in
19	stomatal conductance have been observed primarily in short-term studies, recent data
20	show a long-term and sustained reduction in stomatal conductance under elevated CO_2
21	for a number of species (Ainsworth and Long, 2005; Ellsworth et al., 2004; Gunderson et
22	al., 2002). Instances of increased stomatal conductance have also been observed in
23	response to O3 exposure, suggesting partial stomatal dysfunction after extended periods
24	of exposure (Paoletti and Grulke, 2010; Grulke et al., 2007a; Maier-Maercker, 1998).
25	Important caveats must be raised with regard to the findings presented in published
26	research. The first caveat concerns the distinctly different natures of the exposures to O_3
27	and CO ₂ experienced by plants in the field. Changes in the ambient concentrations of
28	these gases have very different dynamics. In the context of climate change, CO ₂ levels
29	increase relatively slowly (globally 2 ppm/year) and may change little over several
30	seasons of growth. On the other hand, O_3 presents a fluctuating stressor with considerable
31	hour-to-hour, day-to-day and regional variability (Polle and Pell, 1999). Almost all of the
32	evidence presented comes from experimentation involving plants subjected to an abrupt
33	step increase to a higher, steady CO_2 concentration. In contrast, the O_3 exposure
34	concentrations usually varied from day to day. Luo and Reynolds (1999), Hui et al.
35	(2002), and Luo (2001) noted the difficulties in predicting the likely effects of a gradual
36	CO_2 increase from experiments involving a step increase or those using a range of CO_2

concentrations. It is also important to note that the levels of elevated CO ₂ in many of the
studies will not be experienced in the field for 30 or 40 years, but elevated levels of O_3
can occur presently in several areas of the U.S. Therefore, the $CO_2 \times O_3$ interaction
studies may be less relevant for current ambient conditions.
Another caveat concerns the interactions of O ₃ and CO ₂ with other climatic variables,
such as temperature and precipitation. In light of the key role played by temperature in
regulating physiological processes and modifying plant response to increased CO ₂ levels
(Morison and Lawlor, 1999; Long, 1991) and the knowledge that relatively modest
increases in temperature may lead to dramatic consequences in terms of plant
development (Lawlor, 1998), it is important to consider that studying CO_2 and O_3
interactions alone may not create a complete understanding of effects on plants under
future climate change.

9.4.9 Insects and Other Wildlife

9.4.9.1 Insects

13	Insects may respond indirectly to changes in plants (i.e., increased reactive oxygen
14	species, altered phytochemistry, altered nutrient content) that occur under elevated O ₃
15	conditions, or O_3 can have a direct effect on insect performance (Menendez et al., 2009).
16	Effects of O ₃ on insects occur at the species level (i.e., growth, survival, reproduction,
17	development, feeding behavior) and at the population and community-level
18	(i.e., population growth rate, community composition). In general, effects of O_3 on
19	insects are highly context- and species-specific (Lindroth, 2010; Bidart-Bouzat and Imeh-
20	Nathaniel, 2008). Furthermore, plant responses to O_3 exposure and herbivore attack have
21	been demonstrated to share signaling pathways, complicating characterization of these
22	stressors (Lindroth, 2010; Menendez et al., 2010, 2009). Although both species-level and
23	population and community-level responses to elevated O_3 are observed in field and
24	laboratory studies discussed below, there is no consensus on how insects respond to
25	feeding on O ₃ -exposed plants.

Species-level responses

26	In considering insect growth, survival and reproduction in elevated O ₃ conditions, several
27	studies have indicated an effect while others have found no correlation. The performance
28	of five herbivore species (three moths and two weevils) was assessed in an OTC
29	experiment at $2 \times$ ambient concentration (<u>Peltonen et al., 2010</u>). Growth of larvae of the

- 1 Autumnal moth, *Epirrita autumna*, was significantly decreased in the O₃ treatment while 2 no effects were observed in the other species. In an aphid oviposition preference study 3 using birch buds grown in a three year OTC experiment, O_3 had neither a stimulatory or 4 deterring effect on egg-laying (Peltonen et al., 2006). Furthermore, changes in birch bud 5 phenolic compounds associated with the doubled ambient concentrations of O₃ did not 6 correlate with changes in aphid oviposition (Peltonen et al., 2006). Reproduction in 7 *Popillia japonica*, that were fed soybeans and grown under elevated O_3 appeared to be 8 unaffected (O'Neill et al., 2008). In a meta-analysis of effects of elevated O₃ on 22 9 species of trees and 10 species of insects, the rates of survival, reproduction and food 10 consumption were typically unaffected while development times were reduced and pupal 11 masses were increased (Valkama et al., 2007).
- 12 At the Aspen FACE site insect performance under elevated (50-60 ppb) O_3 conditions 13 (approximately $1.5 \times$ background ambient levels of 30-40 ppb O₃) have been considered 14 for several species. Cumulative fecundity of aphids (Cepegillettea betulaefoliae), that 15 were reared on O_3 -exposed paper birch (*Betula papyrifera*) trees, was lower than aphids 16 from control plots (Awmack et al., 2004). No effects on growth, development, adult 17 weight, embryo number and birth weight of newborn nymphs were observed. In a study 18 conducted using three aspen genotypes, performance of the aspen beetle (Chrysomela 19 crochi) decreased across all parameters measured (development time, adult mass and 20 survivorship) under elevated O_3 (Vigue and Lindroth, 2010). There was an increase in the 21 development time of male and female aspen beetle larvae although the percentages varied 22 across genotypes. Decreased beetle adult mass and survivorship was observed across all 23 genotypes under elevated O₃ conditions. Another study from the Aspen FACE site did 24 not find any significant effects of elevated O_3 on performance (longevity, fecundity, 25 abundance) of the invasive weevil (*Polydrusus sericeus*) (Hillstrom et al., 2010b).
- 26 Since the 2006 O_3 AQCD, several studies have considered the effect of elevated O_3 on 27 feeding behavior of insects. In a feeding preference study, the common leaf weevil 28 (*Phyllobius pyri*) consumed significantly more leaf discs from one aspen clone when 29 compared to a second clone under ambient air conditions (Freiwald et al., 2008). In a 30 moderately elevated O_3 environment (1.5 × ambient), this preference for a certain aspen 31 clone was less evident, however, leaves from O_3 -exposed trees were significantly 32 preferred to leaves grown under ambient conditions. Soybeans grown under enriched O_3 33 had significantly less loss of leaf tissue to herbivory in August compared to earlier in the 34 growing season (July) when herbivory was not affected (Hamilton et al., 2005). Other 35 plant-herbivore interactions have shown no effects of elevated O_3 on feeding. Feeding 36 behavior of Japanese beetles (*P. japonica*) appeared to be unchanged when beetles were 37 fed soybean leaves grown under elevated O_3 conditions (O'Neill et al., 2008). At the 38 Aspen FACE site, feeding by the invasive weevil (*Polydrusus sericeus*), as measured by

leaf area consumption, was not significantly different between foliage that was grown under elevated O_3 versus ambient conditions (<u>Hillstrom et al., 2010b</u>).

Population-level and community-level responses

3 Recent data on insects provide evidence of population-level and community-level 4 responses to O₃. Elevated levels of O₃ can affect plant phytochemistry and nutrient 5 content which in turn can alter population density and structure of the associated 6 herbivorous insect communities and impact ecosystem processes (Cornelissen, 2011; 7 Lindroth, 2010). In 72-hour exposures to elevated O_3 , mean relative growth rate of the 8 aphid Diuraphis noxia increased with O₃ concentration suggesting that more rapid 9 population growth may occur when atmospheric O_3 is elevated (Summers et al., 1994). In 10 a long-term study of elevated O_3 on herbivore performance at the Aspen FACE site, 11 individual performance and population-level effects of the aphid C. betulaefoliae were 12 assessed. Elevated O₃ levels had a strong positive effect on the population growth rates of 13 the aphids; although effects were not detected by measuring growth, development, adult 14 weight, embryo number or birth weight of newborn nymphs (Awmack et al., 2004). 15 Conversely, a lower rate of population growth was observed in aphids previously 16 exposed to O_3 in an OTC (Menendez et al., 2010). No direct effects of O_3 were observed; 17 however, nymphs born from adults exposed to and feeding on O_3 exposed plants were 18 less capable of infesting new plants when compared to nymphs in the control plots 19 (Menendez et al., 2010). Elevated O₃ reduced total arthropod abundance by 17% at 20 Aspen FACE, largely as a result of the negative effects on parasitoids, although phloem-21 feeding insects may benefit (Hillstrom and Lindroth, 2008). Herbivore communities 22 affected by O₃ and N were sampled along an air pollution gradient in the Los Angeles 23 basin (Jones and Paine, 2006). Abundance, diversity, and richness of herbivores were not 24 affected. However, a shift in community structure, from phloem-feeding to chewing 25 dominated communities, was observed along the gradient. No consistent effect of 26 elevated O_3 on herbivory or insect population size was detected at SoyFACE (O'Neill et 27 al., 2010; Dermody et al., 2008). 28 Evidence of modification of insect populations and communities in response to elevated

29 O₃ includes genotypic and phenotypic changes. In a study conducted at the Aspen FACE 30 site, elevated O₃ altered the genotype frequencies of the pea aphid (Acyrthosiphon pisum) 31 grown on red clover (Trifolium pratense) over multiple generations (Mondor et al., 32 2005). Aphid color was used to distinguish between the two genotypes. Ozone increased 33 the genotypic frequencies of pink-morph: green-morph aphids from 2:1 to 9:1, and 34 depressed wing-induction responses more strongly in the pink than the green genotype 35 (Mondor et al., 2005). Growth and development of individual green and pink aphids 36 reared as a single genotype or mixed genotypes were unaffected by elevated O_3 (Mondor

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et al., 2010). However, growth of pea aphid populations is not readily predictable using individual growth rates.

9.4.9.2 Wildlife

Herpetofauna

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3	Since the 2006 O ₃ AQCD, direct effects of O ₃ exposure including physiological changes
4	and alterations of ecologically important behaviors such as feeding and thermoregulation
5	have been observed in wildlife. These studies have been conducted in limited laboratory
6	exposures, and the levels of O_3 treatment (e.g., 0.2-0.8 ppm) were often unrealistically
7	higher than the ambient levels. Amphibians may be especially vulnerable to airborne
8	oxidants due to the significant gas exchange that occurs across the skin (Andrews et al.,
9	<u>2008</u> ; <u>Dohm et al., 2008</u>). Exposure to 0.2 ppm to 0.8 ppm O_3 for 4 hours resulted in a
10	decrease of oxygen consumption and depressed lung ventilation in the California tree
11	frog Pseudacris cadaverina (Mautz and Dohm, 2004). Following a single 4-h inhalation
12	exposure to 0.8 ppm O ₃ , reduced pulmonary macrophage phagocytosis was observed at 1
13	and 24 hours postexposure in the marine toad (Bufo marinus) indicating an effect on
14	immune system function (Dohm et al., 2005). There was no difference in macrophage
15	function at 48 hours postexposure in exposed and control individuals.

- 16 Behavioral effects of O₃ observed in amphibians include responses to minimize the 17 surface area of the body exposed to the air and a decrease in feeding rates (Dohm et al., 18 2008; Mautz and Dohm, 2004). The adoption of a low-profile "water conservation 19 posture" during O_3 exposure was observed in experiments with the California tree frog 20 (Mautz and Dohm, 2004). Marine toads, Bufo marinus, exposed to 0.06 µL/L O₃ for 21 4 hours ate significantly fewer mealworms at 1 hour and 48 hours postexposure than 22 control toads (Dohm et al., 2008). In the same study, escape/exploratory behavior as 23 measured by total distance moved was not negatively affected in the O₃-exposed 24 individuals as compared to the controls (Dohm et al., 2008).
- 25 Water balance and thermal preference in herpetofauna are altered with elevated O_3 . 26 Marine toads exposed to 0.8 ppm O_3 for 4 hours exhibited behavioral hypothermia when 27 temperature selection in the toads was assessed at 1, 24 and 48 hours postexposure 28 (Dohm et al., 2001). Ozone-exposed individuals lost almost 5g more body mass on 29 average than controls due to evaporative water loss. At 24 hours after exposure, the 30 individuals that had lost significant body mass selected lower body temperatures (Dohm 31 et al., 2001). Behavioral hypothermia was also observed in reptiles following 4-h 32 exposures to 0.6 ppm O₃. Exposure of the Western Fence Lizard (Sceloporus

1	occidentalis) at 25°C induced behavioral hypothermia that recovered to control
2	temperatures by 24 hours (Mautz and Dohm, 2004). The behavioral hypothermic
3	response persisted in lizards exposed to O_3 at 35°C at 24 hours postexposure resulting in a
4	mean body temperature of 3.3°C over controls.

Soil fauna communities

Ozone has also been shown to alter soil fauna communities (Meehan et al., 2010;
Kasurinen et al., 2007; Loranger et al., 2004). Abundance of Acari (mites and ticks)
decreased by 47% under elevated O_3 at Aspen FACE site, probably due to the higher
secondary metabolites and lower N concentrations in litter and foliage under elevated O_3
(Loranger et al., 2004). In another study from the Aspen FACE site, leaf litter collected
from aspen grown under elevated O_3 conditions was higher in fiber and lignin
concentrations than litter from trees grown under ambient conditions. These chemical
characteristics of the leaves were associated with increased springtail population growth
following 10 weeks in a laboratory microcosm (Meehan et al., 2010). Consumption rates
of earthworms fed on leaf litter for 6 weeks from trees grown under elevated O_3
conditions and ambient air did not vary significantly between treatments (Meehan et al.,
2010). In another study on juvenile earthworms Lumbricus terrestris, individual growth
was reduced when worms were fed high- O_3 birch litter from trees exposed for three years
to elevated O_3 in an OTC system (Kasurinen et al., 2007). In the same study no
significant growth or mortality effects were observed in isopods.

9.4.9.3 Indirect Effects on Wildlife

20	In addition to the direct effects of O_3 exposure on physiological and behavioral endpoints
21	observed in the laboratory, there are indirect effects to wildlife. These effects include
22	changes in biomass and nutritive quality of O_3 -exposed plants (reviewed in Section <u>9.4.4</u>)
23	that are consumed by wildlife. Reduced digestibility of O3-exposed plants may alter
24	dietary intake and foraging strategies in herbivores. In a study using native highbush
25	blackberry (Rubus argutus) relative feed value of the plants decreased in bushes exposed
26	to double ambient concentrations of O_3 (Ditchkoff et al., 2009). Indirect effects of
27	elevated O ₃ on wildlife include changes in chemical signaling important in ecological
28	interactions reviewed below.

Chemical signaling in ecological interactions

1 Ozone has been shown to degrade or alter biogenic VOC signals important to ecological 2 interactions including; (1) attraction of pollinators and seed dispersers; (2) defense 3 against herbivory; and (3) predator-prey interactions (Pinto et al., 2010; McFrederick et 4 al., 2009; Yuan et al., 2009; Pinto et al., 2007a; Pinto et al., 2007b). Each signal released 5 by emitters has an atmospheric lifetime and a unique chemical signature comprised of 6 different ratios of individual hydrocarbons that are susceptible to atmospheric oxidants 7 such as O_3 (Yuan et al., 2009; Wright et al., 2005). Under elevated O_3 conditions, these 8 olfactory cues may travel shorter distances before losing their specificity (McFrederick et 9 al., 2009; McFrederick et al., 2008). Additional non-phytogenic VOC-mediated 10 interrelationships with the potential to be modified by O₃ include territorial marking, 11 pheromones for attraction of mates and various social interactions including scent trails, 12 nestmate recognition and signals involved in aggregation behaviors (McFrederick et al., 13 2009). For example, the alcohols, ketones and aldehydes comprising sex pheromones in 14 moths could be especially vulnerable to degradation by O_3 , since some males travel >100 15 meters to find mates (Carde and Haynes, 2004). In general, effects of O₃ on scentmediated ecological interactions are highly context- and species-specific (Lindroth, 2010; 16 17 Bidart-Bouzat and Imeh-Nathaniel, 2008).

Pollination and seed dispersal

18 Phytogenic VOC's attract pollinators and seed dispersers to flowers and fruits (Dudareva 19 et al., 2006; Theis and Raguso, 2005). These floral scent trails in plant-insect interactions 20 may be destroyed or transformed by O_3 (McFrederick et al., 2008). Using a Lagrangian 21 model, the rate of destruction of phytogenic VOC's was estimated in air parcels at 22 increasing distance from a source in response to increased regional levels of O_3 , hydroxyl 23 and nitrate radicals (McFrederick et al., 2008). Based on the model, the ability of 24 pollinators to locate highly reactive VOCs from emitting flowers may have decreased 25 from kilometers during pre-industrial times to <200 meters at current ambient conditions 26 (McFrederick et al., 2008). Scents that travel shorter distances (0-10 m) are less 27 susceptible to air pollutants, while highly reactive scents that travel longer distances (10 28 to 100's of meters), are at a higher risk for degradation (McFrederick et al., 2009). For 29 example, male euglossine bees can detect bait stations from a distance of at least one 30 kilometer (Dobson, 1994).

Defense against herbivory

1	Ozone can alter the chemical signature of VOCs emitted by plants and these VOCs are
2	subsequently detected by herbivores (Blande et al., 2010; Iriti and Faoro, 2009; Pinto et
3	al., 2007a; Vuorinen et al., 2004; Jackson et al., 1999; Cannon, 1990). These
4	modifications can make the plant either more attractive or repellant to phytophagous
5	insects (Pinto et al., 2010). For example, under elevated O_3 , the host plant preference by
6	forest tent caterpillars increased for birch compared to aspen (Agrell et al., 2005).
7	Ozone-induced emissions from red spruce needles were found to repel spruce budworm
8	larvae (Cannon, 1990). Transcriptional profiles of field grown soybean (Glycine max)
9	grown in elevated O ₃ conditions were altered due to herbivory by Japanese beetles. The
10	herbivory resulted in a higher number of transcripts in the leaves of O ₃ -exposed plants
11	and upregulation of antioxidant metabolism associated with plant defense (Casteel et al.,
12	<u>2008</u>).

13Ozone may modify signals involved in plant-to-plant interactions and plant defense14against pathogens (Blande et al., 2010; Pinto et al., 2010; McFrederick et al., 2009; Yuan15et al., 2009). In a recent study with lima beans, 80 ppb O3 degraded several16herbivore-induced VOCs, reducing the distance over which plant-to-plant signaling17occurred (Blande et al., 2010).

Predator-prey interactions

- 18 Elevated O₃ conditions are associated with disruption of pheromone-mediated 19 interactions at higher trophic levels (e.g., predators and parasitoids of herbivores). In a 20 study from the Aspen FACE site, predator escape behaviors of the aphid (Chatophorus 21 *stevensis*) were enhanced on O_3 -fumigated aspen trees although the mechanism of this 22 response remains unknown (Mondor et al., 2004). The predatory mite *Phytoseiulus* 23 persimilis can distinguish between the VOC signature of ozonated lima bean plants and 24 ozonated lima bean plants simultaneously damaged by T. urticae (Vuorinen et al., 2004) 25 however, other tritrophic interactions have shown no effect (Pinto et al., 2007b).
- 26 There are few studies that consider host location behaviors of parasites under elevated O_3 . 27 In closed chambers funigated with O_3 , the searching efficiency and proportion of the 28 host larval fruit flies parasitized by Asobara tabida declined when compared to filtered 29 air controls (Gate et al., 1995). The host location behavior and rate of parasitism of the 30 wasp (Coesia plutellae) on Plutella xylostella-infested potted cabbage plants was tested 31 under ambient and doubled O₃ conditions in an open-air fumigation system (Pinto et al., 32 2008). The number of wasps found in the field and the percentages of parasitized larvae 33 were not significantly different from controls under elevated O₃.

1 Elevated O₃ has the potential to perturb specialized food-web communication in 2 transgenic crops. In insect-resistant oilseed rape *Brassica napus* grown under 100 ppb O₃ 3 in a growth chamber, reduced feeding damage by Putella xylostella led to deceased 4 attraction of the endoparasitoid (Costesia vestalis), however this tritrophic interaction 5 was influenced by the degree of herbivore feeding (Himanen et al., 2009a; Himanen et 6 al., 2009b). Under chronic O_3 -exposure, the insect resistance trait BT cry1Ac in 7 transgenic *B. napus* was higher than the control (Himanen et al., 2009c). There was a 8 negative relative growth rate of the Bt target herbivore, *P. xylostella*, in all O₃ treatments.

9.4.9.4 Summary

9	Recent information on O ₃ effects on insects and other wildlife is limited to a few species
10	and there is no consensus on how these organisms respond to elevated O3. Studies
11	published since the last review show impacts of elevated O3 on both species-level
12	responses (reproduction, growth, feeding behavior) and community and ecosystem-level
13	responses (population growth, abundance, shift in community structure) in some insects
14	and soil fauna. Changes in ecologically important behaviors such as feeding and
15	thermoregulation have recently been observed with O3 exposure in amphibians and
16	reptiles, however, these responses occur at concentrations of O_3 much higher than
17	ambient levels.
18	Recent information available since the last review considers the effects of O ₃ on chemical

18Recent information available since the last review considers the effects of O_3 on chemical19signaling in insect and wildlife interactions. Specifically, studies on O_3 effects on20pollination and seed dispersal, defenses against herbivory and predator-prey interactions21all consider the ability of O_3 to alter the chemical signature of VOCs emitted during these22pheromone-mediated events. The effects of O_3 on chemical signaling between plants,23herbivores and pollinators as well as interactions between multiple trophic levels is an24emerging area of study that may result in further elucidation of O_3 effects at the species,25community and ecosystem-level.

9.5 Effects-Based Air Quality Exposure Indices and Dose Modeling

9.5.1 Introduction

26 27 Exposure indices are metrics that quantify exposure as it relates to measured plant damage (e.g., reduced growth). They are summary measures of monitored ambient O_3

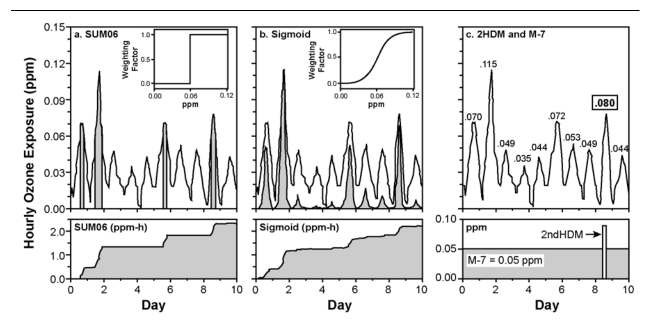
1	concentrations over time, intended to provide a consistent metric for reviewing and
2	comparing exposure-response effects obtained from various studies. Such indices may
3	also provide a basis for developing a biologically-relevant air quality standard for
4	protecting vegetation and ecosystems. Effects on plant growth and/or yield have been a
5	major focus of the characterization of O3 impacts on plants for purposes of the air quality
6	standard setting process (U.S. EPA, 2007b, 1996e, 1986). The relationship of O_3 and
7	plant responses can be characterized quantitatively as "dose-response" or "exposure-
8	response." The distinction is in how the pollutant concentration is expressed: "dose" is
9	the pollutant concentration absorbed by the leaf over some time period, and is very
10	difficult to measure directly, whereas "exposure" is the ambient air concentration
11	measured near the plant over some time period, and summarized for that period using an
12	index. Exposure indices have been most useful in considering the form of the secondary
13	O3 NAAQS, in large part because they only require ambient air quality data rather than
14	more complex indirect calculations of dose to the plant. The attributes of exposure
15	indices that are most relevant to plant damage are the weighting of O_3 concentrations and
16	the daily and seasonal time-periods. Several different types of exposure indices are
17	discussed in Section <u>9.5.2</u> .
18	From a theoretical perspective, a measure of plant O_3 uptake or dose from ambient air
19	(either rate of uptake or cumulative seasonal uptake) might be a better predictor of O_3
20	damage to plants than an exposure index and may be useful in improving risk assessment.
21	An uptake estimate would have to integrate all those environmental factors that influence
22	stomatal conductance, including but not limited to temperature, humidity, and soil water
23	status (Section <u>9.5.4</u>). Therefore, uptake values are generally obtained with simulation
24	models that require knowledge of species- and site-specific values for the variables
25	mentioned. However, a limitation of modeling dose is that environmental variables are
26	poorly characterized. In addition, it has also been recognized that O ₃ detoxification
27	processes and the temporal dynamics of detoxification must be taken into account in dose
28	modeling (Heath et al., 2009) (Section 9.5.4). Because of this, research has focused
29	historically on predictors of O3 damage to plants based only on exposure as a summary
30	measure of monitored ambient pollutant concentration over some integral of time, rather
31	than dose (U.S. EPA, 1996c; Costa et al., 1992; Lee et al., 1988b; U.S. EPA, 1986;
32	Lefohn and Benedict, 1982; O'Gara, 1922).

9.5.2 Description of Exposure Indices Available in the Literature

33	
34	
35	

Mathematical approaches for summarizing ambient air quality information in biologically meaningful forms for O₃ vegetation effects assessment purposes have been explored for more than 80 years (U.S. EPA, 1996b; O'Gara, 1922). In the context of national standards

1	that protect for "known or anticipated" effects on many plant species in a variety of
2	habitats, exposure indices provide a numerical summary of very large numbers of
3	ambient observations of concentration over extended periods. Like any summary statistic,
4	exposure indices retain information on some, but not all, characteristics of the original
5	observations. Several indices have been developed to attempt to incorporate some of the
6	biological, environmental, and exposure factors that influence the magnitude of the
7	biological response and contribute to observed variability (Hogsett et al., 1988). In the
8	1996 O_3 AQCD, the exposure indices were arranged into five categories; (1) One event,
9	(2) Mean, (3) Cumulative, (4) Concentration weighted, and (5) Multicomponent, and
10	were discussed in detail (Lee et al., 1989). Figure 9-9 illustrates how several of the
11	indices weight concentration and accumulate exposure. For example, the SUM06 index
12	(panel a) is a threshold-based approach wherein concentrations below 0.06 ppm are given
13	a weight of zero and concentrations at or above 0.06 ppm are given a weight of 1.0 that is
14	summed, usually over 3 to 6 months. The Sigmoid approach (panel b), which is similar to
15	the W126 index (Lefohn et al., 1988; Lefohn and Runeckles, 1987), is a non-threshold
16	approach wherein all concentrations are given a weight that increases from zero to 1.0
17	with increasing concentration and summed.



(a) SUM06: the upper graphic illustrates an episodic exposure profile; the shaded area under some of the peaks illustrates the concentrations greater than or equal to 0.06 ppm that are accumulated in the index. The insert shows the concentration weighting (0 or 1) function. The lower portion of the graphic illustrates how concentration is accumulated over the exposure period. (b) SIGMOID: the upper graphic illustrates an episodic exposure profile; the variable shaded area under the peaks illustrates the concentration-dependent weights that are accumulated in the index. The insert shows the sigmoid concentration weighting function. This is similar to the W126 function. The lower portion of the graphic illustrates how concentration is accumulated over the exposure period. (c) second HDM and M-7: the upper graphic illustrates an episodic exposure profile. The lower portion of the graphic illustrates that the second HDM considers only a single exposure peak, while the M-7 (average of 7-h daily means) applies a constant exposure value over the exposure period.

Source: Reprinted with permission of Air and Waste Management Association (Tingey et al., 1991).

Figure 9-9 Diagrammatic representation of several exposure indices illustrating how they weight concentration and accumulate exposure.

1	This section will primarily discuss SUM06, W126 and AOTx exposure metrics. Below
2	are the definitions of the three cumulative index forms:
3	• SUM06: Sum of all hourly O ₃ concentrations greater than or equal to
4	0.06 ppm observed during a specified daily and seasonal time window
5	(<u>Figure 9-9a</u>).
6	• AOTx: Sum of the differences between hourly O ₃ concentrations greater than
7	a specified threshold during a specified daily and seasonal time window. For
8	example, AOT40 is sum of the differences between hourly concentarions
9	above 0.04 ppm.
10	 W126: Sigmoidally weighted sum of all hourly O₃ concentrations observed
11	during a specified daily and seasonal time window (Lefohn et al., 1988;
12	Lefohn and Runeckles, 1987), similar to Figure 9-9b). The sigmoidal

1weighting of hourly O3 concentration is given in the equation below, where C2is the hourly O3 concentration in ppm:

$$w_c = \frac{1}{1 + 4403e^{-126C}}$$

Equation 9-1

These indices have a variety of relevant time windows that may be applied and are
discussed in Section <u>9.5.3</u>.

5 Various factors with known or suspected bearing on the exposure-response relationship, 6 including concentration, time of day, respite time, frequency of peak occurrence, plant 7 phenology, predisposition, etc., have been weighted with various functions in a large set 8 of indices. The resulting indices were evaluated by ranking them according to the 9 goodness-of-fit of a regression model of growth or yield response (Lee et al., 1989). The 10 statistical evaluations for each of these indices were completed using growth or yield 11 response data from many earlier exposure studies (e.g., NCLAN). This retrospective 12 approach was necessary because there were no studies specifically designed to test the 13 goodness-of-fit of the various indices. The goodness-of-fit of a set of linear and nonlinear 14 models for exposure-response was ranked as various proposed indices were used in turn 15 to quantify exposure. This approach provided evidence for the best indices. The results of 16 retrospective analyses are described below.

17 Most of the early retrospective studies reporting regression approaches used data from the 18 NCLAN program or data from Corvallis, Oregon or California (Costa et al., 1992; Lee et 19 al., 1988b; Lefohn et al., 1988; Musselman et al., 1988; Lee et al., 1987; U.S. EPA, 20 1986). These studies were previously reviewed by the EPA (U.S. EPA, 1996c; Costa et 21 al., 1992) and were in general agreement that the best fit to the data resulted from using 22 cumulative concentration-weighted exposure indices (e.g., W126, SUM06). Lee et al. 23 (1987) suggested that exposure indices that included all the 24-h data performed better 24 than those that used only 7 hours of data; this was consistent with the conclusions of 25 Heagle et al. (1987) that plants receiving exposures for an additional 5-h/day showed 26 10% greater yield loss than those exposed for 7-h/day. In an analysis using the National 27 Crop Loss Assessment Network (NCLAN) data, Lefohn et al. (1988) found several 28 indices which only cumulated and weighted higher concentrations (e.g., W126, SUM06, 29 SUM08, and AOT40) performed very well. Amongst this group no index had 30 consistently better fits than the other indices across all studies and species (Heagle et al., 31 1994b; Lefohn et al., 1988; Musselman et al., 1988). Lefohn et al. (1988) found that 32 adding phenology weighting to the index somewhat improved the performance of the

indices. The "best" exposure index was a phenologically weighted cumulative index,
 with sigmoid weighting on concentration and a gamma weighting function as a surrogate
 for plant growth stage. This index provided the best statistical fit when used in the models
 under consideration, but it required data on species and site conditions, making
 specification of weighting functions difficult for general use.

- 6Other factors, including predisposition time (Hogsett et al., 1988; McCool et al., 1988)7and crop development stage (Tingey et al., 2002; Heagle et al., 1991) contributed to8variation in the biological response and suggested the need for weighting O39concentrations to account for predisposition time and phenology. However, the roles of10predisposition and phenology in plant response vary considerably with species and11environmental conditions; therefore, specification of a weighting function for general use12in characterizing plant exposure has not been possible.
- 13 European scientists took a similar approach in developing indices describing growth and 14 yield loss in crops and tree seedlings, using OTCs with modified ambient exposures, but 15 many fewer species and study locations were employed in the European studies. There is 16 evidence from some European studies that a lower (Pleijel et al., 1997) or higher (Finnan 17 et al., 1997; Finnan et al., 1996) cutoff value in indices with a threshold may provide a 18 better statistical fit to the experimental data. Finnan et al. (1997) used seven exposure 19 studies of spring wheat to confirm that cumulative exposure indices emphasizing higher 20 O_3 concentrations were best related to plant response and that cumulative exposure 21 indices using weighting functions, including cutoff concentrations, allometric and 22 sigmoidal, provided a better fit and that the ranking of these indices differed depending 23 on the exposure-response model used. Weighting those concentrations associated with 24 sunshine hours in an attempt to incorporate an element of plant uptake did not improve 25 the index performance (Finnan et al., 1997). A more recent study using data from several 26 European studies of Norway spruce, analyzed the relationship between relative biomass 27 accumulation and several cumulative, weighted indices, including the AOT40 (area over 28 a threshold of 40ppb) and the SUM06 (Skarby et al., 2004). All the indices performed 29 relatively well in regressing biomass and exposure index, with the AOT20 and AOT30 30 doing slightly better than others ($r^2 = 0.46-0.47$). In another comparative study of four 31 independent data sets of potato yield and different cumulative uptake indices with different cutoff values, a similarly narrow range of r^2 was observed ($r^2 = 0.3-0.4$) (Pleijel 32 33 et al., 2004b).
- 34In Europe, the cutoff concentration-weighted index AOT40 was selected in developing35exposure-response relationships based on OTC studies of a limited number of crops and36trees (Grunhage and Jager, 2003). The United Nations Economic Commission for Europe37(UNECE, 1988) adopted the critical levels approach for assessment of O3 risk to

1	vegetation across Europe. As used by the UNECE, the critical levels are not like the air
2	quality regulatory standards used in the U.S., but rather function as planning targets for
3	reductions in pollutant emissions to protect ecological resources. Critical levels for O3 are
4	intended to prevent long-term deleterious effects on the most sensitive plant species
5	under the most sensitive environmental conditions, but not intended to quantify O_3
6	effects. A critical level was defined as "the concentration of pollutant in the atmosphere
7	above which direct adverse effects on receptors, such as plants, ecosystems, or materials
8	may occur according to present knowledge" (UNECE, 1988). The nature of the "adverse
9	effects" was not specified in the original definition, which provided for different levels
10	for different types of harmful effect (e.g., visible injury or loss of crop yield). There are
11	also different critical levels for crops, forests, and semi-natural vegetation. The caveat,
12	"according to present knowledge" is important because critical levels are not rigid; they
13	are revised periodically as new scientific information becomes available. For example,
14	the original critical level for O ₃ specified concentrations for three averaging times, but
15	further research and debate led to the current critical level being stated as the cumulative
16	exposure (concentration × hours) over a cutoff concentration of 40 ppb (AOT40) (Fuhrer
17	<u>et al., 1997</u>).

18 More recently in Europe, a decision was made to work towards a flux-based approach 19 (see Section 9.5.4) for the critical levels ("Level II"), with the goal of modeling O_3 20 flux-effect relationships for three vegetation types: crops, forests, and semi-natural 21 vegetation (Grunhage and Jager, 2003). Progress has been made in modeling flux (U.S. 22 EPA, 2006b) and the Mapping Manual is being revised (Ashmore et al., 2004a, b; 23 Grennfelt, 2004; Karlsson et al., 2003). The revisions may include a flux-based approach 24 for three crops: wheat, potatoes, and cotton. However, because of a lack of flux-response 25 data, a cumulative, cutoff concentration-based (AOTx) exposure index will remain in use 26 for the near future for most crops and for forests and semi-natural herbaceous vegetation 27 (Ashmore et al., 2004b).

28 In both the U.S. and Europe, the adequacy of these numerical summaries of exposure in 29 relating biomass and yield changes have, for the most part, all been evaluated using data 30 from studies not necessarily designed to compare one index to another (Skarby et al., 31 2004; Lee et al., 1989; Lefohn et al., 1988). Very few studies in the U.S. have addressed 32 this issue since the 2006 O₃ AQCD. McLaughlin et al. (2007a) reported that the 33 cumulative exposure index of AOT60 related well to reductions in growth rates at forest 34 sites in the southern Appalachian Mountains. However, the authors did not report an 35 analysis to compare multiple indices. Overall, given the available data from previous O_3 36 AQCDs and the few recent studies, the cumulative, concentration-weighted indices 37 perform better than the peak or mean indices. It is still not possible, however, to

1	distinguish the differences in performance among the cumulative, concentration-weighted
2	indices.
3	The main conclusions from the 1996 and 2006 O ₃ AQCDs regarding an index based on
4	ambient exposure are still valid. No information has come forth since the 2006 O ₃ AQCD
5	to alter those conclusions. These key conclusions can be restated as follows:
6	 O₃ effects in plants are cumulative;
7	 higher O₃ concentrations appear to be more important than lower
8	concentrations in eliciting a response;
9	 plant sensitivity to O₃ varies with time of day and plant development stage;
10	and
11	 quantifying exposure with indices that accumulate the O₃ hourly
12	concentrations and preferentially weight the higher concentrations improves
13	the explanatory power of exposure/response models for growth and yield, over
14	using indices based on mean and peak exposure values.
15	Following the 2006 criteria review process (U.S. EPA, 2006b), the EPA proposed an
16	alternative form of the secondary NAAQS for O ₃ using a cumulative, concentration-
17	weighted exposure index to protect vegetation from damage (72 FR 37818). The EPA
18	considered two specific concentration-weighted indices: the cutoff concentration
19	weighted SUM06 and the sigmoid-weighted W126 exposure index (U.S. EPA, 2007b).
20	These two indices performed equally well in predicting the exposure-response
21	relationships observed in the crop and tree seedlings studies (Lee et al., 1989). At a
22	workshop convened to consider the science supporting these indices (Heck and Cowling,
23	<u>1997</u>) there was a consensus that these cumulative concentration-weighted indices being
24	considered were equally capable of predicting plant response. It should be noted that
25	there are some important differences between the SUM06 and W126. When considering
26	the response of vegetation to O_3 exposures represented by the threshold (e.g., SUM06)
27	and non-threshold (e.g., W126) indices, the W126 metric does not have a cut-off in the
28	weighting scheme as does SUM06 and thus it includes consideration of potentially
29	damaging exposures below 60 ppb. The W126 metric also adds increasing weight to
30	hourly concentrations from about 40 ppb to about 100 ppb (Lefohn et al., 1988; Lefohn
31	and Runeckles, 1987). This is unlike cut-off metrics such as the SUM06 where all
32	concentrations above 60 ppb are treated equally. This is an important feature of the W126
33	since as hourly concentrations become higher, they become increasingly likely to
34	overwhelm plant defenses and are known to be more detrimental to vegetation (See
35	Section $9.5.3.1$).

9.5.3 Important Components of Exposure Indices

1	In the previous O ₃ AQCDs it was established that higher hourly concentrations have
2	greater effects on vegetation than lower concentrations (U.S. EPA, 2006b, 1996c).
3	Further, it was determined that the diurnal and seasonal duration of exposure is important
4	for plant response. Weighting of hourly concentrations and the diurnal and seasonal time
5	window of exposure are the most important variables in a cumulative exposure index and
6	will be discussed below. However, these variables should be looked at in the context of
7	plant phenology, diurnal conductance rates, plant canopy structure, and detoxification
8	mechanisms of vegetation as well as the climate and meteorology, all of which are
9	determinants of plant response. These more specific factors will be discussed in the
10	uptake and dose modeling Section $9.5.4$.

9.5.3.1 Role of Concentration

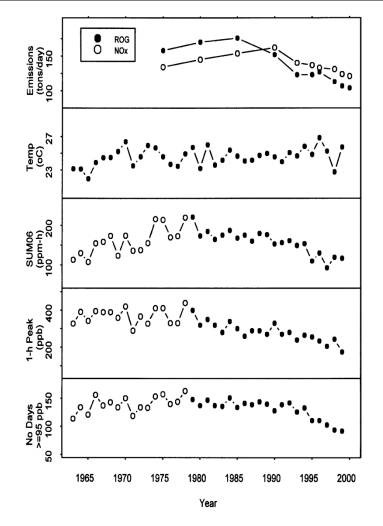
11	The significant role of peak O ₃ concentrations was established based on several
12	experimental studies (U.S. EPA, 1996c). Several studies (Oksanen and Holopainen,
13	2001; Yun and Laurence, 1999; Nussbaum et al., 1995) have added support for the
14	important role that peak concentrations, as well as the pattern of occurrence, plays in
15	plant response to O ₃ . Oksanen and Holopainen (2001) found that the peak concentrations
16	and the shape of the O ₃ exposure (i.e., duration of the event) were important determinants
17	of foliar injury in European white birch saplings, but growth reductions were found to be
18	more related to total cumulative exposure. Based on air quality data from 10 U.S. cities,
19	three 4-week exposure treatments having the same SUM06 value were constructed by
20	Yun and Laurence (1999). The authors used different exposure regimes to explore effects
21	of treatments with variable versus uniform peak occurrence during the exposure period.
22	The authors reported that the variable peak exposures were important in causing injury,
23	and that the different exposure treatments, although having the same SUM06, resulted in
24	very different patterns of foliar injury. Nussbaum et al. (1995) also found peak
25	concentrations and the pattern of occurrence to be critical in determining the measured
26	response. The authors recommended that to describe the effect on total forage yield, peak
27	concentrations >0.11 ppm must be emphasized by using an AOT with higher threshold
28	concentrations.
29	A greater role for peak concentrations in effects on plant growth might be inferred based
30	on air quality analyses for the southern California area (<u>Tingey et al., 2004</u> ; Lee et al.,
31	<u>2003a</u>). In the late 1960s and 1970s, extremely high O_3 concentrations had impacted the
(1/)	

32 San Bernardino National Forest. However, over the past 20+ years, significant reductions
33 in O₃ exposure have occurred (<u>Bytnerowicz et al., 2008; Lee et al., 2003a; Lefohn and</u>

1 Shadwick, 2000; Davidson, 1993). An illustration of this improvement in air quality is 2 shown by the 37-year history of O_3 air quality at the Crestline site in the San Bernardino 3 Mountains (Figure 9-10) (Lee et al., 2003a). Ozone exposure increased from 1963 to 4 1979 concurrent with increased population and vehicular miles, followed by a decline to 5 the present mirroring decreases in precursor emissions. The pattern in exposure was 6 evident in various exposure indices including the cumulative concentration weighted 7 (SUM06), as well as maximum peak event (1-h peak), and the number of days having 8 hourly averaged O_3 concentrations greater than or equal to 95 ppb. The number of days 9 having hourly averaged O₃ concentrations greater than or equal to 95 ppb declined 10 significantly from 163 days in 1978 to 103 days in 1997. The changes in ambient O_3 air quality for the Crestline site were reflected in the changes in frequency and magnitude of 11 12 the peak hourly concentration and the duration of exposure (Figure 9-10). Considering 13 the role of exposure patterns in determining response, the seasonal and diurnal patterns in 14 hourly O₃ concentration did not vary appreciably from year to year over the 37-year 15 period (Lee et al., 2003a).

16 The potential importance of exposure to peak concentrations comes both from results of 17 measures of tree conditions on established plots and from results of model simulations. 18 Across a broad area of the San Bernardino National Forest, the Forest Pest Management 19 (FPM) method of injury assessment indicated an improvement in crown condition from 20 1974 to 1988; and the area of improvement in injury assessment is coincident with an 21 improvement in O₃ air quality (Miller and Rechel, 1999). A more recent analysis of forest 22 changes in the San Bernardino National Forest, using an expanded network of monitoring 23 sites, has verified significant changes in growth, mortality rates, basal area, and species 24 composition throughout the area since 1974 (Arbaugh et al., 2003). A model simulation 25 of ponderosa pine growth over the 40-year period in the San Bernardino National Forest 26 showed a significant impact of O_3 exposure on tree growth and indicates improved 27 growth with reduced O_3 concentrations. This area has also experienced elevated 28 N deposition and based on a number of environmental indicators, it appears that this area 29 is experiencing N saturation (Fenn et al., 1996). To account for this potential interaction, 30 the model simulations were conducted under conditions of unlimited soil N. The actual 31 interactions are not known. The improvement in growth over the years was attributed to 32 improved air quality, but no distinction was made regarding the relative role of 33 "mid-range" and higher hourly concentrations, only that improved growth tracked 34 decreasing SUM06, maximum peak concentration, and number of days of hourly O_3 35 >95 ppb (Tingey et al., 2004). A summary of air quality data from 1980 to 2000 for the 36 San Bernardino National Forest area of the number of "mid-range" hourly concentrations 37 indicated no dramatic changes over this 20-year period, ranging from about 1,500 to 38 2,000 hours per year (Figure 9-11). There was a slow increase in the number of 39 "mid-range" concentrations from 1980 to 1986, which corresponds to the period after

implementation of the air quality standard. Another sharper increase was observed in the late 1990s. This pattern of occurrence of mid-range hourly concentrations suggests a lesser role for these concentration ranges compared to the higher values in either of the ground-level tree injury observations of the model simulation of growth over the 40-year period.



Note: Annual ROG and NO_x emissions data for San Bernardino County were obtained from <u>Alexis et al. (2001a)</u> and the California Air Resource Board's emission inventory available at <u>http://www.arb.ca.gov/html/ds.htm</u> (<u>Cal EPA, 2010</u>). Source: Reprinted with permission of Elsevier Science Ltd. (<u>Lee et al., 2003a</u>).

Figure 9-10 Trends in May to September: 12-hour SUM06, Peak 1-hour ozone concentration and number of daily exceedances of 95 ppb for the Crestline site in 1963 to 1999; in relation to trends in mean daily maximum temperature for Crestline and daily reactive organic gases (ROG) and oxides of nitrogen (NO_X) for San Bernardino County.

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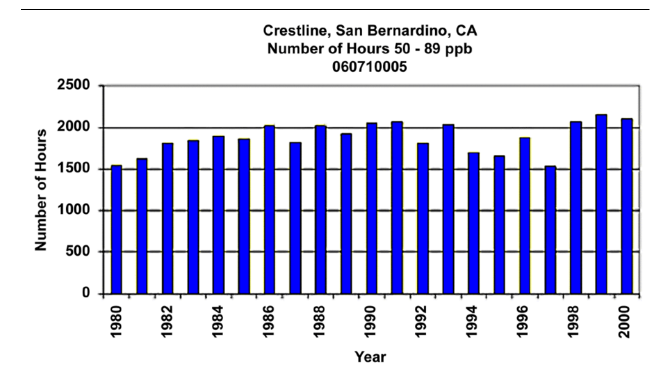


Figure 9-11 The number of hourly average concentrations between 50 and 89 ppb for the period 1980-2000 for the Crestline, San Bernardino County, CA, monitoring site.

9.5.3.2 Diurnal and Seasonal Exposure

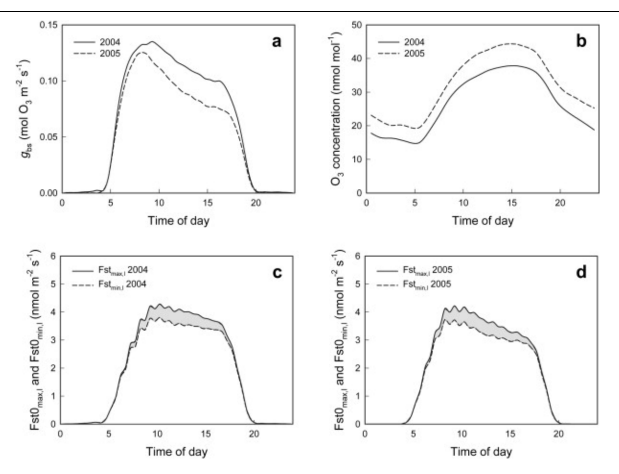
Diurnal Exposure

1	The diurnal patterns of maximal leaf/needle conductance and occurrence of higher
2	ambient concentrations can help determine which hours during the day over a season
3	should be included in an exposure index. Stomatal conductance is species and phenology
4	dependent and is linked to both diurnal and seasonal meteorological activity as well as to
5	soil/site conditions (e.g., VPD, soil moisture). Daily patterns of leaf/needle conductance
6	are often highest in midmorning, whereas higher ambient O ₃ concentrations generally
7	occur in early to late afternoon when stomata are often partially closed and conductances
8	are lower. Total O_3 flux depends on atmospheric and boundary layer resistances, both of
9	which exhibit variability throughout the day. Experimental studies with tree species
10	demonstrated the decoupling of ambient O_3 exposure, peak occurrence, and gas
11	exchange, particularly in areas of drought (Panek, 2004). Several studies have suggested
12	that ponderosa pine trees in the southern and northern Sierra Nevada Mountains may not
13	be as susceptible to high O_3 concentrations as to lower concentrations, due to reduced

1	needle conductance and O ₃ uptake during the period when the highest concentrations
2	occur (Panek et al., 2002; Panek and Goldstein, 2001; Bauer et al., 2000; Arbaugh et al.,
3	<u>1998</u>). Panek et al. (2002) compared direct O_3 flux measurements into a canopy of
4	ponderosa pine and demonstrated a lack of correlation of daily patterns of conductance
5	and O ₃ occurrence, especially in the late season drought period; the authors concluded
6	that a consideration of climate or season was essential, especially considering the role of
7	soil moisture and conductance/uptake. In contrast, Grulke et al. (2002) reported high
8	conductance when O_3 concentrations were high in the same species, but under different
9	growing site conditions. The longer-term biological responses reported by Miller and
10	Rechel (1999) for ponderosa pine in the same region, and the general reduction in recent
11	years in ambient O ₃ concentrations, suggest that stomatal conductance alone may not be a
12	sufficient indicator of potential vegetation injury or damage. Another consideration for
13	the effect of O_3 uptake is the diurnal pattern of detoxification capacity of the plant. The
14	detoxification capacity may not follow the same pattern as stomatal conductance (Heath
15	<u>et al., 2009</u>).
16	The use of a 12-h (8:00 a.m. to 8:00 p.m.) daylight period for a W126 cumulating
17	exposure was based primarily on evidence that the conditions for uptake of O_3 into the
18	plant occur mainly during the daytime hours. In general, plants have the highest stomatal
19	conductance during the daytime and in many areas atmospheric turbulent mixing is
20	greatest during the day as well (Uddling et al., 2010; U.S. EPA, 2006b). However,
21	notable exceptions to maximum daytime conductance are cacti and other plants with
22	crassulacean acid metabolism (CAM photosynthesis) which only open their stomata at
23	night. This section will focus on plants with C3 and C4 photosynthesis, which generally
24	have maximum stomatal conductance during the daytime.
25	Recent reviews of the literature reported that a large number of species had varying
26	degrees of nocturnal stomatal conductance (Caird et al., 2007; Dawson et al., 2007;
27	Musselman and Minnick, 2000). The reason for night-time water loss through stomata is
28	not well understood and is an area of active research (e.g., Christman et al., 2009;
29	<u>Howard et al., 2009</u>). Night-time stomatal opening may be enhanced by O_3 damage that
30	could result in loss of stomatal control, and less complete closure of stomata, than under
31	low O ₃ conditions (<u>Caird et al., 2007;</u> <u>Grulke et al., 2007b</u>). In general, the rate of
32	stomatal conductance at night is much lower than during the day (Caird et al., 2007).
33	Atmospheric turbulence at night is also often low, which results in stable boundary layers
34	and unfavorable conditions for O_3 uptake into vegetation (Finkelstein et al., 2000).
35	Nevertheless, nocturnal turbulence does intermittently occur and may result in
36	non-negligible O ₃ flux into the plants. In addition, plants might be more susceptible to O ₃
37	exposure at night than during the daytime, because of potentially lower plant defenses
38	(Heath et al., 2009; Loreto and Fares, 2007; Musselman et al., 2006; Musselman and

1 Minnick, 2000). For significant nocturnal stomatal flux and O_3 effects to occur, specific 2 conditions must exist. A susceptible plant with nocturnal stomatal conductance and low 3 defenses must be growing in an area with relatively high night-time O₃ concentrations 4 and appreciable nocturnal atmospheric turbulence. It is unclear how many areas there are 5 in the U.S. where these conditions occur. It may be possible that these conditions exist in 6 mountainous areas of southern California, front-range of Colorado (Turnipseed et al., 7 2009) and the Great Smoky Mountains of North Carolina and Tennessee. Tobiessen 8 (1982) found that shade intolerant tree species showed opening of stomata in the dark and 9 did not find this in shade tolerant species. This may indicate shade intolerant trees may be 10 more likely to be susceptible to O_3 exposure at night. More information is needed in 11 locations with high night-time O₃ to assess the local O₃ patterns, micrometeorology and 12 responses of potentially vulnerable plant species.

13 Several field studies have attempted to quantify night-time O_3 uptake with a variety of 14 methods. However, many of these studies have not linked the night-time flux to measured 15 effects on plants. Grulke et al. (2004) showed that the stomatal conductance at night for 16 ponderosa pine in the San Bernardino National Forest (CA) ranged from one tenth to one 17 fourth that of maximum daytime stomatal conductance. In June, at a high-elevation site, it 18 was calculated that 11% of the total daily O_3 uptake of pole-sized trees occurred at night. 19 In late summer, however, O_3 uptake at night was negligible. However, this study did not 20 consider the turbulent conditions at night. Finkelstein et al. (2000) investigated O₃ 21 deposition velocity to forest canopies at three different sites. The authors found the total 22 flux (stomatal and non-stomatal) to the canopy to be very low during night-time hours as 23 compared to day-time hours. However, the authors did note that higher nocturnal 24 deposition velocities at conifer sites may be due to some degree of stomatal opening at 25 night (Finkelstein et al., 2000). Work by Mereu et al. (2009) in Italy on Mediterranean 26 species indicated that nocturnal uptake was from 10 to 18% of total daily uptake during a 27 weak drought and up to 24% as the drought became more pronounced. The proportion of 28 night-time uptake was greater during the drought due to decreases in daytime stomatal 29 conductance (Mereu et al., 2009). In a study conducted in California, (Fares et al., 2011) 30 reported that calculated mean percentages of nocturnal uptake were 5%, 12.5%, 6.9% of total O3 uptake for lemon, mandarin, and orange, respectively. In another recent study at 31 32 the Aspen FACE site in Wisconsin, calculated leaf-level stomatal O₃ flux was near zero 33 from the night-time hours of 8:00 p.m. to 5:00 a.m. (Uddling et al., 2010). This was likely 34 due to low horizontal wind speed (>1 meter/sec) and low O_3 concentrations (<25 ppb) 35 during those same night-time hours (Figure 9-12).



Note: Subscripts "max" and "min" refer to stomatal fluxes calculated neglecting and accounting for potential non-stomatal ozone flux, respectively.

Source: Reprinted with permission of Elsevier Ltd. (Uddling et al., 2010).

Figure 9-12 Diurnal (a) conductance through boundary layer and stomata (gbs), (b) ozone concentration, and leaf-level stomatal ozone flux (Fst0l) in control plots from mid-June through August in (c) 2004 and (d) 2005 in the Aspen FACE experiment.

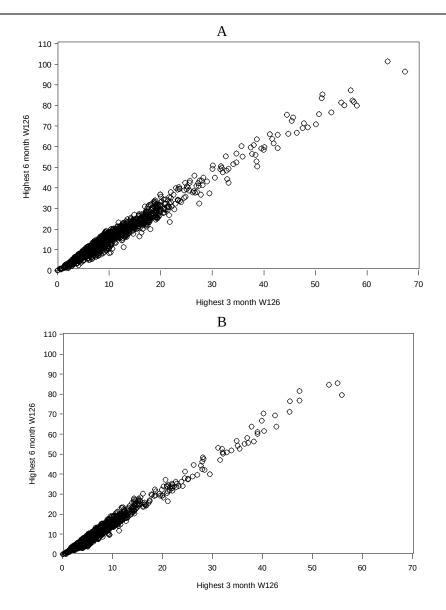
1	A few studies have tested the biological effects of night-time O3 exposure on vegetation
2	in controlled chambers. Biomass of ponderosa pine seedlings was significantly reduced
3	when seedlings were exposed to either daytime or nighttime episodic profiles (Lee and
4	Hogsett, 1999). However, the biomass reductions were much greater with daytime peak
5	concentrations than with nighttime peak concentrations. Similarly, birch cuttings grown
6	in field chambers that were exposed to O_3 at night only, daytime only, and 24 hours
7	showed similar reductions in biomass in night only and day only treatments. Birch
8	seedling showed greater reductions in growth in 24-h exposures than those exposed to O_3
9	at night or day only (Matyssek et al., 1995). Field mustard (Brassica rapa) plants
10	exposed to O_3 during the day or night showed little significant difference in the amounts

1of injury or reduced growth response to O_3 treatment, although the stomatal conductance2was 70-80% lower at night (Winner et al., 1989). These studies show that effects can be3seen with night-time exposures to O_3 but when atmospheric conditions are stable at night,4it is uncertain how these exposures may affect plants and trees with complex canopies in5the field.

Seasonal exposure

- 6 Vegetation across the U.S. has widely varying periods of physiological activity during the 7 year due to variability in climate and phenology. In order for a particular plant to be 8 vulnerable to O_3 pollution, it must have foliage and be physiologically active. Annual 9 crops are typically grown for periods of two to three months. In contrast, perennial 10 species may be photosynthetically active longer (up to 12 months each year for some 11 species) depending on the species and where it is grown. In general, the period of 12 maximum physiological activity and thus, potential O_3 uptake for vegetation coincides 13 with some or all of the intra-annual period defined as the O_3 season, which varies on a 14 state-by-state basis (Figure 3-24). This is because the high temperature and high light 15 conditions that typically promote the formation of tropospheric O_3 also promote 16 physiological activity in vegetation. There are very limited exceptions to this pattern 17 where O_3 can form in the winter in areas in the western U.S. with intense natural gas 18 exploration (Pinto, 2009), but this is typically when plants are dormant and there is little 19 chance of O₃ uptake. Given the significant variability in growth patterns and lengths of 20 growing season among the wide range of vegetation species that may experience adverse 21 effects associated with O₃ exposure, no single time window of exposure can work 22 perfectly for all types of vegetation.
- 23 Various intra-annual averaging and accumulation time periods have been considered for 24 the protection of vegetation. The 2007 proposal for the secondary O_3 standard (75 FR 25 37818) proposed to use the maximum consecutive 3-month period within the O₃ season. 26 The U.S. Forest Service and federal land managers have used a 24-h W126 accumulated 27 for 6 months from April through September (U.S. Forest Service, 2000). However, some 28 monitors in the U.S. are operational for as little as four months and would not have 29 enough data for a 6-month seasonal window. The exposure period in the vast majority of 30 O₃ exposure studies conducted in the U.S. has been much shorter than 6 months. Most of 31 the crop studies done through NCLAN had exposures less than three months with an 32 average of 77 days. Open-top chamber studies of tree seedlings, compiled by the EPA, 33 had an average exposure of just over three months or 99 days. In more recent FACE 34 experiments, SoyFACE exposed soybeans for an average of approximately 120 days per 35 year and the Aspen FACE experiment exposed trees to an average of approximately 36 145 days per year of elevated O_3 , which included the entire growing season at those

particular sites. Despite the possibility that plants may be exposed to ambient O_3 longer than 3 months in some locations, there is generally a lack of exposure experiments conducted for longer than 3 months.



Note: Data are from the AQS and CASTNET monitors for the years 2008 and 2009. (A) W126, 3 month versus 6 month, 2008 (Pearson correlation = 0.99); (B) W126, 3 month versus 6 month, 2009 (Pearson correlation = 0.99).

Figure 9-13 Maximum 3-month, 12-h W126 plotted against maximum 6-month, 12-h W126.

1

2

3

1	In an analysis of the 3- and 6-month maximum W126 values calculated for over 1,200
2	AQS (Air Quality System) and CASTNET (Clean Air Status and Trend Network) EPA
3	monitoring sites for the years 2008-2009, it was found that these 2 accumulation periods
4	resulted in highly correlated metrics (Figure 9-13). The two accumulation periods were
5	centered on the yearly maximum for each monitoring site, and it is possible that this
6	correlation would be weaker if the two periods were not temporally aligned. In the U.S.,
7	W126 cumulated over 3 months, and W126 cumulated over 6 months are proxies of one
8	another, as long as the period in which daily W126 is accumulated corresponds to the
9	seasonal maximum. Therefore, it is expected that either statistic will predict vegetation
10	response equally well. In other words, the strength of the correlation between maximum
11	3-month W126 and maximum 6-month W126 is such that there is no material difference
12	in their predictive value for vegetation response.

9.5.4 Ozone Uptake/Dose Modeling for Vegetation

13 Another approach for improving risk assessment of vegetation response to ambient O_3 is 14 based on estimating the O_3 concentration from the atmosphere that enters the leaf 15 (i.e., flux or deposition). Interest has been increasing in recent years, particularly in 16 Europe, in using mathematically tractable flux models for O_3 assessments at the regional, 17 national, and European scale (Matyssek et al., 2008; Paoletti and Manning, 2007; ICP 18 M&M, 2004; Emberson et al., 2000b; Emberson et al., 2000a). Some researchers have 19 claimed that using flux models can be used to better predict vegetation responses to O_3 20 than exposure-based approaches (Matyssek et al., 2008). However, other research has 21 suggested that flux models do not predict vegetation responses to O₃ better than 22 exposure-based models, such as AOT40 (Gonzalez-Fernandez et al., 2010). While some 23 efforts have been made in the U.S. to calculate O₃ flux into leaves and canopies (Fares et 24 al., 2010a; Turnipseed et al., 2009; Uddling et al., 2009; Bergweiler et al., 2008; Hogg et al., 2007; Grulke et al., 2004; Grantz et al., 1997; Grantz et al., 1995), little information 25 26 has been published relating these fluxes to effects on vegetation. The lack of flux data in 27 the U.S. and the lack of understanding of detoxification processes have made this 28 technique less viable for vulnerability and risk assessments in the U.S. 29 Flux calculations are data intensive and must be carefully implemented. Reducing 30 uncertainties in flux estimates for areas with diverse surface or terrain conditions to 31 within \pm 50% requires "very careful application of dry deposition models, some model 32 development, and support by experimental observations" (Wesely and Hicks, 2000). As 33

an example, the annual average deposition velocity of O_3 among three nearby sites in similar vegetation was found to vary by $\pm 10\%$, presumably due to terrain (Brook et al., Moreover, the authors stated that the actual variation was even greater, because

1	stomatal uptake was unrealistically assumed to be the same among all sites, and flux is
2	strongly influenced by stomatal conductance (Brook et al., 1997; Massman and Grantz,
3	<u>1995; Fuentes et al., 1992; Reich, 1987; Leuning et al., 1979</u>). This uptake-based
4	approach to quantify the vegetation impact of O_3 requires inclusion of those factors that
5	control the diurnal and seasonal O_3 flux to vegetation (e.g., climate patterns, species
6	and/or vegetation-type factors and site-specific factors). The models have to distinguish
7	between stomatal and non-stomatal components of O_3 deposition to adequately estimate
8	actual concentration reaching the target tissue of a plant to elicit a response (Uddling et
9	<u>al., 2009</u>). Determining this O_3 uptake via canopy and stomatal conductance relies on
10	models to predict flux and ultimately the "effective" flux (Grunhage et al., 2004;
11	Massman, 2004; Massman et al., 2000). "Effective flux" has been defined as the balance
12	between O_3 flux and detoxification processes (Heath et al., 2009; Musselman and
13	Massman, 1999; Grunhage and Haenel, 1997; Dammgen et al., 1993). The
14	time-integrated "effective flux" is termed "effective dose." The uptake mechanisms and
15	the resistances in this process, including stomatal conductance and biochemical defense
16	mechanisms, are discussed below. The flux-based index is the goal for the "Level II"
17	critical level for assessment of O_3 risk to vegetation and ecosystems across Europe
18	(<u>Ashmore et al., 2004a</u>).
19	An important consideration in both O ₃ exposure and uptake is how the O ₃ concentration
20	at the top of low vegetation such as, crops and tree seedlings may be lower than the
21	height at which the measurement is taken. Ambient monitor inlets in the U.S. are
22	typically at heights of 3 to 5 meters. During daytime hours, the vertical O_3 gradient can
23	be relatively small because turbulent mixing maintains the downward flux of O ₃ . For
24	example, Horvath et al. (1995) calculated a 7% decrease in O_3 going from a height of 4
25	meters down to 0.5 meters above the surface during unstable (or turbulent) conditions in
26	a study over low vegetation in Hungary [see Section AX3.3.2. of the 2006 O3 AQCD
27	(U.S. EPA, 2006b)]. There have been several studies indicating decreased O_3
28	concentrations under tree canopies (Kolb et al., 1997; Samuelson and Kelly, 1997; Joss
29	and Graber, 1996; Fredericksen et al., 1995; Lorenzini and Nali, 1995; Enders, 1992;
30	Fontan et al., 1992; Neufeld et al., 1992). In contrast, for forests, measured data may
31	underestimate O ₃ concentration at the top of the canopy. The difference between
32	measurement height and canopy height is a function of several factors, the intensity of
33	turbulent mixing in the surface layer and other meteorological factors, canopy height and
34	total deposition to the canopy. Some researchers have used deposition models to estimate
35	O_3 concentration at canopy-top height based on concentrations at measurement height
36	(Emberson et al., 2000a). However, deposition models usually require meteorological
37	data inputs that are not always available or well characterized across large geographical
38	scales.

1	Soil moisture is a critical factor in controlling O ₃ uptake through its effect on plant water
2	status and stomatal conductance. In an attempt to relate uptake, soil moisture, and
3	ambient air quality to identify areas of potential risk, available O3 monitoring data for
4	1983 to 1990 were used along with literature-based seedling exposure-response data from
5	regions within the southern Appalachian Mountains that might have experienced O_3
6	exposures sufficient to inhibit growth (Lefohn et al., 1997). In a small number of areas
7	within the region, O ₃ exposures and soil moisture availability were sufficient to possibly
8	cause growth reductions in some O ₃ sensitive species (e.g., black cherry). The
9	conclusions were limited, however, because of the uncertainty in interpolating O_3
10	exposures in many of the areas and because the hydrologic index used might not reflect
11	actual water stress.
12	The non-stomatal component of plant defenses are the most difficult to quantify, but
13	some studies are available (Heath et al., 2009; Barnes et al., 2002; Plochl et al., 2000;
14	
	Chen et al., 1998; Massman and Grantz, 1995). Massman et al. (2000) developed a
15	<u>Chen et al., 1998; Massman and Grantz, 1995)</u> . <u>Massman et al. (2000)</u> developed a conceptual model of a dose-based index to determine how plant injury response to O_3
15	conceptual model of a dose-based index to determine how plant injury response to O ₃
15 16	conceptual model of a dose-based index to determine how plant injury response to O_3 relates to the traditional exposure-based parameters. The index used time-varying-
15 16 17	conceptual model of a dose-based index to determine how plant injury response to O_3 relates to the traditional exposure-based parameters. The index used time-varying-weighted fluxes to account for the fact that flux was not necessarily correlated with plant
15 16 17 18	conceptual model of a dose-based index to determine how plant injury response to O_3 relates to the traditional exposure-based parameters. The index used time-varying-weighted fluxes to account for the fact that flux was not necessarily correlated with plant injury or damage. The model applied only to plant foliar injury and suggested that
15 16 17 18 19	conceptual model of a dose-based index to determine how plant injury response to O_3 relates to the traditional exposure-based parameters. The index used time-varying-weighted fluxes to account for the fact that flux was not necessarily correlated with plant injury or damage. The model applied only to plant foliar injury and suggested that application of flux-based models for determining plant damage (yield or biomass) would

9.5.5 Summary

22	Exposure indices are metrics that quantify exposure as it relates to measured plant
23	damage (i.e., reduced growth). They are summary measures of monitored ambient O_3
24	concentrations over time intended to provide a consistent metric for reviewing and
25	comparing exposure-response effects obtained from various studies. No recent
26	information is available since 2006 that alters the basic conclusions put forth in the 2006
27	and 1996 O_3 AQCDs. These AQCDs focused on the research used to develop various
28	exposure indices to help quantify effects on growth and yield in crops, perennials, and
29	trees (primarily seedlings). The performance of indices was compared through regression
30	analyses of earlier studies designed to support the estimation of predictive O3 exposure-
31	response models for growth and/or yield of crops and tree (seedling) species.
32	Another approach for improving risk assessment of vegetation response to ambient O_3 is
33	based on determining the O ₃ concentration from the atmosphere that enters the leaf
34	(i.e., flux or deposition). Interest has been increasing in recent years, particularly in

1	Europe, in using mathematically tractable flux models for O ₃ assessments at the regional,
2	national, and European scale (Matyssek et al., 2008; Paoletti and Manning, 2007; ICP
3	M&M, 2004; Emberson et al., 2000b; Emberson et al., 2000a). While some efforts have
4	been made in the U.S. to calculate O ₃ flux into leaves and canopies (Turnipseed et al.,
5	2009; Uddling et al., 2009; Bergweiler et al., 2008; Hogg et al., 2007; Grulke et al., 2004;
6	Grantz et al., 1997; Grantz et al., 1995), little information has been published relating
7	these fluxes to effects on vegetation. There is also concern that not all O ₃ stomatal uptake
8	results in a yield reduction, which depends to some degree on the amount of internal
9	detoxification occurring with each particular species. Those species having high amounts
10	of detoxification potential may, in fact, show little relationship between O ₃ stomatal
11	uptake and plant response (Musselman and Massman, 1999). The lack of data in the U.S.
12	and the lack of understanding of detoxification processes have made this technique less
13	viable for vulnerability and risk assessments in the U.S.
14	The main conclusions from the 1996 and 2006 O3 AQCDs regarding indices based on
15	ambient exposure are still valid. These key conclusions can be restated as follows:
16	 O₃ effects in plants are cumulative;
17	 higher O₃ concentrations appear to be more important than lower
18	concentrations in eliciting a response;
19	 plant sensitivity to O₃ varies with time of day and plant development stage;
20	and
21	 quantifying exposure with indices that accumulate the O₃ hourly
22	concentrations and preferentially weight the higher concentrations improves
23	the explanatory power of exposure/response models for growth and yield, over
24	using indices based on mean and peak exposure values.
25	Various weighting functions have been used, including threshold-weighted
26	(e.g., SUM06) and continuous sigmoid-weighted (e.g., W126) functions. Based on
27	statistical goodness-of-fit tests, these cumulative, concentration-weighted indices could
28	not be differentiated from one another using data from previous exposure studies.
29	Additional statistical forms for O_3 exposure indices have been discussed in Lee et al.
30	(1988b). The majority of studies published since the 2006 O_3 AQCD do not change
31	earlier conclusions, including the importance of peak concentrations, and the duration
32	and occurrence of O_3 exposures in altering plant growth and yield.
33	Given the current state of knowledge and the best available data, exposure indices that
34	cumulate and differentially weight the higher hourly average concentrations and also
35	include the "mid-level" values continue to offer the most defensible approach for use in

developing response functions and comparing studies, as well as for defining future indices for vegetation protection.

9.6 Ozone Exposure-Plant Response Relationships

9.6.1 Introduction

1

3	The adequate characterization of the effects of O ₃ on plants for the purpose of setting air
4	quality standards is contingent not only on the choice of the index used (i.e., SUM06,
5	W126) to summarize O_3 concentrations (Section <u>9.5</u>), but also on quantifying the
6	response of the plant variables of interest at specific values of the selected index. The
7	many factors that determine the response of plants to O3 exposure have been discussed in
8	previous sections. They include species, genotype and other genetic characteristics
9	(Section 9.3), biochemical and physiological status (Section 9.3), previous and current
10	exposure to other stressors (Section $9.4.8$), and characteristics of the exposure itself
11	(Section 9.5). Establishing a secondary air quality standard entails the capability to
12	generalize those observations, in order to obtain predictions that are reliable enough
13	under a broad variety of conditions, taking into account these factors. This section
14	reviews results that have related specific quantitative observations of O_3 exposure with
15	quantitative observations of plant responses, and the predictions of responses that have
16	been derived from those observations through empirical models.
17	For four decades, exposure to O_3 at ambient concentrations found in many areas of the
18	U.S. has been known to cause detrimental effects in plants (U.S. EPA, 2006b, 1996b,
19	<u>1984</u> , <u>1978a</u>). Results published after the 2006 O_3 AQCD continue to support this
20	finding, and the following sections deal with the quantitative characterizations of the
21	relationship, and what new insights may have appeared since 2006. Detrimental effects
22	on plants include visible injury, decreases in the rate of photosynthesis, reduced growth,
23	and reduced yield of marketable plant parts. Most published exposure-response data have
24	reported O_3 effects on the yield of crops and the growth of tree seedlings, and those two
25	variables have been the focus of the characterization of ecological impacts of O_3 for the
26	purpose of setting secondary air quality standards. In order to support quantitative
27	modeling of exposure-response relationships, data should preferably include more than
28	three levels of exposure, and some control of potential confounding or interacting factors
29	should be present in order to model the relationship with sufficient accuracy. Letting
30	potential confounders, such as other stressors, vary freely when generating O_3 exposure-
31	response data might improve the 'realism' of the data, but it also greatly increases the
32	amount of data necessary to extract a clear quantitative description of the relationship.

- 1 Conversely however, experimental settings should not be so exhaustively restrictive as to 2 make generalization outside of them problematic. During the last four decades, many of 3 the studies of the effects of O_3 on growth and yield of plants have not included enough 4 levels of O_3 to parameterize more than the simplest linear model. The majority of these 5 studies have only contrasted two levels, ambient and elevated, or sometimes three by 6 adding carbon filtration in OTC studies, with little or no consideration of quantitatively 7 relating specific values of exposure to specific values of growth or yield. This is not to 8 say that studies that did not include more than two or three levels of O₃ exposure, or 9 studies that were conducted in uncontrolled environments, do not provide exposure-10 response information that is highly relevant to reviewing air quality standards. In fact, 11 they can be essential in verifying the agreement between predictions obtained through the 12 empirical models derived from experiments such as NCLAN, and observations. The 13 consensus of model predictions and observations from a variety of studies conducted in 14 other locations, at other times, and using different exposure methods, greatly increases 15 confidence in the reliability of both. Furthermore, if they are considered in the aggregate, 16 studies with few levels of exposure or high unaccounted variability can provide 17 additional independent estimates of decrements in plant growth and yield, at least within 18 a few broad categories of exposure.
- 19 Extensive exposure-response information on a wide variety of plant species has been 20 produced by two long-term projects that were designed with the explicit aim of obtaining 21 quantitative characterizations of the response of such an assortment of crop plants and 22 tree seedlings to O₃ under North American conditions: the NCLAN project for crops, and 23 the EPA National Health and Environmental Effects Research Laboratory, Western 24 Ecology Division tree seedling project (NHEERL/WED). The NCLAN project was 25 initiated by the EPA in 1980 primarily to improve estimates of yield loss under field 26 conditions and to estimate the magnitude of crop losses caused by O_3 throughout the U.S. 27 (Heck et al., 1991; 1982). The cultural conditions used in the NCLAN studies 28 approximated typical agronomic practices, and the primary objectives were: (1) to define 29 relationships between yields of major agricultural crops and O_3 exposure as required to 30 provide data necessary for economic assessments and development of O_3 NAAQS; (2) to 31 assess the national economic consequences resulting from O_3 exposure of major 32 agricultural crops; and (3) to advance understanding of cause-and-effect relationships that 33 determine crop responses to pollutant exposures.
- 34NCLAN experiments yielded 54 exposure-response curves for 12 crop species, some of35which were represented by multiple cultivars at several of 6 locations throughout the U.S.36The NHEERL/WED project was initiated by EPA in 1988 with similar objectives for tree37species, and yielded 49 exposure-responses curves for multiple genotypes of 11 tree38species grown for up to three years in Oregon, Michigan, and the Great Smoky

1	Mountains National Park. Both projects used OTCs to expose plants to three to five
2	levels of O ₃ . Eight of the 54 crop datasets were from plants grown under a combination
3	of O_3 exposure and experimental drought conditions. <u>Figure 9-14</u> through <u>Figure 9-17</u>
4	summarize some of the NCLAN and NHEERL/WED results.
5	It should be noted that data from FACE experiments might also be used for modeling
6	exposure-response. They only use two levels of O_3 (ambient concentration at the site and
7	a multiple of it), but given that the value of both levels of exposure changes every year,
8	and that they are typically run for many consecutive years, aggregating data over time
9	produces twice as many levels of O_3 as there are years. As described in Section <u>9.2.4</u> ,
10	FACE experiments seek to impose fewer constraints on the growth environment than
11	OTCs. As a consequence, FACE studies have to contend with larger variability,
12	especially year-to-year variability, but the difference in experimental conditions between
13	the two methodologies makes comparisons between their results especially useful.
14	Growth and yield of at least one crop (soybean) has been investigated in yearly
15	experiments since 2001 at a FACE facility in Illinois (UIUC, 2010; Morgan et al., 2006).
16	However, almost all analyses of SoyFACE published so far have been based on subsets
17	of one or two years, and have only contrasted ambient versus elevated O_3 as categorical
18	variables. They have not modeled the response of growth and yield to O_3 exposure
19	continuously over the range of exposure values that have occurred over time. The only
20	exception is a study by Betzelberger et al. (2010), who used a linear regression model on
21	data pooled over 2 years. Likewise, trees of three species (trembling aspen, paper birch,
22	and sugar maple) were grown between 1998 and 2009 in a FACE experiment located in
23	Rhinelander, Wisconsin (Pregitzer et al., 2008; Dickson et al., 2000). The Aspen FACE
24	experiment has provided extensive data on responses of trees beyond the seedling stage
25	under long-term exposure, and also on ecosystem-level responses (Section 9.4), but the
26	only attempt to use those data in a continuous model of the response of tree growth to O_3
27	exposure (Percy et al., 2007) suffered from severe methodological problems, some of
28	which are discussed in Section <u>9.6.3</u> . Finally, one experiment was able to exploit a
29	naturally occurring gradient of O3 concentrations to fit a linear regression model to the
30	growth of cottonwood (Gregg et al., 2006, 2003). Factors such as genotype, soil type and
31	soil moisture were under experimental control, and the authors were able to partition out
32	the effects of potential confounders such as temperature, atmospheric N deposition, and
33	ambient CO_2 .
34	A serious difficulty in assessing results of exposure-response research is the multiplicity
35	of O_3 metrics that have been used in reporting. As described in Section <u>9.5</u> , metrics that
36	entail either weighting or thresholding of hourly values cannot be algebraically converted
27	interpretation entitle entrepretation of the second se

into one another, or into unweighted metrics such as hourly average. When computing O_3

1	exposure using weighted or thresholded metrics, each metric has to be computed
2	separately from the original hourly data. Comparisons of exposure-response models can
3	only be made between studies that used the same metric, and the value of exposure at
4	which a given plant response is expected using one metric of exposure cannot be exactly
5	converted to another metric. Determining the exposure value at which an effect would be
6	observed in a different metric can only be accomplished by first computing the
7	experimental exposures in this metric from the hourly data, then estimating (fitting)
8	model coefficients again. This problem is irremediable, although useful comparisons
9	might be made using categorical exposures such as 'current ambient exposure' or '2050
10	projected exposure', which can serve as a common reference for quantitative values
11	expressed in various metrics. Studies that contained growth or yield exposure-response
12	data at few levels of exposure, and/or using metrics other than W126 are summarized in
13	<u>Table 9-18</u> and <u>Table 9-19</u> .

9.6.2 Estimates of Crop Yield Loss and Tree Seedling Biomass Loss in the 1996 and 2006 Ozone AQCDs

- 14 The 1996 and 2006 O₃ AQCDs relied extensively on analyses of NCLAN and
 - NHEERL/WED by Lee et al. (1994); (1989, 1988b, 1987), Hogsett et al. (1997), Lee and Hogsett (1999), Heck et al. (1984), (Rawlings and Cure, 1985), (Lesser et al., 1990), and (Gumpertz and Rawlings, 1992). Those analyses concluded that a three-parameter Weibull model –

$$Y = \alpha \, e^{-\left(\frac{W126}{\eta}\right)^{\beta}}$$

Equation 9-2

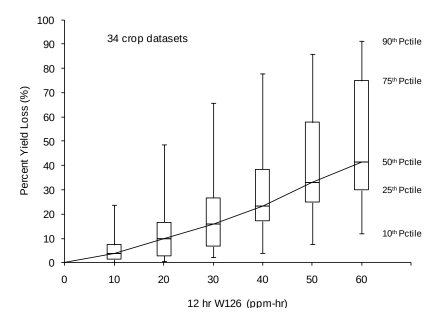
19 is the most appropriate model for the response of absolute yield and growth to O₃ 20 exposure, because of the interpretability of its parameters, its flexibility (given the small 21 number of parameters), and its tractability for estimation. In addition, removing the 22 intercept α results in a model of relative yield (yield relative to [yield at exposure=0]) 23 without any further reparameterization. Formulating the model in terms of relative yield 24 or relative yield loss (yield loss=[1 – relative yield]) is essential in comparing exposure-25 response across species, genotypes, or experiments for which absolute values of the 26 response may vary greatly. In the 1996 and 2006 O₃ AQCDs, the two-parameter model of 27 relative yield was used in deriving common models for multiple species, multiple 28 genotypes within species, and multiple locations.

15

16

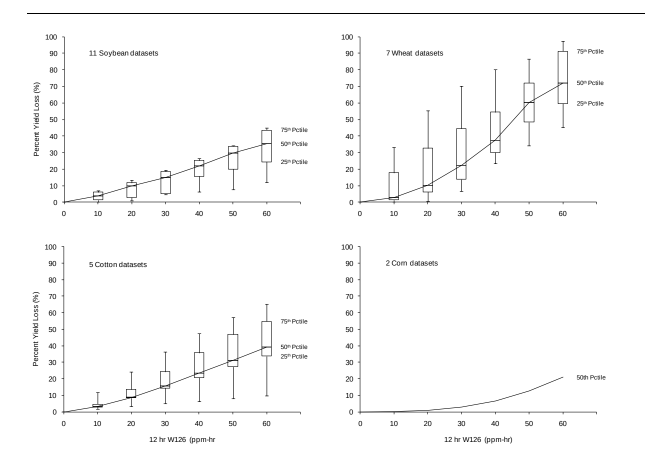
17

1	
1	Given the disparate species, genotypes, and locations that were included in the NCLAN
2	and NHEERL/WED projects, and in the absence of plausible distributional assumptions
3	with respect to those variables, a three step process using robust methods was used to
4	obtain parameter estimates that could be generalized. The models that were derived for
5	each species or group of species were referred to as median composite functions. In the
6	first step, the three parameters of the Weibull model were computed for absolute yield or
7	biomass data from each NCLAN and NHEERL/WED experiment (54 crop datasets and
8	49 tree seedling datasets), using nonlinear regression. When data were only available for
9	three levels of exposure because of experimental problems, the shape parameter β was
10	constrained to 1, reducing the model to an exponential decay model. In the second step, α
11	was dropped, and predicted values of relative yield or biomass were then computed for
12	12-hour W126 exposures between 0 and 60 ppm-h. At each of these W126 exposure
13	values, the 25th, 50th, and 75th percentiles of the response were identified among the
14	predicted curves of relative response. For example, for the 34 NCLAN studies of 12 crop
15	species grown under non-droughted conditions for a complete cropping cycle
16	(Figure 9-14), the 3 quartiles of the response were identified at every integer value of
17	W126 between 0 and 60. The third step fitted a two-parameter Weibull model to those
18	percentiles, yielding the median composite function for the relative yield or biomass
19	response to O ₃ exposure for each grouping of interest (e.g., all crops, all trees, all datasets
20	for one species), as well as composite functions for the other quartiles. In the 1996 and
21	2006 O_3 AQCDs this modeling of crop yield loss and tree seedling biomass loss was
22	conducted using the SUM06 metric for exposure. This section updates those results by
23	using the 12-hour W126 as proposed in 2007 (72 FR 37818) and 2010 (75 FR 2938, page
24	3003). Figure 9-14 through Figure 9-17 present quantiles of predicted relative yield or
25	biomass loss at seven values of the 12-h W126 for some representative groupings of
26	NCLAN and NHEERL/WED results. <u>Table 9-9</u> through <u>Table 9-11</u> give the 90-day 12-h
27	W126 O ₃ exposure values at which 10 and 20% yield or biomass losses are predicted in
28	50 and 75% of crop or tree species using the composite functions.



Note: Quantiles of the predicted relative yield loss at 7 values of 12-hour W126 for 34 Weibull curves estimated using nonlinear regression on data from 34 studies of 12 crop species grown under well-watered conditions for the full duration of 1 cropping cycle. Source of Weibull parameters: Lee and Hogsett (1996).

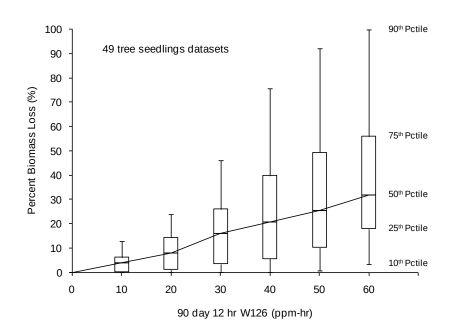
Figure 9-14 Quantiles of predicted relative yield loss for 34 NCLAN crop experiments.



Notes: Quantiles of the predicted relative yield loss at 7 values of 12-h W126 for Weibull curves estimated using nonlinear regression for 4 species grown under well-watered conditions for the full duration of 1 cropping cycle. The number of studies available for each species is indicated on each plot.

Source of Weibull parameters: Lee and Hogsett (1996).

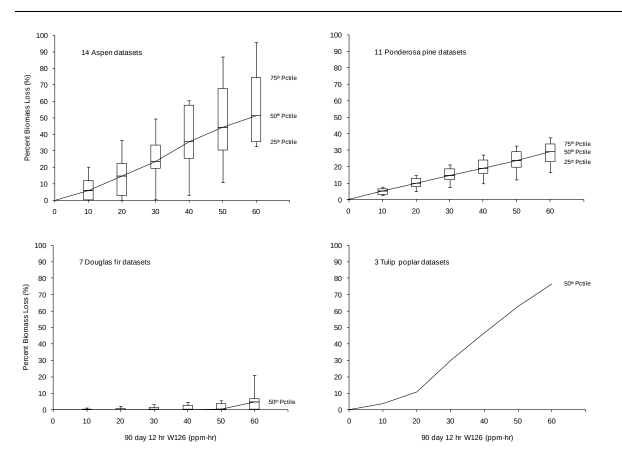
Figure 9-15 Quantiles of predicted relative yield loss for 4 crop species in NCLAN experiments.



Note: Quantiles of the predicted relative above-ground biomass loss at 7 values of 12-h W126 for 49 Weibull curves estimated using nonlinear regression on data from 49 studies of 11 tree species grown under well-watered conditions for 1 or 2 years. Curves were standardized to 90-day W126.

Source of Weibull parameters: Lee and Hogsett (1996).

Figure 9-16 Quantiles of predicted relative biomass loss for 49 tree species in NHEERL/WED experiments.



Note: Quantiles of the predicted relative above-ground biomass loss at 7 exposure values of 12-h W126 for Weibull curves estimated using nonlinear regression on data for 4 tree species grown under well-watered conditions for 1 or 2 year. Curves were standardized to 90-day W126. The number of studies available for each species is indicated on each plot. Source of Weibull parameters: Lee and Hogsett (1996).

Figure 9-17 Quantiles of predicted relative biomass loss for 4 tree species in NHEERL/WED experiments.

Table 9-9Ozone exposures at which 10 and 20% yield loss is predicted for 50
and 75% of crop species.

Predited Yield Loss for Crop Species ^a	90-day 12-h W126 for 10% yield loss (ppm-h)	90-day 12-h W126 for 20% yield loss (ppm-h)
Model for the 50th Percentile of 34 curves		
Relative yield=exp(-(W126/104.82)**1.424)	22	37
Model for the 75th Percentile of 34 curves		
Relative yield=exp(-(W126/78.12)**1.415)	16	27

^aBased on composite functions for the 50th and 75th percentiles of 34 Weibull curves for relative yield loss data from 34 nondroughted NCLAN studies of 12 crop species; curves were standardized to 90-day W126. Source of parameters for the 34 curves: Lee and Hogsett (1996).

Table 9-10Ozone exposures at which 10 and 20% yield loss is predicted for 50
and 75% of crop species (Droughted versus Watered conditions).

Predicted Yield Loss for Crop Species ^a		90 day 12-h W126 for 10% yield loss (ppm-h)	90 day 12-h W126 for 20% yield loss (ppm-h)
Model for the	e 50th Percentile of 2×8 curves		
Watered	Relative yield=exp(-(W126/132.86)**1.170)	19	37
Droughted	Relative yield=exp(-(W126/179.84)**1.713)	48	75
Model for the	e 75th Percentile of 2×8 curves		
Watered	Relative yield=exp(-(W126/90.43)**1.310)	16	29
Droughted	Relative yield=exp(-(W126/105.16)**1.833)	31	46

^aUnder drought conditions and adequate moisture based on composite functions for the 50th and 75th percentiles of 16 Weibull curves for relative yield loss data from 8 NCLAN studies that paired droughted and watered conditions for the same genotype; curves were standardized to 90-day W126.

Source of parameters for the 16 curves: Lee and Hogsett (1996).

Table 9-11Ozone exposures at which 10 and 20% biomass loss is predicted
for 50 and 75% of tree species.

Predicted Biomass Loss for Tree Species ^a	90 day 12 h W126 for 10% yield loss (ppm-h)	90 day 12 h W126 for 20% yield loss (ppm-h)
Model for the 50th Percentile of 49 curves		
Relative yield=exp(-(W126/131.57)**1.242)	21	39
Model for the 75th Percentile of 49 curves		
Relative yield=exp(-(W126/65.49)**1.500)	15	24

^aBased on composite functions for the 50th and 75th percentiles of 49 Weibull curves for relative above-ground biomass loss data from 49 studies of 11 tree species grown under well-watered conditions for 1 or 2 year; curves were standardized to 90-day W126. Source of parameters for the 49 curves: <u>Lee and Hogsett (1996</u>).

9.6.3 Validation of 1996 and 2006 Ozone AQCD Models and Methodology Using the 90 day 12-h W126 and Current FACE Data

1	Since the completion of the NCLAN and NHEERL/WED projects, almost no studies
2	have been published that could provide a basis for estimates of exposure-response that
3	can be compared to those of the 1996 and 2006 O ₃ AQCDs. Most experiments, regardless
4	of exposure methodology, include only two levels of exposure. In addition, very few
5	studies have included measurements of exposure using the W126 metric, or the hourly O_3
6	concentration data that would allow computing exposure using the W126. Two FACE
7	projects, however, were conducted over multiple years, and by adding to the number of
8	exposure levels over time, can support independent model estimation and prediction
9	using the same model and the same robust process as summarized in Section 9.6.2.
10	Hourly O ₃ data were available from both FACE projects.
11	The SoyFACE project is situated near Champaign, IL, and comprises 32 octagonal rings
12	(20m-diameter), 4 of which in a given year are exposed to ambient conditions, and 4 of
13	which are exposed to elevated O_3 as a fixed proportion of the instantaneous ambient
14	concentration (Betzelberger et al., 2010; UIUC, 2010; Morgan et al., 2006; Morgan et al.,
15	2004). Since 2002, yield data have been collected for up to 8 genotypes of soybean
16	grown in subplots within each ring. The Aspen FACE project is situated in Rhinelander,
17	WI, and comprises 12 rings (30m-diameter), 3 of which are exposed to ambient
18	conditions, and 3 of which are exposed to O_3 as a fixed proportion of the instantaneous
19	ambient concentration (Pregitzer et al., 2008; Karnosky et al., 2005; Dickson et al.,
20	2000). In the summer of 1997, half the area of each ring was planted with small (five to
21	seven leaf sized) clonally propagated plants of five genotypes of trembling aspen, which
22	were left to grow in those environments until 2009. Biomass data are currently available

- 1 for the years 1997-2005 (King et al., 2005). Ozone exposure in these two FACE projects 2 can be viewed as a categorical variable with two levels: ambient, and elevated. However, 3 this overlooks the facts that not only do both ambient and elevated exposure vary from 4 year to year, but the proportionality between them also changes yearly. This change has 5 two sources: first, the dispensing of O_3 into the elevated exposure rings varies from the 6 set point for the ambient/elevated proportionality to some extent, and for SoyFACE, the 7 set point changed between years. Second, when using threshold or concentration-8 weighted cumulative metrics (such as AOT40, SUM06 or W126), the proportionality 9 does not propagate regularly from the hourly data to the yearly value. For example, 10 hourly average elevated exposures that are a constant 1.5 times greater than ambient do 11 not result in AOT40, SUM06 or W126 values that are some constant multiple of the 12 ambient values of those indices. Depending on the fraction of hourly values that are 13 above the threshold or heavily weighted, the same average yearly exposure will result in 14 different exposure values when using thresholded or weighted metrics. In some years, 15 elevated exposures in FACE experiments experience many more values above the 16 threshold, or more heavily weighted than the ambient exposures; thus in those years, the 17 distance between ambient and elevated exposure values increases relative to other years. 18 As a consequence, the number of exposure levels in multi-year experiments is twice the 19 number of years. In the case of SoyFACE for the period between 2002 and 2008, ambient 20 exposure in the highest year was approximately equal to elevated exposure in the lowest 21 year, with 14 levels of O₃ exposure evenly distributed from lowest to highest. The 22 particular conditions of the Aspen FACE experiment resulted in 12 exposure levels 23 between 1998 and 2003, but they were not as evenly distributed between minimum and 24 maximum over the 6-year period. 25 There are necessary differences in the modeling of exposure-response in annual plants 26 such as soybean, and in perennial plants such as aspen trees, when exposure takes place over multiple years. In annual plants, responses recorded at the end of the life cycle,
- 27 28 i.e., yearly, are analyzed in relationship to that year's exposure. Yield of soybeans is 29 affected by exposure during the year the crop was growing, and a new crop is planted 30 every year. Thus an exposure-response relationship can be modeled from yearly 31 responses matched to yearly exposures, with those exposure-response data points having 32 been generated in separate years. For perennial organisms, which are not harvested yearly 33 and continue to grow from year to year, such pairing of exposure and response cannot be 34 done without accounting for time. Not only does the size of the organism at the beginning 35 of each year of exposure increase, but size is also dependent on the exposure from 36 previous years. Therefore the relationship of response and exposure must be analyzed 37 either one year at a time, or by standardizing the response as a yearly increment relative 38 to size at the beginning of each year. Furthermore, the relevant measurement of exposure 39 is cumulative, or cumulative yearly average exposure, starting in the year exposure was

1 initiated, up to the end of the year of interest. When analyzing the growth of trees over 2 several years, it would be evidently incorrect to pair the exposure level in every discrete 3 year with absolute size of the trees that year, and posit a direct relationship between them, 4 without taking increasing age into consideration. In the Aspen FACE experiment, for 5 example, one could not establish an exposure-response relationship by matching 6 12 yearly exposures and 12 yearly tree sizes, while disregarding age as if size did not also 7 depend on it. This is the basis of the 2007 study of Aspen FACE data by Percy et al. 8 (2007), which compares the size of trees of various ages as if they were all the same age, 9 and was therefore not informative.

9.6.3.1 Comparison of NCLAN-Based Prediction and SoyFACE Data

10	For this ISA, EPA conducted a comparison between yield of soybean as predicted by the
11	composite function three-step process (Section <u>9.6.2</u>) using NCLAN data, and
12	observations of yield in SoyFACE. The median composite function for relative yield was
13	derived for the 11 NCLAN soybean Weibull functions for non-droughted studies, and
14	comparisons between the predictions of the median composite and SoyFACE
15	observations were conducted as follows.

- 16 For the years 2007 and 2008, SoyFACE yield data were available for 7 and 6 genotypes, 17 respectively. The EPA used those data to compare the relative change in yield observed 18 in SoyFACE in a given year between ambient O_3 and elevated O_3 , versus the relative 19 change in yield predicted by the NCLAN-based median composite function between 20 those same two values of O_3 exposure. The two parameter median composite function for 21 relative yield of soybean based on NCLAN data was used to predict yield response at the 22 two values of exposure observed in SoyFACE in each year, and the change between yield 23 under ambient and elevated was compared to the change observed in SoyFACE for the 24 relevant year (Table 9-12). This approach results in a direct comparison of predicted 25 versus observed change in yield. Because the value of relative response between any two 26 values of O_3 exposure is independent of the intercept α , this comparison does not require 27 prediction of the absolute values of the responses.
- 28 Since comparisons of absolute values might be of interest, the predictive functions were 29 also scaled to the observed data: SoyFACE data were used to compute an intercept α 30 while the shape and scale parameters (β and η) were held at their value in the NCLAN 31 predictive model. This method gives a comparison of prediction and observation that 32 takes all the observed information into account to provide the best possible estimate of 33 the intercept, and thus the best possible scaling (

1	Table 9-13 and Figure 9-18). For the comparison of NCLAN and SoyFACE, this
2	validation was possible for 2007 and 2008, where data for 7 and 6 soybean genotypes,
3	respectively, were available. The median composite function for relative yield was
4	derived for the 11 NCLAN soybean Weibull functions for nondroughted studies, and the
5	values of median yield under ambient exposure at SoyFACE in 2007 and 2008 were used
6	to obtain an estimate of the intercept α for the NCLAN median function in each of the
7	two years. Table 9-12 presents the results of ambient/elevated relative yield comparisons
8	between the NCLAN-derived predictions and SoyFACE observations.
9	Table 9-13 and Figure 9-18 present the results of comparisons between NCLAN-derived
10	predictions and SoyFACE observations of yield, with the predictive function scaled to
11	provide absolute yield values.

Table 9-12Comparison between change in yield observed in the SoyFACE
experiment between elevated and ambient ozone, and change
predicted at the same values of ozone by the median composite
function for NCLAN.

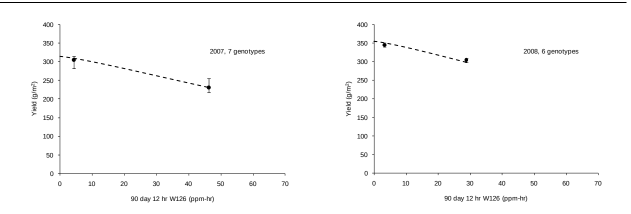
Year	90-day 12-h W126 (ppm-h) observed at SoyFACE			
	Ambient	Elevated	Predicted by NCLAN ^a	Observed at SoyFACE
2007	4.39	46.23	75	76
2008	3.23	28.79	85	88

^aTwo-parameter relative yield model.

Table 9-13Comparison between yield observed in the SoyFACE experiment
and yield predicted at the same values of ozone by the median
composite function for NCLAN.

Year	90-day 12-h W126 (ppm-h) observed at SoyFACE		Yield predicted by NCLAN ^a (g/m ²)		Yield observed at SoyFACE (g/m ²)	
	Ambient	Elevated	Ambient	Elevated	Ambient	Elevated
2007	4.39	46.23	309.2	230.6	305.2	230.6
2008	3.23	28.79	350.3	298.2	344.8	304.4

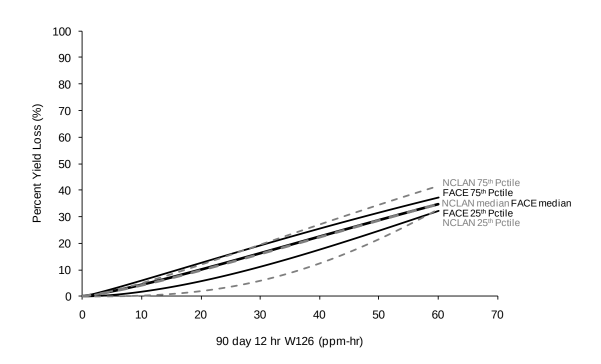
^aThree-parameter absolute yield model with intercept scaled to SoyFACE data.



Note: Black dots are the median of 7 or 6 soybean genotypes in SoyFACE (2007, 2008); bars are Inter-Quartile Range for genotypes; dashed line is median composite model for 11 studies in NCLAN. Source of data: Betzelberger et al. (2010); Morgan et al. (2006); Lee and Hogsett (1996).

Figure 9-18 Comparison of yield observed in SoyFACE experiment in a given year with yield predicted by the median composite function based on NCLAN.

1	Finally, a composite function for the 25th, 50th, and 75th percentiles was developed from
2	SoyFACE annual yield data, and compared to the NCLAN-based function. The process
3	described in Section $9.6.2$ was applied to SoyFACE data for individual genotypes,
4	aggregated over the years during which each was grown; one genotype from 2003 to
5	2007, and six genotypes in 2007 and 2008. First, the three parameter Weibull model
6	described in Section 9.6.2 was estimated using nonlinear regression on exposure-yield
7	data for each genotype separately, over the years for which data were available, totaling
8	seven curves. The 25th, 50th, and 75th percentiles of the predicted values for the two
9	parameter relative yield curves were then identified at every integer of W126 between 0
10	and 60, and a two-parameter Weibull model estimated by regression for the three
11	quartiles. The comparison between these composite functions for the quartiles of relative
12	yield loss in SoyFACE and the corresponding composite functions for NCLAN is
13	presented in Figure 9-19.



Source of data: Betzelberger et al. (2010); Morgan et al. (2006); Lee and Hogsett (1996).

Figure 9-19 Comparison of composite functions for the quartiles of 7 curves for 7 genotypes of soybean grown in the SoyFACE experiment, and for the quartiles of 11 curves for 5 genotypes of soybean grown in the NCLAN project.

As seen in

•	
2	Table 9-13 and <u>Table 9-14</u> , and in <u>Table 9-18</u> , the agreement between predictions based
3	on NCLAN data and SoyFACE observations was notably close in single-year
4	comparisons. Together with the very high agreement between median composite models
5	for NCLAN and SoyFACE (Figure 9-19), it provides very strong mutual confirmation of
6	those two projects' results with respect to the response of yield of soybeans to O ₃
7	exposure. It is readily apparent from these results that the methodology described in
8	Section <u>9.6.2</u> for obtaining predictions of yield or yield loss from NCLAN data is
9	strongly validated by SoyFACE results. As described in Section 9.2, the exposure
10	technologies used in the two projects were in sharp contrast, specifically with respect to
11	the balance each achieved between control of potential interacting factors or confounders,
12	and fidelity to natural conditions. The comparisons that EPA conducted therefore
13	demonstrate that the methodology used in developing the composite functions is resistant
14	to the influence of nuisance variables and that predictions are reliable. They may also

suggest that the aspects in which the two exposure technologies differ have less influence
 on exposure-response than initially supposed. These results are also in agreement with
 comparative studies reviewed in Section <u>9.2.6</u>.

9.6.3.2 Comparison of NHEERL/WED-Based Prediction of Tree Biomass Response and Aspen FACE Data

EPA also conducted two comparisons between prediction of above-ground biomass loss based on NHEERL/WED results and observations from Aspen FACE. The median composite function was developed from NHEERL/WED data for 11 studies that used wild-type seedlings of aspen as well as four clonally propagated genotypes. All plants were grown in OTCs for one growing season before being destructively harvested. Aspen FACE data were from clonally propagated trees of five genotypes grown from 1998 to 2003, with above-ground biomass calculated using allometric equations derived from data for trees harvested destructively in 2000 and 2002 (King et al., 2005).

- 12 The two parameter median composite function for relative biomass was used to predict 13 biomass response under the observed elevated exposure, relative to its value under 14 observed ambient exposure, for each separate year of Aspen FACE. EPA first compared 15 Aspen FACE observations of the change in biomass between ambient and elevated 16 exposure with the corresponding prediction at the same values of exposure. Comparisons 17 between observed and predicted absolute biomass values were then conducted for each 18 year by scaling the predictive function to yearly Aspen FACE data as described for 19 soybean data in Section 9.6.3.1. In all cases, yearly 90 day 12-hour W126 values for 20 Aspen FACE were computed as the cumulative average from the year of planting up to 21 the year of interest. A comparison of composite functions between NHEERL/WED and 22 Aspen FACE, similar to the one performed for NCLAN and SoyFACE, was not possible: 23 as discussed in the introduction to Section 9.6, the pairing of 12 exposure values from 24 separate years and 12 values of biomass cannot be the basis for a model of exposure-25 response, because the trees continued growing for the six-year period of exposure. 26 Because the same trees were used for the entire duration, and continued to grow, data 27 could not be aggregated over years. Table 9-14 presents the results of ambient/elevated 28 relative biomass comparisons between the NHEERL/WED-derived predictions and 29 Aspen FACE observations.
- Table 9-15 and Figure 9-20 present the results of the comparison between
 NHEERL/WED-derived predictions and Aspen FACE observations for absolute biomass,
 using Aspen FACE data to scale the NHEERL/WED-derived composite function.

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Table 9-14Comparison between change in above-ground biomass elevated
and ambient ozone in Aspen FACE experiment in 6 year, and
change predicted at the same values of ozone by the median
composite function for NHEERL/WED.

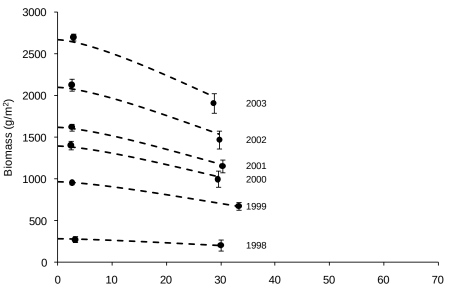
Ambient	Elevated	Predicted by	Observed at Asnen FACE
		NHEERL/WED ^a	Observed at Aspen FACE
3.19	30.08	74	75
2.61	33.85	70	70
2.43	30.16	74	71
2.55	31.00	73	71
2.51	30.27	74	69
2.86	29.12	75	71
	2.61 2.43 2.55 2.51	2.61 33.85 2.43 30.16 2.55 31.00 2.51 30.27	2.61 33.85 70 2.43 30.16 74 2.55 31.00 73 2.51 30.27 74

^aTwo-parameter relative biomass model

Table 9-15Comparison between above-ground biomass observed in AspenFACE experiment in 6 year and biomass predicted by the median
composite function based on NHEERL/WED.

Year	90 day 12-h W126 (ppm-h) Cumulative Average observed at Aspen FACE		Biomass Predicted by NHEERL/WED ^a (g/m ²)		Biomass Observed at Aspen FACE (g/m ²)	
	Ambient	Elevated	Ambient	Elevated	Ambient	Elevated
1998	3.19	30.08	276.0	203.2	274.7	204.9
1999	2.61	33.85	958.7	668.3	955.3	673.3
2000	2.43	30.16	1382.4	1022.8	1400.3	998.6
2001	2.55	31.00	1607.0	1173.7	1620.7	1154.9
2002	2.51	30.27	2079.0	1532.1	2125.9	1468.4
2003	2.86	29.12	2640.1	1981.2	2695.2	1907.8

^aThree-parameter absolute biomass model with intercept scaled to Aspen FACE data.



90 day 12 hr W126 (yearly cumulative average, ppm-hr)

Note: Black dots are aspen biomass/m² for 3 FACE rings filled with an assemblage of 5 clonal genotypes of aspen at Aspen FACE; bars are SE for 3 rings; dashed line is median composite model for 4 clonal genotypes and wild-type seedlings in 11 NHEERL/WED 1-year OTC studies.

Source of data: King et al. (2005); Lee and Hogsett (1996).

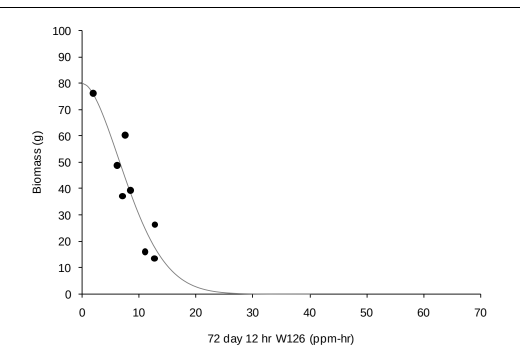
Figure 9-20 Comparison between above-ground biomass observed in Aspen FACE experiment in 6 year and biomass predicted by the median composite function based on NHEERL/WED.

1	As in the comparisons between NCLAN and SoyFACE, the agreement between
2	predictions based on NHEERL/WED data and Aspen FACE observations was very close.
3	The results of the two projects strongly reinforce each other with respect to the response
4	of aspen biomass to O_3 exposure. The methodology used for obtaining the median
5	composite function is shown to be capable of deriving a predictive model despite
6	potential confounders, and despite the added measurement error that is expected from
7	calculating biomass using allometric equations. In addition, the function based on
8	one year of growth was shown to be applicable to subsequent years.
9	The results of experiments that used different exposure methodologies, different
10	genotypes, locations, and durations converged to the same values of response to O_3
11	exposure for each of two very dissimilar plant species, and predictions based on the
12	earlier experiments were validated by the data from current ones. However, in these
13	comparisons, the process used in establishing predictive functions involved aggregating
14	data over variables such as time, locations, and genotypes, and the use of a robust statistic

1	(quartiles) for that aggregation. The validating data, from SoyFACE and Aspen FACE,
2	were in turn aggregated over the same variables. The accuracy of predictions is not
3	expected to be conserved for individual values of those variables over which aggregation
4	occurred. For example, the predicted values for soybean, based on data for five
5	genotypes, are not expected to be valid for each genotype separately. As shown in the
6	validation, however, aggregation that occurred over different values of the same variable
7	did not affect accuracy: composite functions based on one set of genotypes were
8	predictive for another set, as long as medians were used for both sets. A study of
9	cottonwood (Populus deltoides) conducted using a naturally occurring gradient of O3
10	exposure (Gregg et al., 2006, 2003) may provide an illustration of the response of an
11	individual species whose response is far from the median response for an aggregation of
12	species.

9.6.3.3 Exposure-Response in a Gradient Study

13	Gregg et al. (2003) grew saplings of one clonally propagated genotype of cottonwood
14	(<i>Populus deltoides</i>) in seven locations within New York City and in the surrounding
15	region between July and September in 1992, 1993 and 1994, and harvested them 72 days
16	after planting. Owing to regional gradients of atmospheric O_3 concentration, the
17	experiment yielded eight levels of exposure (Figure 9-21), and the authors were able to
18	rule out environmental variables other than O_3 to account for the large differences in
19	biomass observed after one season of growth. The deficit in growth increased
20	substantially faster with increasing O_3 exposure than has been observed in aspen, another
21	species of the same genus (<i>Populus tremuloides</i> , Section <u>9.6.3.2</u>). Using a three
22	parameter Weibull model (Figure 9-21), the biomass of cottonwood at a W126 exposure
23	of 15 ppm-h, relative to biomass at 5 ppm-h, is estimated to be 0.18 (18% of growth at
24	5 ppm-h). The relative biomass of trembling aspen within the same 5-15 ppm-h range of
25	exposure is estimated to be 0.92, using the median composite model for aspen whose
26	very close agreement with Aspen FACE data was shown in Section <u>9.6.3.2</u> . Using a
27	median composite function for all deciduous trees in the NHEERL/WED project (6
28	species in 21 studies) also gives predictions that are very distant from the cottonwood
29	response observed in this experiment. For all deciduous tree species in NHEERL/WED,
30	biomass at a W126 exposure of 15 ppm-h, relative to biomass at 5 ppm-h, was estimated
31	to be 0.87.



Note: Line represents the three-parameter Weibull model. Source: Modified with permission of Nature Publishing Group (<u>Gregg et al., 2003</u>).

Figure 9-21 Above-ground biomass for one genotype of cottonwood grown in seven locations for one season in 3 years.

1	As shown in Section $9.6.2$, the median models available for trembling aspen and soybean
2	have verifiable predictive ability for those particular species. This suggests that the
3	corresponding NCLAN- and NHEERL/WED-based models for multiple crop and tree
4	species can provide reliable estimates of losses for similar assortments of species.
5	However, their predictive ability would likely be poor for individual species not tested.
6	The cottonwood data of Gregg et al. (2003) show an extremely severe response to O_3 .
7	They are consistent with the expectation that among species and genotypes, some are
8	likely to be substantially more sensitive than a median measure, such as the estimate
9	produced by NHEERL/WED (Figure 9-16), but the sensitivity of this particular species
10	has not been studied elsewhere.
11	An alternative hypothesis for the difference between the response of cottonwood in this
12	experiment and deciduous tree species in NHEERL/WED, or the difference between the
13	response of cottonwood and aspen in NHEERL/WED and Aspen FACE, could be the
14	presence of confounding factors in the environments where the experiment was
15	conducted. However, variability in temperature, moisture, soil fertility, and atmospheric

1deposition of N were all ruled out by Gregg et al. (2003) as contributing to the observed2response to O3. In addition, this hypothesis would imply that the unrecognized3confounder(s) were either absent from both OTC and FACE studies, or had the same4value in both. This is not impossible, but the hypothesis that cottonwood is very sensitive5to O3 exposure is more parsimonious, and sufficient.

9.6.3.4 Meta-analyses of growth and yield studies

6	Since the 2006 O ₃ AQCD, five studies have used meta-analytic methods to integrate
7	results from experimental studies of crops or tree species relevant to the U.S. It is
8	possible to obtain exposure-response data for growth and yield from those meta-analyses,
9	but because all of them provided summary measurements of O3 exposure as hourly
10	averages of various lengths of exposures, comparisons with exposure-response results
11	where exposure is expressed as W126 are problematic. Table 9-16 summarizes the
12	characteristics of the five meta-analyses. They all included studies conducted in the U.S.
13	and other locations worldwide, and all of them expressed responses as comparative
14	change between levels of exposure to O ₃ , with carbon filtered air (CF) among those
15	levels. Using hourly average concentration to summarize exposure, CF rarely equates
16	with absence of O ₃ , although it almost always near zero when exposure is summarized as
17	W126, SUM06, or AOT40.

Table 9-16 Meta-analyses of growth or yield studies published since 2005.

Study	Number of articles included	Years of publication surveyed	Crop, species or genera	Response	Number of O ₃ levels	Duration of exposure
<u>Ainsworth</u> (2008)	12	1980-2007	Rice	Yield	2	unreported
<u>Feng et al.</u> (2008b)	53	1980-2007	Wheat	Yield	5	>10 days
<u>Feng and</u> <u>Kobayashi</u> (2009)	All crops together : 81	1980-2007	Potato, barley, wheat, rice, bean, soybean	Yield	3	>10 days
<u>Grantz et al.</u> (2006)	16	1992-2004	34 Herbaceous dicots 21 Herbaceous monocots 5 Tree species	Relative Growth Rate	2	2-24 weeks
<u>Wittig et al.</u> (2009)	All responses:263 Articles that included biomass:unreported	1970-2006	4 Gymnosperm tree genera 11 Angiosperm tree genera	Total biomass	4	>7 days

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1	The only effect of O_3 exposure on yield of rice reported in <u>Ainsworth (2008</u>) was a
2	decrease of 14% with exposure increasing from CF to 62 ppb average concentration.
3	<u>Feng et al. (2008b</u>) were able to separate exposure of wheat into four classes with average
4	concentrations of 42, 69, 97, and 153 ppb, in data where O_3 was the only treatment. Mean
5	responses relative to CF were yield decreases of 17, 25, 49, and 61% respectively. Feng
6	et al. (2008b) observed that wheat yield losses were smaller under conditions of drought,
7	and that Spring wheat and Winter wheat appeared similarly affected. However, mean
8	exposure in studies of Winter wheat was substantially higher than in studies of Spring
9	wheat (86 versus 64 ppb), which suggests that the yield of Spring wheat was in fact more
10	severely affected, since yield was approximately the same, even though Spring wheat was
11	exposed to lower concentrations. Exposures of the six crops considered in Feng and
12	Kobayashi (2009) were classified into two ranges, each compared to CF air. In the lower
13	range of exposure (41-49 ppb), potato studies had the highest average exposure (45 ppb)
14	and wheat and rice the lowest (41 ppb). In the higher range (51-75 ppb), wheat studies
15	had the highest average exposure (65 ppb), and potato, barley and rice the lowest
16	(63 ppb). In other words, across the studies included, all crops were exposed to very
17	similar levels of O ₃ . At approximately 42 ppb, the yield of potato, barley, wheat, rice,
18	bean, and soybean declined by 5.3, 8.9, 9.7, 17.5, 19, and 7.7% respectively, relative to
19	CF air. At approximately 64 ppb O ₃ , declines were 11.9, 12.5, 21.1, 37.5, 41.4, and
20	21.6%. Grantz et al. (2006) reported Relative Growth Rate (RGR) rather than growth,
21	and did not report O_3 exposure values in a way that would allow calculation of mean
22	exposure for each of the three categories of plants for which RGR changes are reported.
23	All studies used only two levels of exposure, with CF air as the lower one, and most used
24	elevated exposure in the range of 40 to 70 ppb. Decline in RGR was 8.2% for the 34
25	herbaceous dicots, 4.5% for the 21 herbaceous monocots, and 17.9% for the 5 tree
26	species. Finally, Wittig et al. (2009) divided the studies analyzed into three classes of
27	comparisons: CF versus ambient, CF versus elevated, and ambient versus elevated, but
28	reported comparisons between three average levels of exposure besides CF: 40 ppb,
29	64 ppb, and 97 ppb. Corresponding decreases in total biomass relative to CF were 7, 17,
30	and 17%.
31	These meta-analyses provide very strong confirmation of EPA's conclusions from
32	previous O ₃ AQCDs: compared to lower levels of ambient O ₃ , current levels in many
33	locations are having a substantial detrimental effect on the growth and yield of a wide
34	variety of crops and natural vegetation. They also confirm strongly that decreases in
35	growth and yield continue at exposure levels higher than current ambient levels.
36	However, direct comparisons with the predictions of exposure-response models that use
37	concentration-weighted cumulative metrics are difficult.

9.6.3.5 Additional exposure-response data

1The studies summarized in Table 9-17 and Table 9-18 contain growth or yield exposure-2response data at too few levels of exposure for exposure-response models, and/or used3metrics other than W126. These tables update Tables AX9-16 through AX9-19 of the42006 O3 AQCD.

9.6.4 Summary

5 None of the information on effects of O_3 on vegetation published since the 2006 O_3 6 AQCD has modified the assessment of quantitative exposure-response relationships that 7 was presented in that document. This assessment updates the 2006 exposure-response 8 models by computing them using the W126 metric, cumulated over 90 days. Almost all 9 of the experimental research on the effects of O₃ on growth or yield of plants published 10 since 2006 used only two levels of exposure. In addition, hourly O_3 concentration data 11 that would allow calculations of exposure using the W126 metric are generally 12 unavailable. However, two long-term experiments, one with a crop species (soybean), 13 one with a tree species (aspen), have produced data that can be used to validate the 14 exposure-response models presented in the 2006 O3 AQCD, and methodology used to 15 derive them.

16 Quantitative characterization of exposure-response in the 2006 O_3 AQCD was based on 17 experimental data generated for that purpose by the National Crop Loss Assessment 18 Network (NCLAN) and EPA National Health and Environmental Effects Research 19 Laboratory, Western Ecology Division (NHEERL-WED) projects, using OTCs to expose 20 crops and trees seedling to O_3 . In recent years, yield and growth results for two of the 21 species that had provided extensive exposure-response information in those projects have 22 become available from studies that used FACE technology, which is intended to provide 23 conditions much closer to natural environments (Pregitzer et al., 2008; Morgan et al., 24 2006; Morgan et al., 2004; Dickson et al., 2000). The robust methods that were used 25 previously with exposure measured as SUM06 were applied to the NCLAN and 26 NHEERL-WED data with exposure measured as W126, in order to derive single-species 27 median models for soybean and aspen from studies involving different genotypes, years, 28 and locations. The resulting models were used to predict the change in yield of soybean 29 and biomass of aspen between the two levels of exposure reported in recent FACE 30 experiments. Results from these new experiments were exceptionally close to predictions 31 from the models. The accuracy of model predictions for two widely different plant 32 species provides support for the validity of the corresponding multiple-species models for 33 crops and trees in the NCLAN and NHEERL-WED projects. However, variability among

1	species in those projects indicates that the range of sensitivity is likely quite wide. This
2	was confirmed by a recent experiment with cottonwood in a naturally occurring gradient
3	of exposure (Gregg et al., 2006), which established the occurrence of species with
4	responses substantially more severe under currently existing conditions than are predicted
5	by the median model for multiple species.
6	Results from several meta-analyses have provided approximate values for responses of
7	yield of soybean, wheat, rice and other crops under broad categories of exposure, relative
8	to charcoal-filtered air (Ainsworth, 2008; Feng et al., 2008b; Morgan et al., 2003).
9	Likewise, Feng and Kobayashi (2009) have summarized yield data for six crop species
10	under various broad comparative exposure categories, while Wittig et al. (2009) reviewed
11	263 studies that reported effects on tree biomass. However, these analyses have proved
12	difficult to compare with exposure-response models, especially given that exposure was
13	not expressed in the same W126 metric.

Table 9-17Summary of studies of effects of ozone exposure on growth and
yield of agricultural crops.

		5 1			
Species Facility Location	Exposure Duration	O ₃ Exposure (Additional Treatment)	Response Measured	Percent Change from CF (Percent Change from Ambient)	Reference
Alfalfa (<i>Medicago sativa</i>) OTC; 0.27m ³ pots Federico, Italy	2 yr, 2005, 2006	AOT40: CF 0 ppm-h 13.9 ppm-h (2005), 10.1 ppm-h (2006) (NaCl: 0.29, 0.65, 0.83, 1.06 deciSiemens/meter)	Total shoot yield	n.s. (N/A)	<u>Maggio et al.</u> (2009)
Bean (<i>Phaseolus vulgaris</i> I. cv Borlotto) OTC; ground- planted Curno, Italy	3 months, 2006	Seasonal AOT40: CF (0.5 ppm-h); ambient (4.6 ppm-h) (N/A)	# Seeds per plant; 100-seed weight	-33 (N/A) n.s. (N/A)	<u>Gerosa et al.</u> (<u>2009</u>)
Big Blue Stem (<i>Andropogon</i> <i>gerardii</i>) OTC Alabama	4 months, 2003	12-h avg: CF (14 ppb), Ambient (29 ppb), Elevated (71 ppb) (N/A)	Final harvest biomass; RVF	n.s. (n.s.) -7 (-7)	<u>Lewis et al.</u> (2006)
<i>Brassica napus</i> cv. Westar Growth chambers Finland	17-26 days	8-h avg: CF (0 ppb), 100 ppb (Bt/non-Bt; herbivory)	Shoot biomass	-30.70 (N/A)	<u>Himanen et al.</u> (2009b)
Corn (<i>Zea may</i> s cv. Chambord) OTC France	33 days	AOT40 ppm-h: 1.1; 1.3; 4.9; 7.2; 9.3; 12.8 (N/A)	Total above- ground biomass	N/A (Highest treatment caused - 26% change)	<u>Leitao et al.</u> (2007a)

Species Facility Location	Exposure Duration	O ₃ Exposure (Additional Treatment)	Response Measured	Percent Change from CF (Percent Change	Reference
				from Ambient)	
Cotton cv. Pima OTC; 9-L pots San Joaquin Valley, CA	8 weeks	12-h avg: 12.8 ± 0.6; 79.9 ± 6.3; 122.7 ± 9.7 (N/A)	Above-ground biomass	-76 (n.s.)	<u>Grantz and</u> <u>Shrestha (2006</u>)
Eastern Gamagrass (<i>Tripsacum</i> <i>dactyloides</i>) OTC Alabama	4 months, 2003	12-h avg: CF (14ppb), Ambient (29 ppb), Elevated (71 ppb) (N/A)	Final harvest biomass; RVF	+68 (+42); -17 (-12)	<u>Lewis et al.</u> (2006)
Grapevine (<i>Vitis vinivera</i>) OTC Austria	3 yr, May-Oct	AOT40 ppm-h: CF (0), Ambient (7-20), Elevated. 1 (20-30), Elevated. 2 (38-48)	Total fruit yield/ Sugar yield	-20 to -80 in different yr (-20 to -90 in different yr)	<u>Soja et al.</u> (2004)
Mustard (<i>Brassica</i> <i>campestris</i>) Chambers; 7.5-cm pots	10 days	CF & 67.8 ppb for 7 h (N/A)	Seeds/plant	n.s. (N/A)	<u>Black et al.</u> (2007)
Oilseed Rape (<i>Brassica napus</i>) OTC Yangtze Delta, China	39 days	Daily avg: 100 ppb, one with diurnal variation and one with constant concentration (N/A)	Biomass and pods per plant	Diurnal variability reduced both biomass and pod number more than constant fumigation (N/A)	<u>Wang et al.</u> (2008)
Peanut (<i>Arachis hypogaea</i>) OTC Raleigh, NC	3 yr	12-h avg: CF (22 ppb), Ambient (46 ppb), Elevated (75ppb) (CO ₂ : 375 ppm; 548 ppm; 730 ppm)	Yield (seed weight, g/m)	-33 (-8)	<u>Burkey et al.</u> (2007)
<i>Poa pratensis</i> OTC Braunschweig, Germany	2000-2002: 4-5 weeks in the Spring	8-h avg: CF+25 (21.7), NF+50 (73.1) (Competition)	Total biomass (g DW/pot)	N/A (n.s.)	Bender et al. (2006)
Potato (Solanum tuberosum) OTC; CHIP	1988,1999. Emergence to harvest	AOT40:CF (0); Ambient (0.27-5.19); NF (0.002-2.93)	Tuber yield averaged across 5 field-sites;	N/A (-27% -+27%, most comparisons n.s.) Linear	Vandermeiren et al. (2005)
6 northern European locations		NF+ (3.10-24.78 (N/A)	Tuber starch content regressed against $[O_3]$ report sig. \pm slope with increasing $[O_3]$	regression slope = -0.0098)	
Rice (<i>Oryza sativa</i>) OTC Raleigh, NC	1997-1998, June- September	12-h mean ppb: CF (27.5), Elevated (74.8) (CO ₂₎	Total biomass; Seed yield	-25(N/A) -13 to 20 (N/A)	Reid and Fiscus (2008)

Species Facility Location	Exposure Duration	O ₃ Exposure (Additional Treatment)	Response Measured	Percent Change from CF (Percent Change from Ambient)	Reference
Rice (<i>Oryza</i> sativa) 20 Asian cultivars OTC Gunma Prefecture, Japan	2008 growing season	Daily avg (ppb): CF (2), 0.8×ambient (23); 1 ×ambient (28); 1.5×ambient (42); 2×ambient (57) (Cultivar comparisons)	Yield	From n.s. to -30 across all cultivars	Sawada and Kohno (2009)
Seminatural grass FACE Le Mouret, Switzerland	5 yr	Seasonal AOT40: Ambient (0.1-7.2 ppm-h); Elevated. (1.8-24.1 ppm-h) (N/A)	Relative annual yield	N/A (2×faster decrease in yield/yr)	<u>Volk et al.</u> (2006)
Soybean OTC; CRA Bari, Italy	2003-2005 growing seasons	Seasonal AOT40 ppm-h: CF (0), Ambient (3.4), High (9.0) (Drought)	Yield	-46 (-9)	Bou Jaoudé et al. (2008b)
Soybean (<i>Glycine max</i> cv. 93B15) SoyFACE Urbana, IL	2002, 2003 growing seasons	8-h avg: Ambient (62 & 50 ppb), Elevated (75 & 63 ppb) (N/A)	Yield	N/A (-15 in 2002; -25 in 2003)	<u>Morgan et al.</u> (2006)
Soybean (<i>Glycine max</i> cv. Essex) Chambers; 21 L Raleigh, NC	2x3 months	12-h avg: CF (28), Elevated (79), Elevated flux (112) (CO ₂ : 365 & 700)	Seed mass per plant	-30 (N/A)	<u>Booker and</u> Fiscus (2005)
Soybean (<i>Glycine max</i> cv. Essex) OTCs; 21-L pots Raleigh, NC	2×3 months	12-h avg: CF (18); Elevated (72) (CO ₂ : 367 & 718)	Seed mass per plant	-34 (N/A)	<u>Booker et al.</u> (2004b)
Soybean (<i>Glycine max</i> cv. Tracaja) Chambers; pots Brazil	20 days	12-h avg: CF & 30 ppb (N/A)	Biomass	-18 (N/A)	Bulbovas et al. (2007)
Soybean (<i>Glycine max</i>) 10 cultivars SoyFACE Urbana, IL	2007 & 2008	8-h avg: Ambient (46.3 & 37.9), Elevated (82.5 & 61.3) (Cultivar comparisons)	Yield	N/A (-17.20)	<u>Betzelberger et</u> <u>al. (2010</u>)
Spring Wheat (<i>Triticum aestivum</i> cv. Minaret; Satu; Drabant; Dragon) OTCs Belgium, Finland, & Sweden	1990-2006	Seasonal AOT40s ranged from 0 to16 ppm-h (N/A)	Seed protein content; 1,000-seed weight regressed across all experiments	N/A (significant negative correlation) N/A (sig negative correlation)	<u>Piikki et al.</u> (2008b)
Strawberry (<i>Fragaria x</i> <i>ananassa</i> Duch. Cv Korona & Elsanta) Growth chambers Bonn, Germany	2 months	8-h avg: CF (0 ppb) & Elevated (78 ppb) (N/A)	Fruit yield (weight/plant)	-16 (N/A)	Keutgen et al. (2005)

Species Facility Location	Exposure Duration	O ₃ Exposure (Additional Treatment)	Response Measured	Percent Change from CF (Percent Change from Ambient)	Reference
Sugarbeet (<i>Beta vulgaris</i> cv. Patriot) OTC Belgium	2003, 2004; 5 months	8-h avg: Ambient (36 ppb); Elevated (62 ppb) (N/A)	Sugar yield	N/A (-9)	<u>De Temmerman</u> <u>et al. (2007</u>)
Sugarcane (<i>Saccharum spp</i>) CSTR San Joaquin Valley, CA	2007; 11-13 weeks.	12-h avg: CF (4 ppb); Ambient (58); Elevated (147) (N/A)	Total biomass (g/plant)	-40 (-30)	Grantz and Vu (2009)
Sweet Potato Growth chambers Bonn, Germany	4 weeks	8-h avg: CF (0 ppb), Ambient (<40 ppb) Elevated (255 ppb) (N/A)	Tuber weight	-14 (-11.5)	<u>Keutgen et al.</u> (2008)
Tomato (<i>Lycopersicon</i> <i>esculentum</i>) OTC Valencia, Spain	133 days in 1998	8-h mean ppb: CF 16.3, NF 30.1, NF+ 83.2 (Various cultivars; early & late harvest)	Yield	n.s (n.s.)	Dalstein and Vas (2005)
<i>Trifolium</i> <i>Subterraneum</i> OTC; 2.5-L pots Madrid, Spain	29 days	12-h avg: CF (<7.9 ± 6.3); Ambient (34.4 ± 10.8); Elevated (56.4 ± 22.3) (N: 5, 15 & 30 kg/ha)	Above-ground biomass	-45 (-35)	<u>Sanz et al.</u> (2005)
Watermelon (<i>Citrullus lanatus</i>) OTC Valencia, Spain	2000, 2001. 90 days	AOT40: CF (0 ppm-h) Ambient (5.7 ppm-h), Elevated (34.1 ppm-h) (N:0, 14.0 & 29.6 g/pot)	total fruit yield (kg)	n.s. (54)	<u>Calatayud et al.</u> (2006)
Yellow Nutsedge OTC; 9-L pots San Joaquin Valley, CA	8 weeks	12-h avg: 12.8 ± 0.6; 79.9 ± 6.3; 122.7 ± 9.7 (N/A)	above-ground biomass	n.s. (n.s.)	<u>Grantz and</u> <u>Shrestha (2006</u>)

In studies where variables other than O_3 were included in the experimental design, response to O_3 is only provided for the control level of those variables.

Species	Exposure	O ₃ Exposure	Response	Response	Reference
Facility Location	Duration	(Additional Treatment)	Measured	·	
Yellow nutsedge (<i>Cyperus</i> <i>esculentus</i>) CSTR San Joaquin Valley, CA	53 days in 2008	12-h mean ppb: CF (4); CF+ (60); CF2+ (115)	Above-ground biomass; tubers (g/plant)	ns; CF(4.1) CF+(3.9) CF2+(2.7)	<u>Grantz et al. (2010a)</u>
35 herbaceous species OTC Corvallis, OR	1999-2002, May-August	4-yr avg; yearly W126 ppm-h: CF (0), CF+ (21), CF 2+ (49.5)	Total community above-ground biomass (35 species) after 4 years	CF (459 g/m ²), CF+ (457 g/m ²), CF2+ (398 g/m ²)	<u>Pfleeger et al. (2010</u>)
Highbush blackberry (<i>Rubus argutus</i>) OTC Auburn, AL	2004, May-August	12-h mean ppb: CF (21.7), Ambient (32.3), Elevated (73.3)	Vegetative regrowth after pruning	CF (75.1 g/plant), Ambient (76.4 g/plant), Elevated (73.1 g/plant)	Ditchkoff et al. (2009)
Horseweed (<i>Conyza canadensis</i>) CSTR San Joaquin Valley, CA	2005, 2 runs, 28 days each (July-Aug, Sept)	W126 ppm-h: CF(0), CF+ (11), CF 2+ (30) (Glyphosate resistance)	Total biomass (g/plant)	Glyphosate sensitive: CF (0.354) CF+ (0.197) CF2+ (0.106) Glyphosate resistant: CF(0.510) CF+ (0.313) CF2+ (0.143)	<u>Grantz et al. (2008</u>)
Red Oak (<i>Quercus</i> <i>rubrum</i>) Forest sites Look Rock & Twin Creeks Forests, TN	2001-2003, April-October	AOT60: 2001 (11.5), 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2001 in year 2002;2003)	-42.8%; +1%	<u>McLaughlin et al.</u> (2007a)
Pine species Forest sites Look Rock Forest, TN	2001-2003, April-October	AOT60: 2001 (11.5), 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2001 in year 2002;2003)	-62.5%; -2.9%	<u>McLaughlin et al.</u> (2007a)

Table 9-18Summary of studies of effects of ozone exposure on growth of
natural vegetation.

Species Facility Location	Exposure Duration	O₃ Exposure (Additional Treatment)	Response Measured	Response	Reference
Hickory species Forest sites Look Rock Forest, TN	2001-2003, April-October	AOT60: 2001 (11.5), 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2001 in year 2002;2003)	-14%; +30%	<u>McLaughlin et al.</u> (2007a)
Chestnut Oak (<i>Quercus prinus</i>) Forest sites Look Rock Forest, TN	2001-2003, April-October	AOT60: 2001 (11.5), 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2001 in year 2002;2003)	+44%; +55%	<u>McLaughlin et al.</u> (2007a)
Black Cherry (<i>Prunus rigida</i>) Forest sites Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2003 in year 2002)	-75%	<u>McLaughlin et al.</u> (2007a)
Shortleaf pine (<i>Pinus echinata</i>) Forest sites Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2003 in year 2002)	-16.8%	<u>McLaughlin et al.</u> (2007a)
Hemlock (<i>Tsuga canadensis</i>) Forest sites Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2003 in year 2002)	-21.9%	<u>McLaughlin et al.</u> (2007a)
Red Maple (<i>Acer rubrum</i>) Forest sites Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2003 in year 2002)	-59.6%	<u>McLaughlin et al.</u> (2007a)

Species Facility Location	Exposure Duration	O₃ Exposure (Additional Treatment)	Response Measured	Response	Reference
Yellow Poplar (<i>Liriodendron tulipifera</i>) Forest sites Look Rock, Oak Ridge, & Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2001 in years 2002; 2003)	-45.9%; -15.25%	<u>McLaughlin et al.</u> (2007a)
Sugar Maple (<i>Acer</i> <i>saccharum</i>) Forest sites Twin Creeks Forest, TN	2002-2003, April-October	AOT60: 2002 (24.0), 2003 (11.7) (Observational study with multiple environmental variables)	Annual circumference increment (change relative to 2003 in year 2002)	-63.8%	<u>McLaughlin et al.</u> (2007a)
Trembling aspen (<i>Populus tremuloides</i>), 5 genotypes Aspen FACE Rhinelander, WI	1998-2004, May-October	Cumulative avg 90-day 12-h W126. Ambient 3.1 ppm-h Elevated: 27.2 ppm-h (Competition with birch, maple)	main stem volume after 7 years	Ambient: 6.22 dm ³ ; Elevated: 4.73 dm ³	<u>Kubiske et al. (2006</u>)
Hybrid Poplar (<i>Populus trichocarpa x Populus deltoides</i>) OTC Seattle, WA	2003, 3 months	Daily mean (µg/g): CF(<9), Elevated (85- 128)	Total biomass	CF to elevated: -12.9%	<u>Woo and Hinckley</u> (2005)

In studies where variables other than O_3 were included in the experimental design, response to O_3 is only provided for the control level of those variables.

9.7 Summary and Conclusions

1	Based on the evidence presented in Chapter $\underline{9}$ and summarized here, O_3 is causally related
2	or likely to be causally related to effects observed on vegetation and ecosystems. The
3	evidence for these effects spans the entire continuum of biological organization, from the
4	cellular and subcellular level to the whole plant, and up to ecosystem-level processes, and
5	includes evidence for effects at lower levels of organization, leading to effects at higher
6	levels. Given the current state of knowledge, exposure indices that cumulate and
7	differentially weight the higher hourly average concentrations and also include the mid-
8	level values are the most appropriate for use in developing response functions and
9	comparing studies. The framework for causal determinations (see Preamble) has been
10	applied to the body of scientific evidence to examine effects attributed to O3 exposure
11	collectively and the determinations are presented in <u>Table 9-19</u> .

Vegetation and Ecosystem Effects	Conclusions from 2006 O ₃ AQCD	Conclusions from 2011 3rd Draft ISA
Visible Foliar Injury Effects on Vegetation	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause impaired aesthetic quality of many native plants and trees by increasing foliar injury.	Causal Relationship
Reduced Vegetation Growth	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased growth and biomass accumulation in annual, perennial and woody plants, including agronomic crops, annuals, shrubs, grasses, and trees.	Causal Relationship
Reduced Productivity in Terrestrial Ecosystems	There is evidence that O_3 is an important stressor of ecosystems and that the effects of O_3 on individual plants and processes are scaled up through the ecosystem, affecting net primary productivity.	Causal Relationship
Reduced Carbon (C) Sequestration in Terrestrial Ecosystems	Limited studies from previous review	Likely to be a Causal Relationship
Reduced Yield and Quality of Agricultural Crops	Data published since the 1996 O_3 AQCD strengthen previous conclusions that there is strong evidence that current ambient O_3 concentrations cause decreased yield and/or nutritive quality in a large number of agronomic and forage crops.	Causal Relationship
Alteration of Terrestrial Ecosystem Water Cycling	Ecosystem water quantity may be affected by O_3 exposure at the landscape level.	Likely to be a Causal Relationship
Alteration of Below-ground Biogeochemical Cycles	Ozone-sensitive species have well known responses to O_3 exposure, including altered C allocation to below-ground tissues, and altered rates of leaf and root production, turnover, and decomposition. These shifts can affect overall C and N loss from the ecosystem in terms of respired C, and leached aqueous dissolved organic and inorganic C and N.	Causal Relationship
Alteration of Terrestrial Community Composition	Ozone may be affecting above- and below -ground community composition through impacts on both growth and reproduction. Significant changes in plant community composition resulting directly from O_3 exposure have been demonstrated.	Likely to be a Causal Relationship

Table 9-19Summary of ozone causal determinations for vegetation and
ecosystem effects

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10 THE ROLE OF TROPOSPHERIC OZONE IN CLIMATE CHANGE AND UV-B EFFECTS

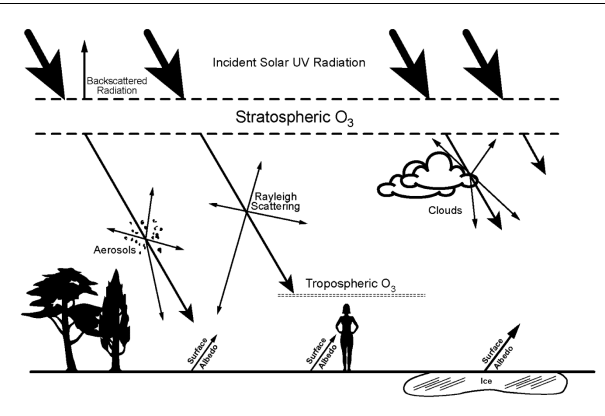
10.1 Introduction

1	Atmospheric O ₃ plays an important role in the Earth's energy budget by interacting with
2	incoming solar radiation and outgoing infrared radiation. Over mid-latitudes,
3	approximately 90% of the total atmospheric O_3 column is located in the stratosphere (Kar
4	et al., 2010; Crist et al., 1994). Therefore, tropospheric O ₃ makes up a relatively small
5	portion (~10%) of the total column of O ₃ over mid-latitudes, but it does play an important
6	role in the overall radiation budget. The next section (Section 10.2) briefly describes the
7	physics of the earth's radiation budget, providing background material for the subsequent
8	two sections assessing how perturbations in tropospheric O ₃ might affect (1) climate
9	through its role as a greenhouse gas (Section 10.3), and (2) health, ecology and welfare
10	through its role in shielding the earth's surface from solar ultraviolet radiation
11	(Section 10.4). The concluding section in this chapter (Section 10.5) includes a summary
12	of effects assessed in this chapter along with their associated causal determinations.

10.2 Physics of the Earth's Radiation Budget

13	Radiant energy from the sun enters the atmosphere in a range of wavelengths, but peaks
14	strongly in the visible (400-750 nm) part of the spectrum. Longer wavelength infrared
15	(750 nm-1 mm) and shorter wavelength ultraviolet (100-400 nm) radiation are also
16	present in the solar electromagnetic spectrum. Since the energy possessed by a photon is
17	inversely proportional to its wavelength, infrared (IR) radiation carries the least energy
18	per photon, and ultraviolet (UV) radiation carries the most energy per photon. UV
19	radiation is further subdivided into classes based on wavelength: UV-A refers to
20	wavelengths from 400-315 nm; UV-B from 315-280 nm; and UV-C from 280-100 nm.
21	By the same argument above describing the relationship between photon wavelength and
22	energy, UV-A radiation is the least energetic and UV-C is the most energetic band in the
23	UV spectrum.
24	The wavelength of radiation also determines how the photons interact with the complex
25	mixture of gases, clouds, and particles present in the atmosphere (see Figure 10-1). UV-A
26	radiation can be scattered but is not absorbed to any meaningful degree by atmospheric
27	gases including O ₃ . UV-B radiation is absorbed and scattered in part within the
28	atmosphere. UV-C is almost entirely blocked by the Earth's upper atmosphere, where it

1	participates in photoionization and photodissociation processes including absorption by
2	stratospheric O ₃ .
3	Since UV-A radiation is less energetic and does not interact with O_3 in the troposphere or
4	the stratosphere and UV-C radiation is almost entirely blocked by stratospheric O ₃ , UV-B
5	radiation is the most important band to consider in relation to tropospheric O ₃ shielding.
6	Furthermore, tropospheric O ₃ plays a "disproportionate" role in absorbing UV-B
7	radiation compared with stratospheric O ₃ on a molecule per molecule basis (Balis et al.,
8	2002; Zerefos et al., 2002; Crist et al., 1994; Bruhl and Crutzen, 1989). This effect results
9	from the higher atmospheric pressure present in the troposphere, resulting in higher
10	concentrations of gas molecules present that can absorb or scatter radiation. For this
11	reason, the troposphere is referred to as a "multiple scattering" regime for UV absorption,
12	compared to the "single scattering" regime in the stratosphere. Thus, careful
13	quantification of atmospheric absorbers and scatterers, along with a well-resolved
14	description of the physics of these interactions, is necessary for predicting the effects of
15	tropospheric O_3 on UV-B flux at the surface.
16	Solar flux at all wavelengths has a temporal dependence, while radiative scattering and
17	absorption have strong wavelength, path length, and gas/particle concentration
18	dependencies. These combine to create nonlinear effects on UV flux at the Earth's
19	surface. Chapter 10 of the 2006 O ₃ AQCD (U.S. EPA, 2006b) describes in detail several
20	key factors that influence the spatiotemporal distribution of ground-level UV radiation
21	flux, including: (1) long-term solar activity including sunspot cycle; (2) solar rotation; (3)
22	the position of the Earth in its orbit around the sun; (4) atmospheric absorption and
23	scattering of UV radiation by gas molecules and aerosol particles; (5) absorption and
24	scattering by stratospheric and tropospheric clouds; and (6) surface albedo. The
25	efficiencies of absorption and scattering are highly dependent on the concentration of the
26	scattering medium, particle size (for aerosols and clouds), and the altitude at which these
27	processes are occurring. These properties are sensitive to meteorology, which introduces
28	additional elements of spatial and temporal dependency in ground-level UV radiation
29	flux.



Source: 2006 O₃ AQCD (<u>U.S. EPA, 2006b</u>).

Figure 10-1 Diagram of the factors that determine human exposure to ultraviolet radiation.

1	About 30% of incoming solar radiation is directly reflected back to space, mainly by
2	clouds or surfaces with high albedo (reflectivity), such as snow, ice, and desert sand.
3	Radiation that does penetrate to the Earth's surface and is absorbed can be re-emitted in
4	the longwave (infrared) portion of the spectrum; the rest goes into evaporating water or
5	soil moisture or emerges as sensible heat. The troposphere is opaque to the outgoing
6	longwave radiation. Polyatomic gases such as water vapor, CO ₂ , CH ₄ , and O ₃ absorb and
7	re-emit the radiation upwelling from the Earth's surface, reducing the efficiency with
8	which that energy returns to space. In effect, these gases act as a blanket warming the
9	Earth's surface. This phenomenon, known as the "Greenhouse Effect," was first
10	quantified in the 19th century (Arrhenius, 1896), and gives rise to the term "greenhouse
11	gas." The most important greenhouse gas is water vapor.

10.3 Effects of Tropospheric Ozone on Climate

10.3.1 Background

1	As a result of its interaction with incoming solar radiation and outgoing longwave
2	radiation, tropospheric O ₃ plays a major role in determining climate, and increases in its
3	abundance may contribute to climate change (<u>IPCC, 2007c</u>). Models estimate that the
4	global average concentration of O_3 in the troposphere has increased 30-70% since the
5	preindustrial era (Gauss et al., 2006), while observations indicate that in some regions
6	tropospheric O ₃ may have increased by factors as great as 4 or 5 (Marenco et al., 1994;
7	Staehelin et al., 1994). These increases are tied to the rise in emissions of O_3 precursors
8	from human activity, mainly fossil fuel consumption and agricultural processes.
9	The effect on climate of the tropospheric O_3 change since preindustrial times has been
10	estimated to be about 25-40% of the anthropogenic CO ₂ effect and about 75% of the
11	anthropogenic CH ₄ effect (<u>IPCC, 2007c</u>), ranking it third in importance behind these two
12	major greenhouse gases. In the 21st century, as the Earth's population continues to grow
13	and energy technology spreads to developing countries, a further rise in the global
14	concentration of tropospheric O ₃ is likely, with associated consequences for human health
15	and ecosystems relating to climate change.
16	To examine the science of a changing climate and to provide balanced and rigorous
17	information to policy makers, the World Meteorological Organization (WMO) and the
18	United Nations Environment Programme (UNEP) formed the Intergovernmental Panel on
19	Climate Change (IPCC) in 1988. The IPCC supports the work of the Conference of
20	Parties (COP) to the United Nations Framework Convention on Climate Change
21	(UNFCCC). The IPCC periodically brings together climate scientists from member
22	countries of WMO and the United Nations to review knowledge of the physical climate
23	system, past and future climate change, and evidence of human-induced climate change.
24	IPCC climate assessment reports are issued every five to seven years.
25	This section draws in part on the fourth IPCC Assessment Report (AR4) (IPCC, 2007c),
26	as well as other peer-reviewed published research. Section $10.3.2$ reviews evidence of
27	climate change in the recent past and projections of future climate change. It also offers a
28	brief comparison of tropospheric O_3 relative to other greenhouse gases. Section <u>10.3.3</u>
29	describes factors that influence the magnitude of tropospheric O_3 effects on climate.
30	Section $10.3.4$ considers the competing effects of O_3 precursors on climate. Finally,
31	Section <u>10.3.5</u> and Section <u>10.3.6</u> describe the effects of changing tropospheric O_3
32	concentrations on past and future climate. Downstream effects resulting from climate
33	change, such as ecosystem responses, are outside the scope of this assessment, which

focuses rather on the effects of changes in tropospheric O_3 concentrations on radiative forcing and climate.

10.3.2 Climate Change Evidence and the Influence of Tropospheric Ozone

10.3.2.1 Climate Change in the Recent Past

- 3 From the end of the Last Ice Age 12,000 years ago until the mid-1800s, observations 4 from ice cores show that concentrations of the long-lived greenhouse gases CO_2 , CH_4 , 5 and N_2O have been relatively stable. Unlike these greenhouse gases, O_3 is not preserved 6 in ice, and no record of it before the late 1800s exists. Models, however, suggest that it, 7 too, has remained relatively constant during this time period (Thompson et al., 1993; 8 Thompson, 1992). The stable mix of these greenhouse gases in the atmosphere, together 9 with water vapor, has kept the global mean temperature of the Earth close to 15°C. 10 Without the presence of greenhouse gases in the atmosphere, the Earth's global mean 11 temperature would be about 30°C cooler, or -15°C. 12 Since the start of the Industrial Revolution, human activity has led to observable
- 13 increases of greenhouse gases in the atmosphere, mainly through fossil fuel combustion. 14 According to the IPCC AR4 (IPCC, 2007c), there is now "very high confidence" that the 15 net effect of anthropogenic greenhouse gas emissions since 1750 has led to warming, and 16 it is "very likely" that human activity contributed to the 0.76°C rise in global mean 17 temperature observed over the last century. The increase of tropospheric O_3 may have 18 contributed 0.1-0.3°C warming to the global climate during this time period (Hansen et 19 al., 2005; Mickley et al., 2004). Global cooling due to anthropogenic aerosols (IPCC, 20 2007c) has likely masked the full warming effect of the anthropogenic greenhouse gases 21 on a global scale.

10.3.2.2 Projections of Future Climate Change

The IPCC AR4 projects a warming of ~0.2°C per decade for the remainder of the 21st century (IPCC, 2007c). Even at constant concentrations of greenhouse gases in the atmosphere, temperatures are expected to increase by about 0.1°C per decade, due to the slow response of oceans to the warming applied so far. It is likely that the Earth will experience longer and more frequent heat waves in the 21st century, together with more frequent droughts and/or heavy precipitation events in some regions, due to perturbations in the hydrological cycle that result from changing temperatures. Sea levels could

1

2

- increase by 0.3-0.8 meters by 2300 due to thermal expansion of the oceans. The extent of
 Arctic sea ice is expected to decline, and contraction of the Greenland ice sheet could
 further contribute to the sea level rise (<u>IPCC, 2007c</u>).
- 4 Projections of future climate change are all associated with some degree of uncertainty. A 5 major uncertainty involves future trends in the anthropogenic emissions of greenhouse 6 gases or their precursors. For the IPCC AR4 climate projections, a set of distinct 7 "storylines" or emission pathways was developed (IPCC, 2000). Each storyline took into 8 account factors such as population growth, mix of energy technologies, and the sharing of 9 technology between developed and developing nations, and each resulted in a different 10 scenario for anthropogenic emissions. When these trends in emissions are applied to 11 models, these scenarios yield a broad range of possible climate trajectories for the 21st 12 century.
- 13 A second factor bringing large uncertainty to model projections of future climate is the 14 representation of climate and, especially, climate feedbacks. A rise in surface 15 temperatures would perturb a suite of other processes in the earth-atmosphere-ocean 16 system, which may in turn either amplify the temperature increase (positive feedback) or 17 diminish it (negative feedback). One important feedback involves the increase of water 18 vapor content of the atmosphere that would accompany higher temperatures (Bony et al., 19 2006). Water vapor is a potent greenhouse gas; accounting for the water vapor feedback 20 may increase the climate sensitivity to a doubling of CO₂ by nearly a factor of two (Held 21 and Soden, 2000). The ice-albedo feedback is also strongly positive; a decline in snow 22 cover and sea ice extent would diminish the Earth's albedo, allowing more solar energy 23 to be retained at the surface (Holland and Bitz, 2003; Rind et al., 1995). A final example 24 of a climate feedback involves the effects of changing cloud cover in a warming 25 atmosphere. Models disagree on the magnitude and even the sign of this feedback on 26 surface temperatures (Soden and Held, 2006).

10.3.2.3 Metrics of Potential Climate Change

27 Two metrics frequently used to estimate the potential climate effect of some perturbation 28 such as a change in greenhouse gas concentration are: (1) radiative forcing; and (2) global 29 warming potential (GWP). These metrics differ in a fundamental way as described below. 30 Radiative forcing is a change in the radiative balance at a particular level of the 31 atmosphere or at the surface when a perturbation is introduced in the earth-atmosphere-32 ocean system. In the global mean, radiative forcing of greenhouse gases at the tropopause 33 (top of the troposphere) is roughly proportional to the surface temperature response 34 (Hansen et al., 2005; NRC, 2005). It thus provides a useful metric for policymakers for

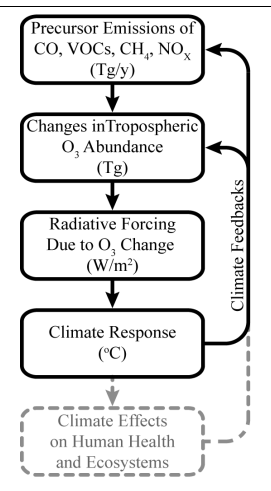
1	assessing the response of the earth's surface temperature to a given change in the
2	concentration of a greenhouse gas. Positive values of radiative forcing indicate warming
3	in a test case relative to the control; negative values indicate cooling. The units of
4	radiative forcing are energy flux per area, or W/m^2 .
5	Radiative forcing requires just a few model years to calculate, and it shows consistency
6	from model to model. However, radiative forcing does not take into account the climate
7	feedbacks that could amplify or dampen the actual surface temperature response,
8	depending on region. Quantifying the change in surface temperature requires a climate
9	simulation in which all important feedbacks are accounted for. As some of these
10	processes are not well understood, the surface temperature response to a given radiative
11	forcing can be highly uncertain and can vary greatly among models and even from region
12	to region within the same model.
13	GWP indicates the integrated radiative forcing over a specified period (usually 100 years)
14	from a unit mass pulse emission of a greenhouse gas or its precursor, and is reported as

121314141616161614from a unit mass pulse emission of a greenhouse gas or its precursor, and is reported as15the magnitude of this radiative forcing relative to that of CO_2 . GWP is most useful for16comparing the potential climate effects of long-lived gases, such as N₂O or CH₄. Since17tropospheric O₃ has a lifetime on the order of weeks to months, GWP is not seen as a18valuable metric for quantifying the importance of O₃ on climate (Forster et al., 2007).19Thus, this assessment focuses on radiative forcing as the metric of climate influence20resulting from changes in tropospheric O₃.

10.3.2.4 Tropospheric Ozone as a Greenhouse Gas

21	Tropospheric O ₃ differs in important ways from other greenhouse gases. It is not emitted
22	directly, but is produced through photochemical oxidation of CO, CH ₄ , and nonmethane
23	volatile organic compounds (VOCs) in the presence of nitrogen oxide radicals
24	$(NO_X = NO + NO_2; \text{ see Chapter } \underline{3}, \text{ Section } \underline{3.2} \text{ for further details on the chemistry of } O_3$
25	formation). It is also supplied by vertical transport from the stratosphere. The lifetime of
26	O_3 in the troposphere is typically a few weeks, resulting in an inhomogeneous
27	distribution that varies seasonally; the distribution of the long-lived greenhouse gases like
28	CO_2 and CH_4 are much more uniform. The longwave radiative forcing by O_3 is mainly
29	due to absorption in the 9.6 μ m window, where absorption by water vapor is weak. It is
30	therefore less sensitive to local humidity than the radiative forcing by CO ₂ or CH ₄ , for
31	which there is much more overlap with the water absorption bands (Lenoble, 1993). And
32	unlike other major greenhouse gases, O_3 absorbs in the shortwave as well as the
33	longwave part of the spectrum.

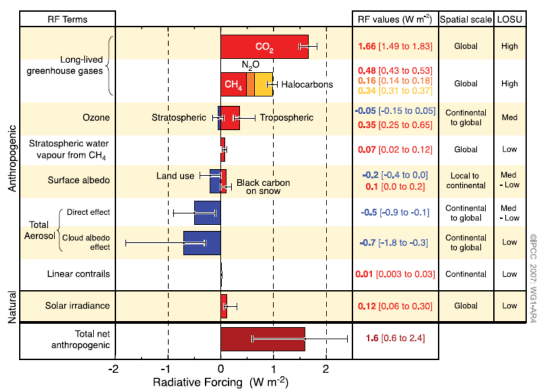
1	<u>Figure 10-2</u> shows the main steps involved in the influence of tropospheric O_3 on climate.
2	Emissions of O ₃ precursors including CO, VOCs, CH ₄ , and NO _X lead to production of
3	tropospheric O_3 . A change in the abundance of tropospheric O_3 perturbs the radiative
4	balance of the atmosphere, an effect quantified by the radiative forcing metric. The earth-
5	atmosphere-ocean system responds to the radiative forcing with a climate response,
6	typically expressed as a change in surface temperature. Finally, the climate response
7	causes downstream climate-related health and ecosystem effects, such as redistribution of
8	diseases or ecosystem characteristics due to temperature changes. Feedbacks from both
9	the climate response and downstream effects can, in turn, affect the abundance of
10	tropospheric O_3 and O_3 precursors through multiple mechanisms. Direct feedbacks are
11	discussed further in Section $10.3.3.4$; the downstream climate effects and their feedbacks
12	are extremely complex and outside the scope of this assessment.



Note: Figure includes the relationship between precursor emissions, changes in tropospheric ozone abundance, radiative forcing, climate response, and climate effects. Units shown are those typical for each quantity illustrated. Feedbacks from both the climate response and climate effects can, in turn, affect the abundance of tropospheric ozone and ozone precursors through multiple feedback mechanisms. Climate effects and their feedbacks are deemphasized in the figure since these downstream effects are extremely complex and outside the scope of this assessment.

Figure 10-2 Schematic illustrating the effects of tropospheric ozone on climate.

1	The <u>IPCC (2007c</u>) reported a radiative forcing of 0.35 W/m^2 for the change in
2	tropospheric O_3 since the preindustrial era, ranking it third in importance after the
3	greenhouse gases CO_2 (1.66 W/m ²) and CH_4 (0.48 W/m ²). Figure 10-3 shows the global
4	average radiative forcing estimates and uncertainty ranges in 2005 for anthropogenic
5	CO ₂ , CH ₄ , O ₃ and other important agents and mechanisms. The error bars encompassing
6	the tropospheric O_3 radiative forcing estimate in the figure range from 0.25 to 0.65 W/m ² ,
7	making it relatively more uncertain than the long-lived greenhouse gases.



RADIATIVE FORCING COMPONENTS

Note: Figure shows the typical geographical extent (spatial scale) of the radiative forcing and the assessed level of scientific understanding (LOSU). The net anthropogenic radiative forcing and its range are also shown. These require summing asymmetric uncertainty estimates from the component terms, and cannot be obtained by simple addition. Additional radiative forcing factors not included here are considered to have a very low LOSU.

Source: Reprinted with permission of Cambridge University Press (IPCC, 2007c).

Figure 10-3 Global average radiative forcing (RF) estimates and uncertainty ranges in 2005 for anthropogenic CO₂, CH₄, ozone and other important agents and mechanisms.

10.3.3 Factors that Influence the Effect of Tropospheric Ozone on Climate

1	This section describes the main factors that influence the magnitude of the climate
2	response to changes in tropospheric O ₃ . They include: (1) trends in the concentration of
3	tropospheric O_3 ; (2) the effect of surface albedo on O_3 radiative forcing; (3) the effect of
4	vertical distribution on O_3 radiative forcing; (4) feedback factors that can alter the climate
5	response to O_3 radiative forcing; and (5) the indirect effects of tropospheric O_3 on the
6	carbon cycle. Trends in stratospheric O ₃ may also affect temperatures at the Earth's
7	surface, but aside from issues relating to stratospheric-tropospheric exchange discussed in
8	Chapter <u>3</u> , Section <u>3.4.1.1</u> , stratospheric O_3 assessment is beyond the scope of this
9	document.

10.3.3.1 Trends in the Concentration of Tropospheric Ozone

1	To first order, the effect of tropospheric O_3 on global surface temperature is proportional
2	to the change in tropospheric O ₃ concentration. The earth's surface temperatures are most
3	sensitive to O_3 perturbations in the mid to upper troposphere. This section therefore
4	focuses mainly on observed O_3 trends in the free troposphere or in regions far from O_3
5	sources, where a change in O_3 concentrations may indicate change throughout the
6	troposphere. Data from ozonesondes, mountaintops, and remote surface sites are
7	discussed, as well as satellite data.

Observed Trends in Ozone since the Preindustrial Era

8 Measurements of O_3 at two European mountain sites dating from the late 1800s to early 9 1900s show values at about 10 ppb, about one-fifth the values observed today at similar 10 sites (Pavelin et al., 1999; Marenco et al., 1994). The accuracy of these early 11 measurements is questionable however, in part because they exhibit O_3 concentrations 12 equivalent to or only a couple of parts per billion greater than those observed at nearby 13 low-altitude sites during the same time period (Mickley et al., 2001; Volz and Kley, 14 <u>1988</u>). A larger vertical gradient in tropospheric O_3 would be expected because of its 15 stratospheric source and its longer lifetime aloft. In another study, Staehelin et al. (1994) 16 revisited observations made in the Swiss mountains during the 1950s and found a 17 doubling in O₃ concentrations from that era to 1989-1991.

- 18 Routine observations of O_3 in the troposphere began in the 1970s with the use of balloon-19 borne ozonesondes, but even this record is sparse. Trends from ozonesondes have been 20 highly variable and dependent on region (Logan et al., 1999). Over most sites in the U.S., 21 ozonesondes reveal little trend. Over Canada, observations show a decline in O_3 between 22 1980 and 1990, then a rebound in the following decade (Tarasick et al., 2005). 23 Ozonesondes over Europe give a mixed picture. European ozonesondes showed increases 24 in the 1970s and 1980s, with smaller increases or even declines since then (Oltmans et 25 al., 2006; Logan et al., 1999). Over Japan, O_3 in the lower troposphere increased about 26 0.2-0.4 ppb/year during the 1990s (Naja and Akimoto, 2004).
- 27Ground-based measurements in remote regions provide a record of tropospheric O3, but28like ozonesonde data are sparse before the 1970s. Springtime O3 observations from29several mountain sites in the western U.S. show a positive trend of about of 0.5-300.7 ppb/year since the 1980s (Cooper et al., 2010; Jaffe et al., 2003). Ship-borne O331measurements for the time period 1977 to 2002 indicate increases of 0.1-0.7 ppb/year32over much of the Atlantic south of 40°N, but no appreciable change north of 40°N
- 33 (Lelieveld et al., 2004). The lack of trend for the North Atlantic would seem at odds with

1	O_3 observations at Mace Head (53°N) on the west coast of Ireland, which show a
2	significant positive trend of about 0.5 ppb/year from 1987 to 2003 (Simmonds et al.,
3	2004). Over Japan, O_3 at a remote mountain site has increased 1 ppb/year from 1998 to
4	2003 (Tanimoto, 2009), a rate more than double that recorded by ozonesondes in the
5	lower troposphere over Japan during the 1990s (<u>Naja and Akimoto, 2004</u>). At Zugspitze,
6	a mountain site in Germany, O_3 increased by 12% per decade during the 1970s and
7	1980s, consistent with European ozonesondes (Oltmans et al., 2006). Since then, O_3
8	continues to increase at Zugspitze, but more slowly. What little data exist for the
9	Southern Hemisphere point to measurable increases in tropospheric O_3 in recent decades,
10	as much as $\sim 15\%$ at Cape Grim in the 1989-2004 time period (<u>Oltmans et al., 2006</u>).
10	as much as $\sim 15\%$ at Cape Ormi in the 1969-2004 time period (Orthans et al., 2000).
11	The satellite record is now approaching a length that can be useful for diagnosing trends
12	in the total tropospheric O ₃ column (details on the use of satellites to measure
13	tropospheric O_3 are covered in Chapter <u>3</u> , Section <u>3.5.5.5</u>). In contrast to the surface data
14	from ships, tropospheric O_3 columns from the Total Ozone Mapping Spectrometer
15	(TOMS) show no trend over the tropical Atlantic for the period 1980-1990 (Thompson
16	and Hudson, 1999). Over the Pacific, a longer, 25 year record of TOMS data again
17	reveals no trend over the tropics, but shows increases in tropospheric column O_3 of about
18	2-3 Dobson Units $(DU)^1$ at mid-latitudes in both hemispheres (Ziemke et al., 2005).
19	Interpreting these recent trends in tropospheric O_3 is challenging. The first difficulty is
20	reconciling apparently contradictory trends in the observations, e.g., over tropical oceans.
21	A second difficulty is that the O ₃ trends depend on several factors, not all of which can be
22	well characterized. These factors include (1) trends in emissions of O_3 precursors,
23	(2) variation in the stratospheric source of O_3 , (3) changes in solar radiation resulting
24	from stratospheric O_3 depletion, and (4) trends in tropospheric temperatures (Fusco and
25	Logan, 2003). Recent positive trends in the western U.S. and over Japan are consistent
26	with the rapid increase in emissions of O_3 precursors from mainland Asia and transport of
27	pollution across the Pacific (Cooper et al., 2010; Tanimoto, 2009). The satellite trends
28	over the northern mid-latitudes are consistent with this picture as well (Ziemke et al.,
29	<u>2005</u>). Increases in tropospheric O_3 in the Southern Hemisphere are also likely due to
30	increased anthropogenic NO_X emissions, especially from biomass burning (Fishman et
31	<u>al., 1991</u>). Recent declines in summertime O_3 over Europe can be partly explained by
32	decreases in O_3 precursor emissions there (Jonson et al., 2005), while springtime
33	increases at some European sites are likely linked to changes in stratospheric dynamics
34	(Ordonez et al., 2007). Over Canada, Fusco and Logan (2003) found that O_3 depletion in

¹ The Dobson Unit is a typical unit of measure for the total O_3 in a vertical column above the Earth's surface. One DU is equivalent to the amount of O_3 that would exist in a 1 μ m (10⁻⁵ m) thick layer of pure O_3 at standard temperature (0°C) and pressure (1 atm), and corresponds to a column of O_3 containing 2.69 × 10²⁰ molecules/m². A typical value for the amount of ozone in a column of the Earth's atmosphere, although highly variable, is 300 DU and approximately 10% (30 DU) of that exists in the troposphere at mid latitudes.

1	the lowermost stratosphere may have reduced the stratospheric flux of O_3 into the
2	troposphere by as much as 30% from the early 1970s to the mid 1990s, consistent with
3	the trends in ozonesondes there.

Calculation of Ozone Trends for the Recent Past

- Simulations of trends in tropospheric O₃ provide a means for testing current knowledge
 of O₃ processes and predicting with greater confidence trends in future O₃ concentrations.
 Time-dependent emission inventories of O₃ precursors have also been developed for
 1850-2000 (Lamarque et al., 2010) and for 1890-1990 (VanAardenne et al., 2001). These
 inventories allow for the calculation of changing O₃ concentration over time.
- 9 One recent multi-model study calculated an increase in the O₃ concentration since 10 preindustrial times of 8-14 DU, or about 30-70% (Gauss et al., 2006). The large spread in 11 modeled estimates reveals the limitations in knowledge of processes in the pristine 12 atmosphere. Models typically overestimate the late nineteenth and early twentieth century 13 observations available in surface air and at mountain sites by 50-100% (Lamarque et al., 14 2005; Shindell et al., 2003; Mickley et al., 2001; Kiehl et al., 1999). Reconciling the 15 differences between models and measurements will require more accurate simulation of 16 the natural sources of O₃ (Mickley et al., 2001) and/or implementation of novel sinks 17 such as bromine radicals, which may reduce background O_3 in the pristine atmosphere by 18 as much as 30% (Yang et al., 2005c).
- 19 For the more recent past (since 1970), application of time-dependent emissions reveals an 20 equatorward shift in the distribution of tropospheric O_3 in the Northern Hemisphere due 21 to the industrialization of societies at low-latitudes (Lamarque et al., 2005; Berntsen et 22 al., 2000). By constraining a model with historical (1950s-2000) observations, Shindell 23 and Faluvegi (2002) calculated a large increase of 8.2 DU in tropospheric O₃ over 24 polluted continental regions since 1950. This trend is not captured in standard chemistry 25 models, but is consistent with the change in tropospheric O_3 since preindustrial times 26 implied by the observations from the late 1800s (Pavelin et al., 1999; Marenco et al., 27 1994).

10.3.3.2 The Effect of Surface Albedo on Ozone Radiative Forcing

28The Earth's surface albedo plays a role in O3 radiative forcing. Through most of the29troposphere, absorption of incoming shortwave solar radiation by O3 is small relative to30its absorption of outgoing longwave terrestrial radiation. However, over surfaces31characterized by high albedo (e.g., over snow, ice, or desert sand), incoming radiation is

1	more likely to be reflected than over darker surfaces, and the probability that O_3 will
2	absorb shortwave solar radiation is therefore larger. In other words, energy that would
3	otherwise return to space may instead be retained in the atmosphere. Several studies have
4	shown that transport of O ₃ to the Arctic from mid-latitudes leads to radiative forcing
5	estimates greater than 1.0 W/m^2 in the region, especially in summer (Shindell et al., 2006;
6	Liao et al., 2004b; Mickley et al., 1999). Both the high surface albedo of the Arctic and
7	the large solar zenith angles there (which increase the path length of incoming sunlight)
8	lead to strong shortwave forcing in the region. Because the Arctic is especially sensitive
9	to radiative forcing through the ice-albedo feedback, the large contribution in the
10	shortwave solar spectrum to the total radiative forcing in the region may be important.

10.3.3.3 The Effect of Vertical Distribution on Ozone Radiative Forcing

11	In the absence of feedbacks, O_3 increments near the tropopause produce the largest
12	increases in surface temperature (Lacis et al., 1990; Wang et al., 1980). This is a result of
13	the colder temperature of the tropopause relative to the rest of the troposphere and
14	stratosphere. Since radiation emitted by the atmosphere is approximately proportional to
15	the fourth power of its temperature ¹ , the colder the added O_3 is relative to the earth's
16	surface, the weaker the radiation emitted and the greater the "trapping" of longwave
17	radiation in the troposphere.

10.3.3.4 Feedback Factors that Alter the Climate Response to Changes in Ozone Radiative Forcing

18	Estimates of radiative forcing provide a first-order assessment of the effect of
19	tropospheric O_3 on climate. In the atmosphere, climate feedbacks and transport of heat
20	alter the sensitivity of Earth's surface temperature to addition of tropospheric O_3 .
21	Assessment of the full climate response to increases in tropospheric O ₃ requires use of a
22	climate model to simulate these interactions.
23	Due to its short lifetime, O ₃ is heterogeneously distributed through the troposphere. Sharp
24	horizontal gradients exist in the radiative forcing of O ₃ , with the greatest radiative forcing
25	since preindustrial times occurring over the northern mid-latitudes (more on this in
26	Section $10.3.5$ and Section $10.3.6$). If climate feedbacks are particularly powerful, they
27	may obscure or even erase the correlation between regional radiative forcing and climate

¹ As described by the Stefan-Boltzmann law, an ideal blackbody--which the atmosphere approximates--absorbs at all wavelengths and re-radiates proportional to the fourth power of its temperature.

- 1 response (Harvey, 2004; Boer and Yu, 2003). The transport of heat through the 2 atmosphere, though not technically a feedback, may also weaken the correlation between 3 forcing and climate response. Several model studies have reported that the horizontal 4 pattern of surface temperature response from 2000-2100 trends in predicted short-lived 5 species (including O₃) closely matches the pattern from the trends in the long-lived 6 greenhouse gases over the same time period (Levy et al., 2008; Shindell et al., 2008; 7 Shindell et al., 2007). This correspondence occurs even though the patterns of radiative 8 forcing for the short-lived and long-lived species differ substantially. In a separate paper, 9 Shindell et al. (2007) found that Arctic temperatures are especially sensitive to the mid-10 latitude radiative forcing from tropospheric O₃.
- 11 Other studies have found that the signature of warming due to tropospheric O_3 does show 12 some consistency with the O_3 radiative forcing. For example, Mickley et al. (2004) 13 examined the change in O_3 since preindustrial times and found greater warming in the 14 Northern Hemisphere than in the Southern Hemisphere ($+0.4^{\circ}$ C versus $+0.2^{\circ}$ C), as well 15 as higher surface temperatures downwind of Europe and Asia and over the North 16 American interior in summer. For an array of short-lived species including O₃, Shindell 17 and Faluvegi (2009) found that radiative forcing applied over northern mid-latitudes yield more localized responses due to local cloud, water vapor, and albedo feedbacks than 18 19 radiative forcing applied over the tropics.
- 20 Climate feedbacks can also alter the sensitivity of surface temperature to the vertical 21 distribution of tropospheric O_3 . The previous section (Section 10.3.3.3) described the 22 greater effect of O_3 added to the upper troposphere (near the tropopause) on radiative 23 forcing, relative to additions in the mid- to lower troposphere. However, warming 24 induced by increased O_3 in the upper troposphere could stabilize the atmosphere to some 25 extent, limiting the transport of heat to the Earth's surface and mitigating the effect of the 26 added O_3 on surface temperature (Joshi et al., 2003; Christiansen, 1999). Hansen et al. 27 (1997) determined that allowing cloud feedbacks in a climate model meant that O_3 28 enhancements in the mid-troposphere had the greatest effect on surface temperature.
- 29 Finally, climate feedbacks can amplify or diminish the climate response of one 30 greenhouse gas relative to another. For example, Mickley et al. (2004) found a greater 31 temperature response to CO_2 radiative forcing than to an O_3 radiative forcing of similar 32 global mean magnitude, due in part to the relatively weak ice-albedo feedback for O_3 . 33 Since CO_2 absorbs in the same bands as water vapor, CO_2 radiative forcing saturates in 34 the middle troposphere and is also shifted toward the drier poles. A poleward shift in 35 radiative forcing amplifies the ice-albedo feedback in the case of CO_2 , and the greater 36 mid-troposphere radiative forcing allows for greater surface temperature response, 37 relative to that for O_3 .

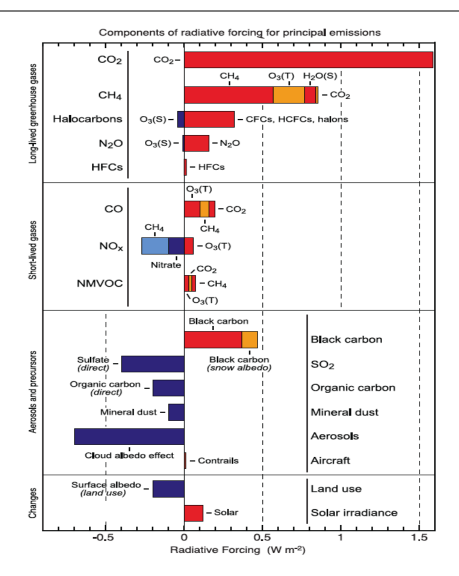
10.3.3.5 Indirect Effects of Tropospheric Ozone on the Carbon Cycle

1 A proposed indirect effect of tropospheric O ₃ on climate involves the carbo	oon cycle. By
2 directly damaging plant life in ways discussed in Chapter 9, increases in tr	ropospheric O ₃
3 may depress the land-carbon sink of CO_2 , leading to accumulation of CO_2	$\frac{1}{2}$ in the
4 atmosphere and ultimately warming of the Earth's surface. Sitch et al. (200	07) calculated
5 that this indirect warming effect of O_3 on climate has about the same magn	nitude as the O ₃
6 direct effect. Their results suggest a doubled sensitivity of surface tempera	atures to O ₃
7 radiative forcing, compared to current model estimates.	

10.3.4 Competing Effects of Ozone Precursors on Climate

8	Changes in O_3 precursors can affect the radiative balance of the atmosphere through
9	multiple (and sometimes competing) mechanisms. For example, the O ₃ precursor CH ₄ is
10	itself a powerful greenhouse gas. O ₃ and its other precursors also exert a strong control on
11	the oxidizing capacity of the troposphere, and so can affect the lifetime of gases such as
12	CH ₄ (Derwent et al., 2001). For example, an increase in CO or VOCs would lead to a
13	decrease in hydroxyl (OH) concentrations. Since OH is a major sink for CH ₄ , a decline in
14	OH would lengthen the CH_4 lifetime, enhance the CH_4 concentration, and amplify
15	surface warming. A rise in NO _X emissions, on the other hand, could lead to an increase in
16	OH in certain locations, shortening the CH ₄ lifetime and causing surface cooling
17	(Fuglestvedt et al., 1999). O_3 can itself generate OH through (1) photolysis leading to
18	excited oxygen atoms followed by reaction with water vapor and (2) reaction with HO_2 .
19	Figure 10-4 shows the radiative forcing associated with a suite of anthropogenic
20	emissions, including O_3 precursors (<u>IPCC, 2007b</u>). The emission-based radiative forcing
21	for CH ₄ , which includes the CH ₄ effect on O_3 production, is +0.9 W/m ² , or nearly double
22	that of the CH ₄ abundance-based radiative forcing shown in Figure 10-3. Figure 10-4 also
23	shows a warming from anthropogenic CO and VOC emissions of $+0.27 \text{ W/m}^2$ and a net
24	cooling of -0.21 W/m^2 for NO _X emissions. The net cooling for NO _X occurs mainly due to
25	the links between NO _X and CH ₄ . Consistent with these results, Shindell and Faluvegi
26	(2009) calculated positive (+0.25 W/m^2) radiative forcing from the increase in
27	anthropogenic emissions of CO and VOCs since preindustrial times, as well as for CH_4
28	(+1 W/m ²). In contrast, <u>Shindell and Faluvegi (2009</u>) found negative (-0.29 W/m ²)
29	radiative forcing from anthropogenic emissions of NO _X . Other studies have found a near
30	cancellation of the positive O_3 radiative forcing and the negative CH_4 radiative forcing
31	that arise from an incremental increase in anthropogenic NO _X emissions (Naik et al.,
32	2005; Fiore et al., 2002; Fuglestvedt et al., 1999). The net effect of aircraft NO _X on
33	climate is especially complex (Isaksen et al., 2001; Wild et al., 2001). Stevenson (2004)

1 2	calculated that aircraft NO_X leads to short-term net warming via O_3 production in the cool upper troposphere, but long-term net cooling because of CH_4 loss.
3	OH production from O ₃ precursors can also affect regional sulfate air quality and climate
4	forcing by increasing gas-phase oxidation rates of SO ₂ . Using the A1B scenario in the
5	IPCC AR4, Unger (2006) reported that by 2030, enhanced OH from the A1B O_3
6	precursors may increase surface sulfate aerosol concentrations by up to 20% over India
7	and China, relative to the present-day, with a corresponding increase in radiative cooling
8	over these regions. In this way, O ₃ precursors may impose an indirect cooling via sulfate
9	(<u>Unger, 2006</u>).
10	Taken together, these results point out the need for careful assessment of net radiative
11	forcing involving multiple pollutants in developing climate change policy (Unger et al.,
12	<u>2008</u>). Many studies point to CH_4 as a particularly attractive target for emissions control
13	since CH_4 is itself an important precursor of O_3 (West et al., 2007; Fiore et al., 2002).
14	Fiore et al. (2002) found that reducing anthropogenic CH_4 emissions by 50% would lead
15	to a global negative (-0.37 W/m^2) radiative forcing, mostly from CH ₄ . In later research,
16	Fiore et al. (2008) reported that CH_4 reductions would most strongly affect tropospheric
17	O_3 column amounts in regions of strong downwelling from the upper troposphere
18	(e.g., around 30°N) and in regions of NO _X -saturated conditions.
19	The magnitude of the radiative forcing from the change in tropospheric O_3 since the
20	preindustrial era is uncertain. This uncertainty derives in part from the scarcity of early
21	measurements and in part from limited knowledge regarding processes in the natural
22	atmosphere. As noted previously, the IPCC AR4 reports a radiative forcing of 0.35 W/m^2
23	from the change in tropospheric O_3 since 1750 (Forster et al., 2007), ranking it third in
24	importance behind the greenhouse gases CO ₂ and CH ₄ . The O ₃ radiative forcing could, in
25	fact, be as large as 0.7 W/m^2 , if reconstructions of preindustrial and mid-20th century O_3
26	based on the measurement record are valid (Shindell and Faluvegi, 2002; Mickley et al.,
27	<u>2001</u>). In any event, <u>Unger et al. (2010</u>) showed that present-day O_3 radiative forcing can
28	be attributed to emissions from many economic sectors, including on-road vehicles,
29	household biofuel, power generation, and biomass burning. As much as one-third of the
30	radiative forcing from the 1890 to 1990 change in tropospheric O_3 could be due to
31	increased biomass burning (<u>Ito et al., 2007a</u>).
32	These calculated radiative forcing estimates can be compared to those obtained from
33	satellite data. Using data from TOMS, Worden et al. (2008) estimated a reduction in
34	clear-sky outgoing longwave radiation of 0.48 W/m^2 by O_3 in the upper troposphere over
35	oceans in 2006. This radiative forcing includes contributions from both anthropogenic
36	and natural O_3 . Assuming that the concentration of O_3 has roughly doubled since
37	preindustrial times (Gauss et al., 2006), the total O_3 radiative forcing estimated with

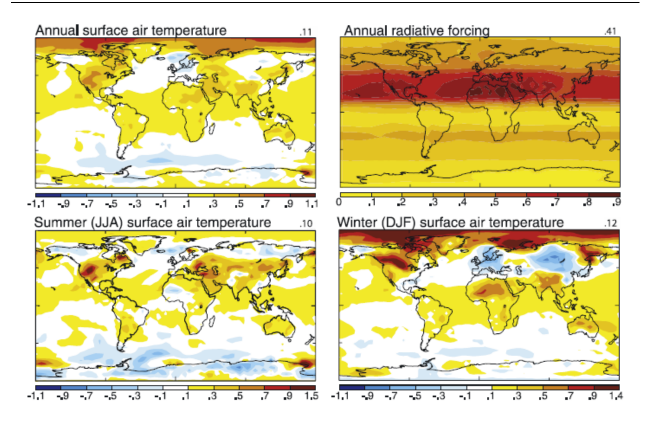


Note: Values represent radiative forcing in 2005 due to emissions and changes since 1750. (S) and (T) next to gas species represent stratospheric and tropospheric changes, respectively. Source: Reprinted with permission of Cambridge University Press (<u>IPCC, 2007b</u>).

Figure 10-4 Components of radiative forcing for emissions of principal gases, aerosols, aerosol precursors, and other changes.

10.3.5 Calculating Radiative Forcing and Climate Response to Past Trends in Tropospheric Ozone

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Note: Figure includes the input radiative forcing (W/m²), as computed by the NASA GISS chemistry-climate model. Values are surface temperature trends for the annual average (top left), June–August (bottom left), and December-February (bottom right) and annual average tropopause instantaneous radiative forcing from 1880 to 1990 (top right). Temperature trends greater than about 0.1°C are significant over the oceans, while values greater than 0.3°C are typically significant over land, except for northern middle and high latitudes during winter where values in excess of about 0.5°C are significant. Values in the top right corner give area-weighted global averages in the same units as the plots.

Source: Reprinted with permission of American Geophysical Union (Shindell et al., 2006).

Figure 10-5 Ensemble average 1900-2000 radiative forcing and surface temperature trends (°C per century) in response to tropospheric ozone changes.

10.3.6 Calculating Radiative Forcing and Climate Response to Future Trends in Tropospheric Ozone

1	Future trends in tropospheric O_3 concentrations depend in large part on what pathways in
2	energy technology the world's societies will follow in coming decades. The trends in O_3
3	will also depend on the changes in a suite of climate-sensitive factors, such as the water
4	vapor content of the atmosphere. This section describes the following issues:
5	(1) projected trends in the anthropogenic emissions of O_3 precursors; (2) the effects of
6	these emissions on the tropospheric O_3 concentrations; (3) the effects of changing climate

on tropospheric O_3 ; and (4) radiative forcing and climate response to 21^{st} century trends in tropospheric O_3 .

10.3.6.1 Emissions of Anthropogenic Ozone Precursors Across the 21st Century

3 The IPCC SRES effort devised scenarios for short-lived O₃ precursors as well as the 4 well-mixed greenhouse gases including NO_x, CO, and VOCs (IPCC, 2000). Using the 5 IMAGE socioeconomic model, Streets et al. (2004) provided speciation for NO_X and 6 VOCs and allocated the trends in emissions over 17 regions and 8 economic sectors for 7 the 2000-2050 time period. The worst-case IPCC scenario, A2, features continued 8 dependence on fossil fuels, rapid population growth, and little sharing of technology 9 between developed and developing nations. By 2100 in this scenario, global NO_X , CO 10 and CH_4 emissions increase by a factor of 3.5, 2.6, and 2.9, respectively, relative to 2000 11 (IPCC, 2000). Most of these increases in emissions occur over developing countries. For 12 example over Asia, NO_x emissions in the A2 scenario increase by more than a factor of 13 four by 2100. The more moderate A1B scenario has global NO_X and CO emissions 14 increasing by 25% and 90%, respectively by 2100, but global CH₄ emissions decreasing 15 by 10%. In the B1 scenario, with its emphasis on clean and efficient technologies, global 16 emissions of NO_x , CO, and CH₄ all decrease by 2100 relative to the present day (-40%, 17 -60%, and -30%, respectively).

- 18Other emissions scenarios have been recently developed to describe trends in the short-19term (up to 2030). The Current Legislation (CLE) scenario provides trends consistent20with existing air quality regulations; the Maximum Feasible Reduction (MFR) scenario21seeks to reduce emissions of O_3 precursors to the maximum extent possible. Emission22source changes relative to the present day for CLE, MFR, and A2 are given in Stevenson23et al. (2006).
- 24 For the Fifth Assessment Report (IPCC AR5), a new set of climate futures has been 25 developed: the Representative Concentration Pathways (RCPs) (Moss et al., 2010). The 26 RCPs will explore for the first time approaches to climate change mitigation. The RCPs are designed to achieve radiative forcing targets of 2.6, 4.5, 6.0 and 8.5 W/m^2 by 2100, 27 28 and have been designated RCP 2.6, RCP 4.5, RCP 6.0, and RCP 8.5, respectively (RCP 29 2.6 is also known as RCP3-PD.) The trends in O₃ precursors for the RCP scenarios were 30 determined by climate policies implicit in each scenario and by plausible assumptions 31 regarding future air quality regulations. These scenarios were chosen to map the wide 32 range of climate outcomes presented in the literature and represent only four of many 33 possible scenarios that would lead to the specific radiative forcing targets; a wide range

1

2

1of socioeconomic conditions could be consistent with each forcing pathway (Moss et al.,22010). Therefore, they should not be interpreted as forecasts of future conditions, but3rather as plausible climate and socio-economic futures.

4 Plots and comparisons of the RCP trends are available on the RCP website (RCP, 2009). In all RCPs, global anthropogenic NO_x emissions decline 30-50% during the 21st century, 5 though RCP 8.5 shows a peak during the 2020s at a value ~15% greater than that of 6 7 2000. Global anthropogenic VOC and CO emissions are relatively flat during the 2000-8 2050 time range, and then decline by 30-50% by the end of the century. For CH₄, global 9 mean emission trends for the four RCP projections differ substantially across the 21st 10 century, with RCP 8.5 showing a tripling of emissions by 2100, and RCP 2.6 showing the 11 emissions cut by half in this time range. RCP 4.5 and 6.0 show a peak in CH₄ emissions 12 in the middle of the century before dropping by the end of the century to just below 2000 13 emission levels. All these global trends, however, contain some regional variation. For 14 example, Asian emissions of both NO_X and VOCs show large increases in the near term 15 (2030s to 2050s).

10.3.6.2 Impact of 21st Century Trends in Emissions on Tropospheric Ozone

16	Due to its short lifetime, tropospheric O ₃ will respond readily to changes in
17	anthropogenic emissions of its precursors. As shown in <u>Table 10-1</u> , a recent multi-model
18	study found increases in the tropospheric O ₃ concentration of 15% and 6% for the IPCC
19	A2 and CLE scenarios respectively for the 2000-2030 time period, and a decrease for the
20	MFR scenario of 5% (Stevenson et al., 2006). These results indicate that the growth in
21	tropospheric O ₃ between 2000 and 2030 could be reduced or even reversed, depending on
22	emission controls. For the relatively moderate A1B emissions scenario over the 2000-
23	2050 time period, <u>Wu et al. (2008a</u>) calculated a change in O_3 concentration of about
24	20%.
25	As noted above, the RCP scenarios show large variations in their future projections of
26	global mean CH ₄ emissions, but mainly declines in the emissions of the other O ₃
27	precursors across the 21st century. In one of the first efforts to assess the effect of these
28	emission trends on global O_3 abundances, <u>Lamarque et al. (2011</u>) found that the large
29	CH ₄ increase in the RCP 8.5 scenario would drive a 15% enhancement of the
30	tropospheric O_3 burden by 2100, relative to the present-day, leading to a global mean
31	radiative forcing of +0.2 W/m^2 . By contrast, the global O ₃ burden would decrease in the
32	other three RCPs, with declines in forcing ranging from -0.07 to -0.2 W/m^2 .

Table 10-12000-2030 changes in anthropogenic emissions, and CH4 and
tropospheric ozone concentrations, and the associated
tropospheric ozone forcing for three scenarios.

Scenario	IPCC A2 ^a	Current Legislation (CLE) ^a	Maximum Feasible Reduction (MFR) ^a	
Percent change in NO _x emissions	+96%	+18%	-53%	
Percent change in CO emissions	+62%	-16%	-53%	
Percent change in CH ₄ concentration	+23%	+19%	0%	
Percent change in tropospheric O ₃ concentration	+15%	+6%	-5%	
Radiative forcing due to O_3 change ^b (W/m ²)	0.3	0.18	-0.05	

^aValues are ensemble means.

^bIncludes radiative forcing due to corresponding CH₄ change.

Source: Adapted from Stevenson et al. (2006).

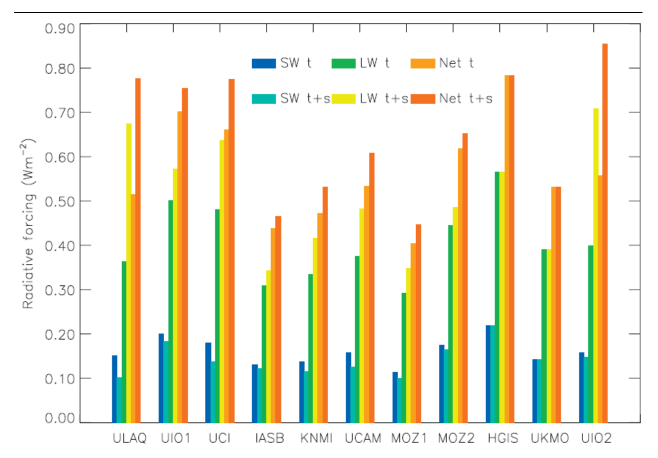
10.3.6.3 Impact of 21st Century Climate on Tropospheric Ozone

1 2 3 4 5 6 7 8	For the time period from the 1800s to the present-day, most of the increase in the concentration of tropospheric O_3 can be traced to changing emissions. Model studies show that climate change so far has likely had little effect on the tropospheric O_3 (e.g., Grenfell et al., 2001). In the future, however, climate change is expected to bring large changes in a suite of variables that could affect O_3 production, loss, and transport. For example, increased water vapor in a warming atmosphere is expected to enhance OH concentrations, which in remote, NO _X -poor regions will accelerate O_3 loss rates (Johnson et al., 1999).
9	In the 2050s A1B climate, <u>Wu et al. (2008b</u>) calculated a 5 ppb decrease in surface O_3
10	over oceans. A rise in temperatures will also likely promote emissions of isoprene, an
11	important biogenic precursor of O ₃ . Model studies have calculated 21 st -century increases
12	in isoprene emissions ranging from 25-50%, depending on climate scenario and time
13	horizon (Wu et al., 2008a and references therein). These studies however did not take
14	into account the effects of changing climate and CO ₂ concentration on vegetation extent,
15	which could have large consequences for biogenic emissions (Heald et al., 2008;
16	Sanderson et al., 2003). In any event, enhanced isoprene emissions will increase O_3
17	concentrations in VOC-limited regions, but decrease O_3 in NO_X -limited regions (<u>Wu et</u>
18	al., 2008a; Pyle et al., 2007; Sanderson et al., 2003). Convection frequencies and
19	lightning flash rates will also likely change in a changing climate, with consequences for
20	lightning NO_X emissions and O_3 concentrations in the upper troposphere (Sinha and
21	Toumi, 1997; Price and Rind, 1994). While Wu et al. (2008a) calculated an increase in
22	lightning NO _X by 2050 due to enhanced deep convection, <u>Jacobson and Streets (2009</u>)

1	projected a decrease in lightning NO_X due to a declining cloud ice in their future
2	atmosphere. Finally, changes in transport processes will almost certainly accompany
3	global climate change. For the 2050 A1B climate, Wu et al. (2008b) showed that
4	flattening of the meridional temperature gradient in a warming world would lead to
5	slower intercontinental transport of tropospheric O ₃ . For the A2 climate in 2100, Zeng
6	and Pyle (2003) projected an 80% increase in the flux of stratospheric O ₃ into the
7	troposphere, relative to the present-day.
8	Taken together, these climate-driven processes could have appreciable effects on the
9	concentration and distribution of tropospheric O ₃ . As shown in <u>Wu et al. (2008b</u>), model
10	projections of the change in O_3 concentration due solely to future climate change range
11	from -12% to $+3\%$, depending on the model, scenario, and time horizon.

10.3.6.4 Radiative Forcing and Climate Response from 21st Century Trends in Tropospheric Ozone

12	In the near term (2000-2030), Stevenson et al. (2006) estimated an O ₃ forcing of near zero
13	for MFR, 0.18 W/m ² for CLE, and +0.3 W/m ² for the A2 scenario (<u>Table 10-1</u>). <u>Menon et</u>
14	al. (2008), following the moderate A1B scenario, calculated a radiative forcing of
15	0.12 W/m^2 from the 2000-2030 change in tropospheric O ₃ , about the same as that derived
16	by Stevenson et al. (2006) for the CLE scenario. Over the longer term (2000 to 2100) for
17	the A1B scenario, Gauss et al. (2003) reported large positive radiative forcing (0.40 to
18	0.78 W/m ²) due to the change in tropospheric O_3 , as shown in Figure 10-6. Normalized
19	radiative forcing for these model calculations fell within a relatively narrow range, 0.032
20	to 0.040 W/m^2 DU, indicating that the largest uncertainty lies in the model-calculated
21	changes in O ₃ concentration. Applying the A2 scenario, Chen et al. (2007b) estimated a
22	global mean radiative forcing of 0.65 W/m^2 from tropospheric O ₃ by 2100, consistent
23	with the Gauss et al. (2003) results. These studies took into account only the effect of
24	changing emissions on tropospheric O_3 . In their calculations of the 2000-2100 radiative
25	forcing from O_3 in the A2 scenario, <u>Liao et al. (2006</u>) found that inclusion of climate
26	effects on tropospheric O_3 reduced their radiative forcing estimate by 20%.



Note: Shown are the components of radiative forcing in W/m2. SW = shortwave component; LW = longwave component; Net = total forcing; t = tropospheric ozone changes only; and t + s = both tropospheric and stratospheric changes. Source: Reprinted from <u>Gauss et al. (2003</u>), American Geophysical Union.

Figure 10-6 Global mean radiative forcing estimates calculated by a set of models for the 2000-2100 change in tropospheric ozone.

1	Several studies have included tropospheric O ₃ in their investigations of the response in
2	the future atmosphere to a suite of short-lived species (e.g., <u>Levy et al., 2008</u> ; <u>Shindell et</u>
3	al., 2008; Shindell et al., 2007). Few studies, however, have calculated the climate
4	response to changes in tropospheric O_3 alone in the future atmosphere. For the A2
5	atmosphere, Chen et al. (2007b) estimated a global mean surface temperature increase of
6	+0.34°C by 2100 in response to the change in O ₃ . The largest temperature increases in
7	this study, as much as 5°C, occurred over the populous regions of Asia and the Middle
8	East and downwind of biomass burning regions in South Africa and South America.

10.4 UV-B Related Effects and Tropospheric Ozone

10.4.1 Background

1	UV radiation emitted from the Sun contains sufficient energy when it reaches the Earth to
2	break (photolyze) chemical bonds in molecules, thereby leading to damaging effects on
3	living organisms and materials. Atmospheric O ₃ plays a crucial role in reducing exposure
4	to solar UV radiation at the Earth's surface. Stratospheric O_3 is responsible for the
5	majority of this shielding effect, as approximately 90% of total atmospheric O ₃ is located
6	there over mid-latitudes (Kar et al., 2010; Crist et al., 1994). Investigation of the
7	supplemental shielding of UV-B radiation provided by tropospheric O ₃ is necessary for
8	quantifying UV-B exposure and the incidence of related human health effects, ecosystem
9	effects, and materials damage. The role of tropospheric O ₃ in shielding of UV-B radiation
10	is discussed in this section.

10.4.2 Human Exposure and Susceptibility to Ultraviolet Radiation

11	The factors that potentially influence UV radiation exposure were discussed in detail in
12	Chapter 10 of the 2006 O_3 AQCD (<u>U.S. EPA, 2006b</u>) and are summarized here. These
13	factors included outdoor activity, occupation, age, gender, geography, and protective
14	behavior. Outdoor activity and occupation both influenced the amount of time people
15	spend outdoors during daylight hours, the predominant factor for exposure to solar UV
16	radiation. Age and gender were found to be factors that influence human exposure to UV
17	radiation, particularly by influencing other factors of exposure such as outdoor activity
18	and risk behavior. Studies indicated that females generally spent less time outdoors and,
19	consequently, had lower UV radiation exposure on average compared to males.
20	Geography influences the degree of solar UV flux to the surface, and hence exposure to
21	UV radiation. Higher solar flux at lower latitudes increased the annual UV radiation dose
22	for people living in southern states relative to northern states. Altitude was also found to
23	influence personal exposure to UV radiation. Protective behaviors such as using
24	sunscreen, wearing protective clothing, and spending time in shaded areas were shown to
25	reduce exposure to UV radiation. Given these and other factors that potentially influence
26	UV radiation exposure, the 2006 O ₃ AQCD (U.S. EPA, 2006b) listed the following
27	subpopulations potentially at risk for higher exposures to UV radiation:
28	 Individuals who engage in high-risk behavior (e.g., sunbathing);
29	 Individuals who participate in outdoor sports and activities;

1 2	 Individuals who work outdoors with inadequate shade (e.g., farmers, construction workers, etc.); and
3 4	 Individuals living in geographic areas with higher solar flux including lower latitudes (e.g., Honolulu, HI) and higher altitudes (e.g., Denver, CO).
5 6	The risks associated with all these factors are, of course, highly dependent on season and region (<u>Sliney and Wengraitis, 2006</u>).

10.4.3 Human Health Effects due to UV-B Radiation

7	Chapter 10 of the 2006 O_3 AQCD (U.S. EPA, 2006b) covered in detail the human health
8	effects associated with solar UV-B radiation exposure. These effects include erythema,
9	skin cancer, ocular damage, and immune system suppression. These adverse effects,
10	along with protective effects of UV radiation through increased production of vitamin D
11	are summarized in this section. For additional details, the reader is referred to Chapter 10
12	of the 2006 O_3 AQCD (U.S. EPA, 2006b) and references therein.
13	The most conspicuous and well-recognized acute response to UV radiation is erythema,
14	or the reddening of the skin. Erythema is likely caused by direct damage to DNA by UV
15	radiation. Many studies discussed in Chapter 10 of the 2006 O ₃ AQCD (U.S. EPA,
16	2006b) found skin type to be a significant risk factor for erythema. Skin cancer is another
17	prevalent health effect associated with UV radiation. Exposure to UV radiation is
18	considered to be a major risk factor for all forms of skin cancer. Ocular damage from UV
19	radiation exposure includes effects on the cornea, lens, iris, and associated epithelial and
20	conjunctival tissues. The region of the eye affected by exposure to UV radiation depends
21	on the wavelength of the incident UV radiation. Depending on wavelength, common
22	health effects associated with UV radiation include photokeratitis (snow blindness; short
23	wavelengths) and cataracts (opacity of the lens; long wavelengths).
24	Experimental studies reviewed in Chapter 10 of the 2006 O ₃ AQCD (U.S. EPA, 2006b)
25	have shown that exposure to UV radiation may suppress local and systemic immune
26	responses to a variety of antigens. Results from human clinical studies suggest that
27	immune suppression induced by UV radiation may be a risk factor contributing to skin
28	cancer induction. There is also evidence that UV radiation has indirect involvement in
29	viral oncogenesis through the human papillomavirus, dermatomyositis, human
30	immunodeficiency virus and other forms of immunosuppression.
31	A potential health benefit of increased UV-B exposure relates to the production of
32	vitamin D in humans. Most humans depend on sun exposure to satisfy their requirements
33	for vitamin D. Vitamin D deficiency can cause metabolic bone disease among children

1	and adults, and also may increase the risk of many common chronic diseases, including
2	type I diabetes mellitus and rheumatoid arthritis. Substantial in vitro and toxicological
3	evidence also support a role for vitamin D activity against the incidence or progression of
4	various forms of cancer. In some studies, UV-B related production of vitamin D had
5	potential beneficial immunomodulatory effects on multiple sclerosis, insulin-dependent
6	diabetes mellitus, and rheumatoid arthritis. More details on UV-B protective studies are
7	provided in Chapter 10 of the 2006 O_3 AQCD (U.S. EPA, 2006b).
8	In establishing guidelines on limits of exposure to UV radiation, the International
9	commission on Non-ionizing Radiation Protection (ICNIRP) agreed that some low-level
10	exposure to UV radiation has health benefits (ICNIRP, 2004). However, the adverse
11	health effects of higher UV exposures necessitated the development of exposure limits
12	for UV radiation. The ICNIRP recognized the challenge in establishing exposure limits
13	that would achieve a realistic balance between beneficial and adverse health effects. As
14	concluded by ICNIRP (2004), "[t]he present understanding of injury mechanisms and
15	long-term effects of exposure to [UV radiation] is incomplete, and awaits further
16	research."

10.4.4 Ecosystem and Materials Damage Effects Due to UV-B Radiation

17	A 2009 progress report on the environmental effects of O_3 depletion from the UNEP,
18	Environmental Effects Assessment Panel (UNEP, 2009) lists many ecosystem and
19	materials damage effects from UV-B radiation. An in-depth assessment of the global
20	ecosystem and materials damage effects from UV-B radiation per se is out of the scope of
21	this assessment. However, a brief summary of some mid-latitude effects is provided in
22	this section to provide context for UV-B related issues pertaining to tropospheric O ₃ . The
23	reader is referred to the UNEP report (UNEP, 2009) and references therein for further
24	details. All of these UV-B related ecosystem and materials effects can also be influenced
25	by climate change through temperature and other meteorological alterations, making
26	quantifiable predictions of UV-B effects difficult.
27	Terrestrial ecosystem effects from increased UV-B radiation include reduced plant
28	productivity and plant cover, changes in biodiversity, susceptibility to infection, and
29	increases in natural UV protective responses. In general, however, these effects are small
30	for moderate UV-B increases at mid-latitudes. A field study on wheat in southern Chile
31	found no substantial changes in crop yield with moderate increases in UV-B radiation
32	(Calderini et al., 2008). Similarly, field studies on silver birch (Betula pendula) in
33	Einland found no massurable offects in photosymphotic function with increases in LIV D
	Finland found no measurable effects in photosynthetic function with increases in UV-B

have also been linked to increases in UV-B radiation (Mazza et al., 2010; Obara et al.,
 2008; Wahl, 2008). Some plants have natural coping mechanisms for dealing with
 changes in UV-B radiation (Favory et al., 2009; Jenkins, 2009; Brown and Jenkins, 2008;
 Ioki et al., 2008), but these defenses may have costs in terms of reduced growth (Snell et
 al., 2009; Clarke and Robinson, 2008; Semerdjieva et al., 2003; Phoenix et al., 2000).

6 Aquatic ecosystem effects from increased UV-B radiation include sensitivity in 7 growth, immune response, and behavioral patterns of aquatic organisms. One study 8 looking at coccolithophores, an abundant phytoplankton group, found a 25% reduction in 9 cellular growth with UV-B exposure (Gao et al., 2009a). Exposure to relevant levels of 10 UV-B radiation has been shown to modify immune response, blood chemistry, and 11 behavior in certain species of fish (Markkula et al., 2009; Holtby and Bothwell, 2008; 12 Jokinen et al., 2008). Adverse effects on growth and development from UV-B radiation 13 have also been observed for amphibians, sea urchins, mollusks, corals, and zooplankton 14 (Garcia et al., 2009; Romansic et al., 2009; Croteau et al., 2008b; Croteau et al., 2008a; 15 Marquis et al., 2008; Marquis and Miaud, 2008; Oromi et al., 2008). Increases in the flux 16 of UV-B radiation may also result in an increase in the catalysis of trace metals including 17 mercury, particularly in clear oligotrophic lakes with low levels of dissolved organic 18 carbon to stop the penetration of UV-B radiation (Schindler et al., 1996). This could then 19 alter the mobility of trace metals including the potential for increased mercury 20 volatilization and transport within and among ecosystems.

- 21 **Biogeochemical cycles**, particularly the carbon cycle, can also be influenced by 22 increased UV-B radiation. A study on high latitude wetlands found UV-induced increases 23 in CO₂ uptake through soil respiration (Haapala et al., 2009) while studies on arid 24 terrestrial ecosystems found evidence for UV-induced release of CO₂ through 25 photodegradation of above-ground plant litter (Brandt et al., 2009; Henry et al., 2008; 26 Caldwell et al., 2007; Zepp et al., 2007). Changes in solar UV radiation may also have 27 effects on carbon cycling and CO₂ uptake in the oceans (Brewer and Peltzer, 2009; 28 Meador et al., 2009; Fritz et al., 2008; Zepp et al., 2008; Hader et al., 2007) as well as 29 release of dissolved organic matter from sediment and algae (Mayer et al., 2009; 30 Riggsbee et al., 2008). Additional studies showing effects on these and additional 31 biogeochemical cycles including the water cycle and halocarbon cycle can be found in 32 the UNEP report (UNEP, 2009) and references therein.
- 33 Materials damage from increased UV-B radiation include UV-induced
- 34photodegradation of wood (Kataoka et al., 2007) and plastics (Pickett et al., 2008). These35studies and others summarizing photo-resistant coatings and materials designed to reduce36photodegradation of materials are summarized in the UNEP report (UNEP, 2009) and37references therein.

10.4.5 UV-B Related Effects Associated with Changes in Tropospheric Ozone Concentrations

1	There are multiple complexities in attempting to quantify the relationship between
2	changes in tropospheric O_3 concentrations and UV-B exposure. The 2006 O_3 AQCD
3	(<u>U.S. EPA, 2006b</u>) described a handful of studies addressing this relationship, but none
4	reported quantifiable effects of tropospheric O_3 concentration fluctuations on UV-B
5	exposure at the surface. Further quantifying the relationship between UV-B exposure and
6	health or welfare effects is complicated by the uncertainties involved in the selection of
0 7	an action spectrum and appropriate characterization of dose (e.g., peak or cumulative
8	levels of exposure, timing of exposures, etc.) The lack of published studies that critically
9	examined these issues togetherthat is the incremental health or welfare effects
10	attributable specifically to UV-B changes resulting from changes in tropospheric O_3
10	
11	concentrationslead to the prior conclusion that the effect of changes in surface-level O_3
	concentrations on UV-induced health outcomes could not be critically assessed within
13	reasonable uncertainty (U.S. EPA, 2006b). ¹
14	A recent study by Madronich et al. (2011) used CMAQ to estimate UV radiation
15	response to changes in tropospheric O ₃ under different control scenarios projected out to
16	2020. This study focused on southeastern U.S. and accounted for spatial and temporal
17	variation in tropospheric O ₃ reductions, an important consideration since most controls
18	are focused on reducing O_3 in populated urban areas. The contrasting control strategies
19	considered in this study included a historical scenario designed to meet an 84 ppb 8-h
20	daily max standard and a reduced scenario designed to bring areas predicted to exceed a
21	similarly designed 70 ppb standard into attainment. A biologically effective irradiance
22	was estimated by multiplying the modeled UV irradiance by a sensitivity function (action
23	spectrum) for the induction of nonmelanoma skin cancer in mice corrected for human
24	skin transmission, then integrating over UV wavelengths. The average relative change in
25	skin cancer-weighted surface UV radiation between the two scenarios was $0.11 \pm 0.03\%$
26	over June, July and August. Weighting by population, this estimate increased to
27	$0.19 \pm 0.06\%$. Madronich et al. (2011) report that their estimated UV radiation increment
28	is an order of magnitude less than that reported in an earlier study by Lutter and Wolz
29	(1997) with the main reason for the discrepancy coming from the unrealistic uniform
30	10 ppb reduction in O_3 assumed in the former study. <u>Madronich et al. (2011</u>) did not
31	attempt to link their predicted increase in UV radiation to a predicted increase in skin
32	cancer incidence, however, due to several remaining and substantial uncertainties.
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¹ The reader is referred to the U.S. EPA 2003 Final Response to Court Remand (<u>U.S. EPA, 2003</u>) for detailed discussions of the data and scientific issues associated with the determination of public health benefits resulting from the attenuation of UV-B by surface-level O_3 .

1	Quantitatively estimating human health and welfare effects directly attributed to changes
2	in UV-B penetration resulting from changes in ground-level O3 concentrations will
3	require both (a) a solid understanding of the multiple factors that define the extent of
4	exposure to UV-B, and (b) well-defined and quantifiable links between UV-B exposure
5	and human disease and welfare effects. Detailed information does not exist regarding the
6	relevant type (e.g., peak or cumulative) and time period (e.g., developmental, lifetime, or
7	current) of exposure, wavelength dependency of biological responses, and
8	inter-individual variability in UV resistance.
9	Although the UV-B related health effects attributed to marginal reductions in
	e
10	tropospheric or ground-level O_3 have not been directly assessed to date, they would be
10 11	
	tropospheric or ground-level O ₃ have not been directly assessed to date, they would be
11	tropospheric or ground-level O_3 have not been directly assessed to date, they would be expected to be small based on current information indicating a negligibly small effect of
11 12	tropospheric or ground-level O_3 have not been directly assessed to date, they would be expected to be small based on current information indicating a negligibly small effect of potential future changes in tropospheric O_3 concentrations on ground-level UV-B

10.5 Summary and Causal Determinations

10.5.1 Summary of the Effects of Tropospheric Ozone on Climate

16	Tropospheric O ₃ is a major greenhouse gas, third in importance after CO ₂ and CH ₄ . While
17	the developed world has successfully reduced emissions of O ₃ precursors in recent
18	decades, many developing countries have experienced large increases in precursor
19	emissions and these trends are expected to continue, at least in the near term. Projections
20	of radiative forcing due to changing O3 over the 21st century show wide variation, due in
21	large part to the uncertainty of future emissions of source gases. In the near-term (2000-
22	2030), projections of O_3 radiative forcing range from near zero to +0.3 W/m ² , depending
23	on the emissions scenario (Stevenson et al., 2006). Reduction of tropospheric O_3
24	concentrations could therefore provide an important means to slow climate change in
25	addition to the added benefit of improving surface air quality.
26	It is clear that increases in tropospheric O_3 lead to warming. However the precursors of
27	O ₃ also have competing effects on the greenhouse gas CH ₄ , complicating emissions
28	reduction strategies. A decrease in CO or VOC emissions would enhance OH
29	concentrations, shortening the lifetime of CH_4 , while a decrease in NO_X emissions could
30	depress OH concentrations in certain regions and lengthen the CH ₄ lifetime.

1 Abatement of CH4 emissions would likely provide the most straightforward means to 2 address climate change since CH4 is itself an important precursor of background O3 3 (West et al., 2007; West et al., 2006; Fiore et al., 2002). A reduction of CH4 emissions 4 would also improve air quality on its own right. A set of global abatement measures 5 identified by West and Fiore (2005) could reduce CH4 emissions by 10% at a cost 6 savings, decrease background O3 by about 1 pb in the Northern Hemisphere summer, 7 and lead to a global net cooling of 0.12 W/m ² . West et al. (2007) explored further the 8 benefits of CH4 abatement, finding that a 20% reduction in global CH4 emissions would 9 lead to greater cooling per unit reduction in surface O3, compared to 20% reductions in 10 VOCs or CO. 11 Important uncertainties remain regarding the effect of tropospheric O3 on future climate 12 change. To address these uncertainties, further research is needed to: (1) improve 13 knowledge of the natural atmosphere; (2) interpret observed trends of O3 in the free 14 troposphere and remote regions; (3) improve understanding of the CH4, budget, especially 15 emissions reductions that would act to limit future climate change. 16 between regional O3 radiative forcing		
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9lead to greater cooling per unit reduction in surface O3, compared to 20% reductions in VOCs or CO.11Important uncertainties remain regarding the effect of tropospheric O3 on future climate change. To address these uncertainties, further research is needed to: (1) improve knowledge of the natural atmosphere; (2) interpret observed trends of O3 in the free troposphere and remote regions; (3) improve understanding of the CH4 budget, especially emissions from wetlands and agricultural sources, (4) understand the relationship between regional O3 radiative forcing and regional climate change; and (5) determine the optimal mix of emissions reductions that would act to limit future climate change.18The effect of the tropospheric O3 change since preindustrial times on climate has been estimated to be about 25-40% of anthropogenic CO2 effect and about 75% of anthropogenic CH4 effect according to the IPCC. There are large uncertainties in the radiative forcing estimate attributed to tropospheric O3, making the effect of tropospheric O3 on climate more uncertain than the effect of the long-lived greenhouse gases. Overall, the evidence supports a causal relationship between changes in tropospheric O3 the evidence supports a causal relationship between changes in tropospheric O3 concentrations and radiative forcing.25Radiative forcing does not take into account the climate feedbacks that could amplify or dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given radiative forcing is highly uncertain and can vary greatly among models and from region to region within the same model. In light of thes	7	and lead to a global net cooling of 0.12 W/m ² . West et al. (2007) explored further the
10VOCs or CO.11Important uncertainties remain regarding the effect of tropospheric O3 on future climate change. To address these uncertainties, further research is needed to: (1) improve knowledge of the natural atmosphere; (2) interpret observed trends of O3 in the free troposphere and remote regions; (3) improve understanding of the CH4 budget, especially emissions from wetlands and agricultural sources, (4) understand the relationship between regional O3 radiative forcing and regional climate change; and (5) determine the optimal mix of emissions reductions that would act to limit future climate change.18The effect of the tropospheric O3 change since preindustrial times on climate has been estimated to be about 25-40% of anthropogenic CO2 effect and about 75% of anthropogenic CH4 effect according to the IPCC. There are large uncertainties in the radiative forcing estimate attributed to tropospheric O3, making the effect of tropospheric O3 on climate more uncertain than the effect of the long-lived greenhouse gases. Overall, the evidence supports a causal relationship between changes in tropospheric O3 concentrations and radiative forcing.25Radiative forcing does not take into account the climate feedbacks that could amplify or dampen the actual surface temperature response. Quantifying the change in surface temperature requires a complex climate simulation in which all important feedbacks and interactions are accounted for. As these processes are not well understood or easily modeled, the surface temperature response to a given radiative forcing is highly uncertain and can vary greatly among models and from region to region within the same model. In light of these uncertainties, the evidence indicates that there is likely to be a causal acausal acausal relationship between changes in tropospheric o3	8	benefits of CH ₄ abatement, finding that a 20% reduction in global CH ₄ emissions would
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30and can vary greatly among models and from region to region within the same model. In31light of these uncertainties, the evidence indicates that there is likely to be a causal32relationship between changes in tropospheric O3 concentrations and effects on	28	interactions are accounted for. As these processes are not well understood or easily
 31 light of these uncertainties, the evidence indicates that there is likely to be a causal 32 relationship between changes in tropospheric O₃ concentrations and effects on 	29	modeled, the surface temperature response to a given radiative forcing is highly uncertain
32 relationship between changes in tropospheric O ₃ concentrations and effects on	30	and can vary greatly among models and from region to region within the same model. In
	31	light of these uncertainties, the evidence indicates that there is likely to be a causal
33 climate.		relationship between changes in tropospheric O_3 concentrations and effects on
	33	climate.

10.5.2 Summary of UV-B Related Effects on Human Health, Ecosystems, and Materials Relating to Changes in Tropospheric Ozone Concentrations

1	UV radiation emitted from the Sun contains sufficient energy when it reaches the Earth to
2	break (photolyze) chemical bonds in molecules, thereby leading to damaging effects on
3	living organisms and materials. Atmospheric O3 plays a crucial role in reducing exposure
4	to solar UV radiation at the Earth's surface. Ozone in the stratosphere is responsible for
5	the majority of this shielding effect, as approximately 90% of total atmospheric O_3 is
6	located there over mid-latitudes. Ozone in the troposphere provides supplemental
7	shielding of radiation in the wavelength band from 280-315 nm, referred to as UV-B
8	radiation. UV-B radiation has important effects on human health and ecosystems, and is
9	associated with materials damage.

10There is a lack of published studies that critically examine the incremental health or11welfare effects (adverse or beneficial) attributable specifically to changes in UV-B12exposure resulting from perturbations in tropospheric O3 concentrations. While the13effects are expected to be small, they cannot yet be critically assessed within reasonable14uncertainty. Overall, the evidence is inadequate to determine if a causal relationship15exists between changes in tropospheric O3 concentrations and effects on health16and welfare related to UV-B shielding.

10.5.3 Summary of Ozone Causal Determinations

17	The evidence reviewed in this chapter describes the recent findings regarding the climate
18	and UV-B related effects of changes in tropospheric O_3 concentrations. <u>Table 10-2</u>
19	provides an overview of the causal determinations for each of the categories evaluated.

Table 10-2Summary of ozone causal determinations for climate and UV-B
effects.

Effects	Causal Determination
Radiative Forcing	Causal relationship
Climate Change	Likely to be a causal relationship
Health and Welfare Effects Related to UV-B Shielding	Inadequate to determine if a causal relationship exists

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