Appendix C

Key Particulate Matter (PM) Epidemiologic Findings Related to PM NAAQS Decisions

- C.1 Overview of Key Findings Supporting 1997 PM NAAQS Decisions
- C.2 Prospective Cohort Studies of Long-Term Ambient PM Exposure Effects

1 C.1. Overview of Key Findings Supporting 1997 PM NAAQS Decisions

2 In promulgating the 1997 PM NAAQS (Federal Register, 1997), EPA relied mainly on the 3 relative risk (RR) levels for increased risks of mortality or morbidity associated with acute 4 (short-term) and chronic long-term measures of PM exposure reported in U.S. and Canadian PM 5 epidemiology studies, which provide the most directly pertinent quantitative risk estimates as inputs to U.S. PM NAAQS decisions. These included (a) relative risk (RR) estimates for 6 mortality or morbidity associated with 50 μ g/m³ increases in 24-h PM₁₀ concentrations (Table C-7 1) or with variable increases in fine particle indicators, e.g., 25 μ g/m³ increment in 24-h PM₂₅ 8 concentrations (Table C-2); and (b) analogous relative risk estimates for health effects related to 9 10 specified increments in long-term (e.g., annual mean or median) levels of fine particle indicators 11 (Table C-3). The study results summarized in these tables reproduced from Chapter 13 of the PM CD (U.S. EPA, 1996a)¹ were found to provide sufficient evidence for concluding that 12 13 significant associations of increased mortality and morbidity risks were likely attributable to fine 14 particles, as indexed by various fine particle indicators, e.g., PM₂₅, sulfates (SO₄), etc.; but 15 possible toxic effects of the coarse fraction of PM₁₀ (i.e., PM_{10-2.5}) could not be ruled out. Some inhalable coarse fraction particles subsumed under PM₁₀ do reach the lower respiratory tract, and 16 17 some health effects of concern are suggested by some epidemiology results.

18 Both the PM CD (U.S. EPA, 1996a) and Staff Paper (U.S. EPA, 1996b) noted the very 19 limited extent of available toxicologic findings by which (a) to identify key PM constituents of 20 urban ambient air mixes that may be causally related to mortality/morbidity effects observed in the 21 community epidemiologic studies; or (b) to delineate plausible biological mechanisms by which 22 such effects could be induced at the relatively low ambient PM concentrations evaluated in the epidemiologic studies. As discussed in the PM CD, several types of mechanisms have been shown 23 24 to underlie toxic effects observed with acute or chronic exposures to various PM species or 25 mixtures (e.g., acute lung inflammation; impaired respiratory function; impaired pulmonary 26 defense mechanisms, etc.), but generally at much higher PM levels than now typically 27 encountered in U.S. ambient air. As also discussed in the 1996 PM CD, several fine particle 28 constituents were hypothesized as being likely important contributors to ambient PM effects, e.g., 29 acid aerosols (indexed by sulfates; H+ ions, etc.); transition metals (e.g., Fe, Mn, etc.); and 30 ultrafine particles. Nevertheless, despite the lack of more definitive characterization of pertinent 31 underlying biological mechanisms, several aspects of the epidemiologic evidence (e.g., the 32 consistency and coherence of the epidemiologic findings), as discussed in the PM CD, support the 33 conclusion that exposure to ambient PM, acting alone or in combination with other air pollutants,

¹Full reference citations for each study identified in Tables C-1, C-2, and C-3 can be obtained in the bibliographic listing for Chapter 13 in U.S. EPA (1996a).

- is probably a key causal agent contributing to the increased mortality and morbidity risks observed
 in the epidemiology studies. Figure C-1, from the PM Staff Paper (1996b), illustrates the
- 3 consistency and coherence of the relative risk findings for PM_{10} .
- 4 Relative risk estimates shown in Table C-2 for mortality and morbidity effects associated 5 with short-term ambient PM exposures provided the key bases for derivation of the new 65 μ g/m³ PM_{2.5} (24-hr) NAAQS set by EPA in 1997 to protect sensitive human population groups from 6 7 adverse effects of short-term exposures to fine particles. Of particular importance in 8 substantiating the need for fine particle standards were analyses of Harvard Six City Study data 9 reported by Schwartz et al. (1996a) showing stronger, more consistently statistically significant, 10 associations between acute (24-h) PM_{2.5} concentrations and increased mortality risks than for 24-h concentrations of inhalable coarse fraction particles (PM_{15-2.5}) in the same cities (see 11 12 Figure C-2).
- 13 However, as indicated in Chapter 5 of this document, there is little evidence substantiating 14 the occurrence of health effects due to acute (<24-hr) exposures to diesel emissions containing 15 DPM at ambient or near-ambient concentrations. Note that 300 μ g/m³ is the lowest DPM 16 concentration at which mild irritation and inflammation of respiratory tract tissues (but not pulmonary function decrements) were observed with 1-hr controlled human exposures of healthy 17 18 adult volunteers to diesel exhaust (see Chapter 5). In contrast, various noncancer (respiratory 19 system) effects have been shown to occur in numerous mammalian species as the result of 20 controlled long-term (subchronic, chronic) exposures to DPM. Thus, key elements forming the basis for derivation of the 15 μ g/m³ PM_{2.5} annual-average NAAQS set in 1997 to protect against 21 22 health effects associated with long-term fine particle exposures are far more germane here in 23 attempting to relate ambient fine particle health risk estimates to potential ambient DPM exposure 24 risks.
- As noted in Chapter 6 of this document, the derivation of the 15 μ g/m³ PM_{2.5} annualaverage standard was based, in part, on the assumption that increased mortality and morbidity effects associated with acute (24-h) PM_{2.5} exposures were most likely due to PM_{2.5} concentrations above the annual mean values for the cities evaluated. Also, it was noted in Chapter 6 that annual mean PM_{2.5} values typically exceeded 15 μ g/m³ for cities where 24-h PM_{2.5} levels were found to be statistically significantly related to increased mortality and/or morbidity risks, as shown by several key studies (Schwartz et al., 1996; Thurston et al., 1994; Neas et al., 1995).
- Other key elements contributing to the derivation of the annual average PM_{2.5} NAAQS
 were several new prospective cohort studies (published in the 1990's) that evaluated associations
 between long-term exposures to ambient PM and increased risks of mortality or morbidity. The
 most salient points of the PM CD (U.S. EPA, 1996a) assessment of such prospective cohort

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C.2. Prospective Cohort Studies of Long-Term Ambient PM Exposure Effects

studies are summarized in Section C.2 below. These are augmented by discussion of pertinent

findings from recent new follow-up analyses for one of the subject prospective cohort studies.

Newer prospective cohort studies (Abbey et al., 1991; Dockery et al., 1993; and Pope
et al., 1995) were considered in the PM CD (1996a) as providing more credible evidence on
PM-health effects relationships than numerous previous cross-sectional studies. Salient features
of those three key prospective studies are summarized in Table C-4 (reproduced from Chapter 12
of the 1996 PM CD).

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C.2.1. Harvard Six U.S. Cities Study

12 Dockery et al. (1993) analyzed survival probabilities among 8,111 adults first recruited in 13 the mid-1970s in mid-western and eastern U.S. cities, including: Topeka, KS; Portage, WI (a 14 small town north of Madison); St. Louis, MO; Steubenville, OH; (an industrial community on 15 W. VA-PA border); Kingston-Harriman, TN (small towns southwest of Knoxville) and Watertown, MA (western suburb of Boston). These locations comprise a transect across the 16 17 Northcentral and Northeastern United States, from the upper Midwest through Appalachia, to 18 suburban Boston. In each community, about 2,500 adults (white, aged 25 to 74, at enrollment) 19 were selected randomly, but the final cohorts numbered 1,400 to 1,800 persons in each city. 20 Follow-up periods ranged from 14 to 16 years, during which 13 to 22% of the enrollees died. 21 Of the 1,430 death certificates, 98% of the decedents were located, including persons who had 22 moved away and died elsewhere, but no information was provided on actual locations of death. 23 The analyses reported were mainly based on all-cause mortality; no mention was made of 24 subtracting external causes.

25 Air monitoring data obtained from routine sampling stations and special instruments set up 26 by the research team were used. Individual characteristics of the cohort subjects (and thus of the 27 decedents) considered in statistical analyses included: smoking habits, an index of occupational 28 exposure, body mass index, and completion of high school education. The Cox proportional 29 hazards model was used to estimate coefficients for individual risk factors after stratifying by gender and age (5-year groups). The effects of air pollution were evaluated (a) by estimating the 30 31 relative risks of residence in each city relative to Portage (the city with the lowest pollution levels 32 for most indices) and (b) by including the community-average air quality levels directly in the 33 models. Since only six different long-term average values were available for each pollutant, the 34 effective degrees of freedom are small. Most of the air quality measures were averaged over the 35 period of study, in an effort to study long-term (chronic) exposure effects; the specific averaging

1 periods varied by pollutant. Steubenville, Kingston-Harriman, and St. Louis were the most

2 polluted cities and also had the oldest and least educated cohorts and the heaviest rates of3 smoking among the six cities.

4 No consideration was given to possible independent effects of occupation classification, other personal lifestyle variables such as diet or physical activity, migration, or income. 5 6 Presumably, each subject was characterized by his status at entry to the study; follow-up data on 7 possible changes in risk factors over time were not mentioned. Since the air quality data used in 8 this study were largely obtained from "private" monitoring rather than from public archives, 9 comparisons of the average levels with routine monitoring data were of some interest; and no 10 serious disagreements were found, except that it might have been preferable to consider peak rather than average levels of ozone, as is more typical in most studies of acute O₃ effects on 11 12 mortality. Also, it is notable that collection of size-classified PM data began in 1980, whereas 13 TSP data began in 1974 and from 1974 to 1980 there were large reductions in TSP (and likely the 14 size-classified particles as well), so that the size-classified data may be less representative than 15 TSP of cumulative exposures. Sulfate appeared to be intermediate in this regard.

A more complete breakdown of relative risk estimates by city, sex, smoking status, education, and body mass index is given in Table C-5. The mean PM_{2.5} values are provided for reference, but the adjusted relative risks used only age, smoking, education, and body mass as covariates. The RR values for men and women combined are plotted in Figure C-3 for each pollutant. Note that the apparently linear relationship between fine particles and risk is less linear if plotted separately for men and for women, and the confidence intervals also become wider due to smaller sample sizes.

23 Substantial differences in survival rates (expected based on statewide mortality data) were 24 observed across the study's transect of the Northcentral and Northeastern U.S. The long-term 25 average mortality rate in Topeka was 9.7 deaths per 1,000 person-years and in Steubenville was 16.2, yielding a range in average (crude) relative risk of 67% among the six cities. After 26 27 individual adjustment for age, smoking status, education, and body-mass index, the range in 28 average relative risk was reduced to 26%. The relative importance of adjustments for age, 29 smoking, education, and body mass in determining the final ranks of the cities may be seen from 30 the Table C-5. Also, there is more scatter for men and women separately than when combined, 31 presumably because of the reduction in sample size.

Dockery et al. (1993) report that "mortality was more strongly associated with the levels
 of fine, inhalable, and sulfate particles" than with the other pollutants, which they attributed

1 primarily to factors of particle size. They provided relative risk estimates and confidence limits 2 based on the differences between air quality in Steubenville and in Portage for these three PM 3 indicators. However, it is relatively simple to independently estimate coefficients from the adjusted risks and pollutants levels in each of the six communities. These estimates obtained (see 4 5 Table C-6) correspond well to those of Dockery et al. (1993), based on output from the Cox 6 proportional hazards model. However, because there are only 6 different values for the air quality 7 data, the resulting confidence limits are considerably wider than those for the risk factors having 8 individual data. The estimates given in Table C-6, allow comparisons of results for various 9 pollutants and combination of pollutants. As in the original paper, the relative risks are based on 10 the difference in air pollution between Steubenville and Portage. The data for 1970 TSP (corresponding to a lag of about 12 years) were obtained from Lipfert (1978), assuming that 11 12 Madison could represent Portage, WI, as was done in the analysis of Schwartz et al. (1996b).

13 Table C-6 shows only small differences among many pollutants, including SO₂ and NO₂, owing in part to the strong collinearity present. Note that relative risk elevations for the PM₁₅ and 14 15 fine particle indicators (PM_{25} , SO_4) were statistically significant. The non-sulfate portion of PM_{25} 16 had the tightest confidence limits. In contrast, TSP and the coarse particle variables created by 17 subtracting PM₁₅ from TSP and PM₂₅ from PM₁₅ were not significant, suggesting that particles 18 \geq 15 μ m in aerodynamic diameter may be less important; this outcome may reflect in part greater 19 spatial variability within the communities for coarse versus fine particles. Note also that the 20 estimated 1970 TSP variable performed slightly better than the TSP data (ca. 1982) used by 21 Dockery et al., thus suggesting a role for previous pollution exposure. Dockery et al. noted that 22 mean ozone levels varied little among cities; but this may have been less so if a measure of peak 23 (e.g., 1- or 8-hr) O₃ levels had been used instead of daily (24-h) averages. Also, no relationship 24 was found for aerosol acidity (H⁺), but only limited data were available. Both sulfate and 25 non-sulfate fine particles effects seem rather similar, as shown in Figure C-2, making it plausible 26 that there may be PM effects related to particle size independent of sulfate content or particle 27 acidity.

In comparing the most and least polluted cities, Dockery et al. also reported elevated risks for cardiopulmonary causes (RR 1.37; 95% CL 1.11 to 1.68) and lung cancer (RR 1.37; 95% CL 0.81 to 2.31, not significant). The relative risk for all other causes of death was 1.01 (0.79 to 1.30). When the six cities were considered individually, only Steubenville showed a statistically significant (p < 0.05) elevated risk with respect to the least polluted city (Portage).

Comparison of pollution risks among the various cohort subsets considered is one of the most useful outcomes of a study on individuals. Such comparisons must account for the higher variability among subgroups, however, and the study was not capable of distinguishing excess risks between subgroups less than about 18% (i.e., an excess risk of 1.18 cannot be distinguished

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from one of 1.36, for example). Although none of these subgroup differences were statistically significant, the mortality risks associated with area of residence (and thus air pollution) were higher for females and for smokers, as were risks for those occupationally exposed compared to the nonexposed. Because of reduced uncertainties about exposures of non-smokers and non-occupationally exposed persons to air pollution not reflected in the outdoor monitoring data used in this study, the relative risk estimates for those subgroups might be the most reliable estimates (1.19 and 1.17, respectively).

Issues concerning possible residual confounding, age adjustment, and smoking controls 8 9 were raised, and Dockery and Pope (1994) agreed that confounding is a potential concern but did 10 not address the possibility that variables other than the ones they considered might be important. They dealt with the age adjustment issue quantitatively and pointed out that the air pollution risk 11 12 estimates were reasonably stable over different subgroups by smoking status. Age is a potentially 13 important covariate because it measures both susceptibility to health effects and cumulative 14 exposure to pollutants. There is also a possible interaction involving age, air pollution, and time 15 of death, since air pollution concentrations in some communities such as Steubenville and St. 16 Louis decreased substantially during the years preceding and during the period of the study.

17 The authors of the Harvard Six City Study were cautious in their conclusions, stating only 18 that the results suggest that fine-particulate air pollution "contributes to excess mortality in certain 19 U.S. cities." One further caveat is warranted before placing quantitative reliance on the specific 20 relative risk values generated by the study. If the responses to air pollution truly are chronic in 21 nature, it is logical to expect that cumulative exposure would be the preferred metric. Pollution 22 levels 10 years before the Six City study began were much higher in Steubenville and St. Louis, as 23 indexed by TSP from routine monitoring networks; and atmospheric visibility data suggest that 24 previous fine particle levels may have been higher in winter, but not necessarily in summer. These 25 uncertainties argue for caution in accepting and using the quantitative regression results based solely on coincident monitoring data. For example, annual average TSP in 1965 in Steubenville 26 27 was about three times the value used by Dockery et al.; inclusion of older data in the exposure 28 indices would have reduced implied regression coefficients and relative risk estimates.

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C.2.2. American Cancer Society (ACS) Study

Pope et al. (1995) analyzed 7-year survival data (1982 to 1989) obtained by the American Cancer Society (ACS) for about 550,000 adult volunteers. The Cox proportional hazards model was used to define individual risk factors for age, sex, race, smoking (including passive smoke exposure), occupational exposure, alcohol consumption, education, and body-mass index. The deaths (about 39,000 in all) were assigned to geographic locations using 3-digit zip codes for residences listed at enrollment into the ACS study in 1982. Relative risks were then computed for

- 1 151 metropolitan areas defined by these zip codes and compared to corresponding air quality data 2 (ca. 1980). The sources of air quality data used were (a) the EPA AIRS system data for sulfates, 3 obtained from high-volume sampler filters for 1980, and (b) the Inhalable Particulate Network 4 data for fine particles (PM_{25}) obtained from dichotomous samplers during 1979-81. Pope et al. 5 used the values from this data base reported by Lipfert et al., 1988, but only 50 PM_{2.5} locations 6 could be matched with the death data. The correlation between the two pollutants was 0.73. 7 Causes of death considered included all causes, cardiopulmonary causes (ICD-9 401-440, 460-519), lung cancer (ICD-9 162), and all other causes. 8
- 9 This study took great care with potential confounding factors for which data were 10 available. Several different active smoking measures were considered, as was time exposed to passive smoke. The occupational exposure variable was specific to (any of) chemicals/solvents, 11 12 asbestos, coal or stone dusts, coal tar/pitch/asphalt, diesel exhaust, or formaldehyde. The 13 education variable was an indicator for having less than a high-school education, and alcohol use 14 and body-mass index were considered as linear predictors of survival. Pope et al. (1995) did not 15 report relative risk coefficients they obtained for these cofactors, which does not allow 16 comparison of findings for the non-pollution variables with exogenous estimates from 17 independent studies. Risk factors not considered by Pope et al. (1995) include: income, 18 employment status, dietary factors, drinking water hardness and physical activity levels (all shown 19 to affect longevity); and they did not discuss possible influences of other air pollutants.
- 20 The ACS cohort is not a random sample of the U.S. population; it is 94% white and better 21 educated than the general public, with a lower percentage of smokers than in the Six City Study. 22 The (crude) death rate during the 7.25 years of follow-up was just under 1% per year, which is 23 about 20% lower than expected for the white population of the U.S. in 1985, at the average age 24 reported by Pope et al. In contrast, the corresponding rates for the Six- Cities Study (Dockery 25 et al., 1993) discussed above tended to be higher than the U.S. average. In spite of these 26 differences, the cause specific ratios for smoking are not significantly different between the ACS and Six-Cities studies. 27
- 28 No mention was made of residence histories for the decedents; matching was done on 29 residence location at time of study entry. The 1979 to 1981 pollution values were assumed to be representative of long-term cumulative exposures, in keeping with the goal of analyzing chronic 30 31 effects. However, the previous decade was one of extensive pollution cleanup in most of the 32 nation's dirtiest cities (TSP dropped by a factor of 2 in New York City, for example); but PM 33 levels remained relatively constant in cities that already met the standards. Thus, it is reasonable to expect that the contrast between "clean" and "dirty" cities would have been greater in 1970 34 35 than in 1980. For example, the ranges of TSP and SO₄ across the U.S. in 1970 were from 40 to 36 224 and from 3 to 28 μ g/m³, respectively (Lipfert, 1978). In 1980, these ranges decreased to

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- 1 41-142 and 2-17 μ g/m³ (Lipfert, 1984), suggesting that the dirtiest cities became cleaner while the 2 "clean" cities stayed about the same. The change in pollution range is about a factor of 1.8. If the 3 excess mortality found in the ACS study were in fact due to cumulative exposures, the regression 4 coefficients would have been biased upward (in terms of relative risk per μ g/m³) by only using the 5 more recent data. The typically long latency period for lung cancer (ca. 20 yr.) suggests that data 6 on prior exposures may be particularly important for this cause of death.
- 7 The adjusted total mortality risk ratios (computed for the range of the pollution variables) were 1.15 (95% CL = 1.09 to 1.22) for sulfates and 1.17 (95% CL = 1.09 to 1.26) for $PM_{2.5}$, 8 9 suggesting that particle chemistry may be relatively unimportant as an independent risk factor. 10 Pope et al. (1995) found that the PM pollution coefficients were reduced by 10 to 15% when variables for climate extremes were added to the model. No significant excess mortality for the 11 12 "other" causes of death was attributed to air pollution in this study. Note that Pope et al. found 13 very consistent pollution risks for males and females and for ever-smokers and never-smokers for 14 all-cause mortality. However, the relative risks for air pollution were slightly higher for females 15 for cardiopulmonary causes of death and the sulfate-lung cancer association was only statistically 16 significant for males, except for male never-smokers.
- The results of the ACS prospective study were qualitatively consistent with those of the 17 18 Six City Study with regard to their findings for sulfates and fine particles; but relative standard 19 errors were smaller, as expected because of the substantially larger ACS database. However, no 20 other copollutants (e.g., O₃, CO, NO₂, etc.) were investigated in the ACS analysis, so that it was 21 not possible to provide an analogous type of pollutant comparison given earlier in Table C-6 for 22 the Six Cities Study. In addition, the ACS regression coefficients were about 1/4 to 1/2 of the 23 corresponding Six City values and were much closer to the corresponding values obtained in 24 various acute mortality studies.
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C.2.3. California Seventh-Day Adventists Study

27 In the Abbey et al. (1991) prospective study (the Adventist Health Study of Smog or 28 "AHSMOG"), 6,338 long-term California residents (all white, non-Hispanic, and nonsmoking) 29 were followed for 6 to 10 years, beginning in 1976. Ambient air quality data dating back to 1966 30 were used in analyses restricted to those who lived within 5 miles of their current residence for at 31 least 10 years. Subjects lived either within the 3 major California air basins (San Diego, Los 32 Angeles, or San Francisco) or else were part of a random 10% sample of Adventist Health Study 33 participants in the rest of California. Individual exposure profiles (duration of exposure to 34 specific minimum concentration levels) were created for each participant, by interpolating to their 35 zip code centroids based on the 3 nearest monitoring stations. Monitored pollutants were mainly 36 limited to TSP and O₃ in this paper; but, total oxidant concentrations were used in the early part

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1 of the monitoring record. Health endpoints evaluated and the numbers of cases included: 2 (a) newly diagnosed cancers (incidence at any site) for males, 115; (b) any cancer site for females, 3 175; (c) respiratory cancer, 17; (d) definite myocardial infarction, 62; (e) mortality from any 4 external cause, 845; and (f) respiratory symptoms, 272. The Cox proportional hazards model was 5 used, considering age, sex, past smoking, education, and presence of definite symptoms of 6 asthma, chronic bronchitis, or emphysema of airway obstructive disease (AOD) in 1977 as 7 individual risk factors, together with various exposure indices for TSP or O₃ (considered 8 separately). Data on occupational exposures and history of high blood pressure were available 9 but not used in the mortality model; nor were data available on climate, body mass, income, 10 migration, physical activity levels or diet.

11 Of the above endpoints, only respiratory symptoms and female cancers (any site) were 12 reported by Abbey et al. (1991) to be statistically associated with TSP exposure. Neither heart 13 attacks or nonexternal mortality were associated with either TSP or O_3 / oxidants. The authors 14 stated that possible errors in their estimated exposures to air pollution may have contributed to 15 the lack of significant findings, and a later version of the data base included estimates of 16 attenuation resulting from time spent indoors (Abbey et al., 1993), but mortality was not 17 considered in the 1993 paper. Follow-up analyses (Abbey et al., 1995) considered exposures to 18 PM₁₀ (estimated from site-specific regressions on TSP), PM_{2.5} (estimated from visibility), sulfates (SO₄), and visibility per se (extinction coefficient). No significant associations with nonexternal 19 20 mortality were reported, and only high levels of TSP or PM₁₀ were associated with AOD or 21 bronchitis symptoms.

22 This study used an unique air quality data base developed explicitly for studying effects of 23 long-term cumulative exposures to community air pollution. The technique provided spatial 24 interpolations that were somewhat better for O₃ than for TSP, in keeping with the regional nature 25 of O₃. TSP may have been an inadequate index of exposure to inhalable particles, especially in 26 this relatively arid region where a large fraction of non-inhalable crustal particles could be 27 expected. Also, no attention was given to temporal matching of air quality and health; the 28 analyses using this data base were intended to evaluate the hypothesis that health is affected by 29 cumulative long-term pollution exposure at some undetermined time, as opposed to acute or 30 coincident exposures. Note that the data base began in 1966 and the mortality follow-up began 31 10 years later. Because air quality generally improved during this period, highest pollutant 32 concentrations likely occurred in the earlier part of the record; and one would not expect 33 spatially-based correlations to also reflect the sum of acute effects, as when air quality and health 34 data are also matched in time.

The PM CD (U.S. EPA 1996a) noted that the finding of Abbey et al. (1991, 1995) of no association between long-term cumulative exposure to ambient TSP or O_3 (or to SO_4 or estimated

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PM₁₀ or PM_{2.5}) concentrations and all natural-cause mortality could be interpreted as showing the
 absence of chronic responses after 10 years but not necessarily the absence of (integrated) acute
 responses, since coincident air pollution exposures or integrated exposures over the preceding few
 years were not considered. It is also possible that the exposure measurements or estimates used
 were inadequate or that the latency period for chronic effects may exceed 10 years and that
 additional follow-up might still reveal chronic effects.

Further such follow-up analyses of the same California AHSMOG database have been
reported recently by Abbey et al. (1999). These analyses (not considered in the 1996 PM CD or
1997 PM NAAQS decisions) do provide some evidence indicative of increased risk of mortality
from contributing non-malignant respiratory causes being associated with long-term PM
exposures. Other recent AHSMOG analyses reported by Abbey et al. (1999) and Beeson et al.
(1998) are also suggestive of increased risk of mortality from lung cancer possibly being

13 associated with long-term PM_{10} exposures, as summarized below.

14 Abbey et al. (1999) evaluated the mortality status of AHSMOG subjects after ca. 15-years 15 of follow-up (1977-1992), finding 1,628 deaths (989 female, 639 male) in the cohort. There were 16 1,575 deaths from all natural (non-external) causes, of which 1,029 were cardiopulmonary deaths, 17 135 were non-malignant respiratory deaths (ICD9 codes 460-529), and 30 were lung cancer 18 deaths (ICD9 code 162). Abbey et al. (1999) also created an additional death category, 19 "contributing respiratory causes" (CRC). CRC included any mention of nonmalignant respiratory 20 death as either an underlying cause or a contributing cause on the death certificate CRC coded by 21 an exposure-blinded nosologist (the other groups listed only underlying causes), with 410 deaths 22 (246 female and 164 male) being found. Numerous analyses were done for the CRC category, 23 due to the large numbers and relative specificity of respiratory causes as a factor in the deaths. 24 Education was used as an index of socio-economic status, rather than income. Physical activity 25 and occupational exposure to dust were also used as covariates. Migration was not a major 26 concern in this residentially stable cohort.

27 A number of exposure indicators were used: mean values of PM₁₀ (imputed from TSP in 28 the earlier years of the study), SO₄, SO₂, O₃, and NO₂; and "threshold" indicators (i.e., days per year with $PM_{10} > 100 \ \mu g/m^3$; and hours per year with $O_3 > 100 \text{ ppb}$). In summary tables that 29 follow below, the "standard" increments used for PM_{10} and SO_4 are (a) the same as used earlier 30 for the short-term mortality studies (50 μ g/m³ for PM₁₀ and 15 μ g/m³ for SO₄) and (b) 30 days 31 per year for exceedances of PM₁₀ above 100 μ g/m³. The mean values for PM₁₀ and SO₄ during 32 33 the study period were 51 and 7.2 μ g/m³ respectively, and 31 days per year for PM₁₀ exceedances 34 over 100 μ g/m³. The means were much larger than the inter-quartile ranges (IQR) of 24 and 35 3.0 μ g/m³. IQR is the increment used for other variables. RR and confidence limits using IQR

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from Abbey et al. (1999) are shown to 2 decimal places; those estimated for standard increments
 are shown to 3 decimal places.

- Cox proportional hazard models adjusted for a variety of covariates, or stratified by sex,
 were used in the models. The "time" variable used in most of the models was survival time from
 date of enrollment, except that age on study was used for lung cancer effects due to the expected
 lack of short-term effects. A large number of covariate adjustments were evaluated, as shown in
 Table C-7 and described by Abbey et al. (1999).
- 8 The CRC RR estimates for 30 days per year with $PM_{10} > 100 \ \mu g/m^3$ for males and females 9 combined are shown in Table C-7. Positive and statistically significant effects are found for 10 almost all models that include age, pack-years of smoking, and body-mass index (BMI) as covariates. Subsets of the cohort also often had elevated risks. Former smokers had higher 11 12 relative risks than never-smokers (RR for PM₁₀ exceedances for never-smokers was marginally 13 significant by itself, in spite of the reduced sample size). Subjects with low intake of anti-oxidant 14 vitamins A, C, E had significantly elevated risk of response to PM₁₀ whereas those with adequate intake did not, suggesting that dietary factors (or possibly other socio-economic or life style 15 16 factors for which they are a surrogate) may be important covariates. There also appears to be a 17 gradient of PM₁₀ risk with respect to time spent outdoors, with individuals who had spent at least 18 16 hours per week outside at distinctly elevated risk from PM_{10} exceedances. The extent to which 19 time spent outdoors is a surrogate for other variables or is a modifying factor reflecting temporal 20 variation in exposure to ambient air pollution is not certain. For example, males spend about twice 21 as much time outdoors as females, so that outdoor exposure time is confounded with gender.

22 A considerably different picture is shown when the analyses are broken down by gender. 23 Table C-8 shows much lower RR for female CRC deaths for all co-pollutants, with all female 24 RR positive, but not statistically significant. The CRC for males remains significant only for PM_{10} 25 exceedances, but not for other air pollution metrics. The PM₁₀ exceedance effect for CRC for 26 both sexes is roughly the average of that for males and females. Personal monitoring was not 27 conducted on this part of the cohort, and other factors (e.g., occupational exposure) for which the 28 questionnaire was not adequate may also account for male vs. female differences, along with 29 gender differences in the amount of time spent outdoors. Finally, it is not surprising that individuals reporting respiratory symptoms in 1977 may be at greater risk to PM₁₀ or other 30 31 environmental insults presumably involved in subsequent CRC deaths, and prior health status may 32 also be gender-related.

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Table C-9 shows much lower RR for female non-external deaths for all co-pollutants, with
 no female RR positive nor statistically significant. Deaths from non-external causes for males

- 1 remains statistically significant for PM_{10} exceedances, but not for other air pollution metrics.
- 2 However, the RR estimates for males for other air pollutant metrics are relatively large.
- Table C-10 shows much lower RR for female cardio-pulmonary deaths for all
 co-pollutants, with only the female RR for mean SO₂ positive and none statistically significant.
 The RR for deaths from cardiopulmonary causes for males is no longer statistically significant for
 PM₁₀ exceedances, nor for other air pollution metrics (although the RR estimates for males for air
 pollutant metrics are relatively large).
- Table C-11 shows a confusing welter of results obtained for lung cancer mortality.
 For example, the RR's for lung cancer deaths are significant for males for PM₁₀ and O₃ metrics,
 but not for females. In contrast lung cancer deaths are significant for mean NO₂ for females, but
 not for males, but lung cancer metrics for mean SO₂ are significant for both males and females.
 This pattern is not readily interpretable, but may be attributable to the very small numbers of
- 13 cancer-related deaths (18 for females; 12 for males), resulting in wide RR confidence intervals.
- 14 In general, this study (Abbey et al., 1999) suggests a pattern of mortality from diverse 15 causes (e.g., CRC, lung cancer) in males, but provides little evidence for female mortality from 16 these causes. The male causes primarily appear to be associated with exposures to PM_{10} and 17 especially to $PM_{10} > 100 \ \mu g/m^3$. Some other air pollutants (SO₂, NO₂) appear to be associated 18 with lung cancer deaths in females.
- 19 The analyses reported here attempted to separate PM_{10} effects from those of the other 20 pollutants by use of two-pollutant models, but none of the quantitative findings from these models 21 were reported. The Abbey et al. (1999) text mentions that the PM₁₀ coefficient for CRC remained 22 stable or increased when other pollutants were added to the model. Lung cancer mortality models 23 for males were evaluated for co-pollutant effects in detail. NO₂ remained nonsignificant in all 24 two-pollutant models, and the other pollutant coefficients were stable in magnitude. The PM_{10} 25 and O_3 effects remained stable when SO_2 was added, suggesting that their effects are independent. 26 However, the effects of PM₁₀ and O₃ were hard to separate because
- these pollutants were highly correlated in this study. When both exceedances $PM_{10} > 100 \ \mu g/m^3$
- and $O_3 > 100$ ppb were used in the model, both RR were reduced in magnitude, but the O_3
- exceedance RR remained more significant than the RR for the PM_{10} exceedance. The possibility
- 30 that the finding of a significant PM_{10} effect is partially attributable to correlation with other
- 31 pollutants such as O_3 cannot be precluded. The SO₂ coefficient for lung cancer mortality in
- 32 females remained stable in two-pollutant models when PM_{10} and O_3 exceedances were included.
- This suggests that the significance of the SO_2 effect for females may not be an artifact wholely
- 34 attributable to collinearity with these co-pollutants.
- 35

36 Beeson et al. (1998)

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1 This study used essentially the same data as did Abbey et al. (1999), but concentrates on 2 lung cancer incidence (1977-1992) as an endpoint. There were only 20 female cases and 16 male 3 cases of lung cancer among the 6,338 AHSMOG subjects. The exposure metrics were 4 constructed to be specifically relevant to cancer, being the annual average of the monthly 5 exposure indices from January, 1973 through the following months, but ending 3 years before the 6 date of diagnosis of the case. This represents a 3-year lag between exposure and diagnosis of 7 lung cancer, allowing for a latency period. Therefore, statistical indices for exposure have somewhat different statistics than in Abbey et al. (1999), such as the IQR and mean. 8

9 The covariates in the Cox proportional hazards model were pack-years of smoking and education, and the time variable was attained age. A number of additional covariates were 10 11 evaluated for inclusion in the model, but only 'current use of alcohol' met the criteria for inclusion in the final model. Individual pollutants evaluated were PM₁₀, SO₂, NO₂, and O₃. No interaction 12 13 terms with the pollutants proved to be significant, including outdoor exposure times. Gender-14 specific relative risk estimates were reported for the various risk factors. Results are shown in Table C-12 for males and Table C-13 for females. Standard increments were used for PM₁₀ mean 15 (50 μ g/m³) and exceedances of PM₁₀ > 100 μ g/m³ (30 d/y). The RR estimates and confidence 16 limits using IQR from Beeson et al. (1998) are shown to 2 decimal places, those estimated for 17 18 standard increments are shown to 3 decimal places.

19 The RR estimates for the male lung cancer cases are: positive and statistically significant 20 for all PM₁₀ indicators; positive and predominantly significant for O₃ indicators, except for mean 21 O_3 , number of O_3 exceedances > 60 ppb, and in former smokers; and are positive and significant 22 for mean SO₂, except when restricted to proximate monitors. The RR for mean NO₂ is positive 23 but not significant. The very high RR for mean PM_{10} for males (31.1) may be attributable to the 24 small number of cases (N = 16) and the large standard increment (50 μ g/m³) used. When data are 25 restricted to subjects with at least 80 percent A/B quality data (within 32 km of the residence), the RR is reduced to 9.26 over 50 μ g/m³. The RR over the IQR of 24 μ g/m³ in the full data set is 26 27 5.21, so that the use of the IQR may be more appropriate for the exposure in long-term studies.

The female lung cancer RR estimates reported by Beeson et al. (Table C-13) are much smaller than those for males, not being statistically significant for any indicator of PM_{10} or O_3 and statistically significant only for mean SO_2 .

Extensive multi-pollutant analyses were also carried out. Regression coefficients for PM_{10} and SO_2 were not reduced when O_3 or NO_2 were added to the single-pollutant models for males. The regression coefficients for the two-pollutant model with PM_{10} and SO_2 remained highly positive and significant, which the authors suggest may be associated with independent effects of PM_{10} and SO_2 on lung cancer incidence. PM_{10} was more strongly correlated with lung cancer in males than the other pollutants. For females, the SO_2 coefficient remained significant when

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1 co-pollutants were added one at a time, and was the air pollutant most strongly associated with

2 female lung cancer cases.

3 The results of Abbey et al. (1999) and Beeson et al. (1998) are somewhat different than those of earlier studies using the same cohort. Abbey et al. (1991) reported completely 4 5 non-significant relationships between total ('all natural causes') mortality and air pollution. The RR for 1000 h/y of TSP > 200 μ g/m³ was 0.99 (CI 0.87-1.13), and for 500 h/y of O₃ > 100 ppb 6 was 1.00 (CI 0.89-1.12), after 10 years of follow-up. Also, Abbey et al. (1991) reported no 7 8 statistically significant increases in all malignant neoplasms for males attributable to air pollution. The RR for 1000 h/y of TSP > 200 μ g/m³ was 0.96 (CI 0.68-1.36), and for 500 h/y of O3 > 100 9 ppb was 1.09 (CI 0.80-1.47), after 10 years of follow-up. However, there was a statistically 10 11 significant increase in all malignant neoplasms for females. The RR for females attributed to 1000 12 h/y of TSP > 200 μ g/m³ was 1.37 (CI 1.05-1.80). Neoplasms in females attributed to 500 h/y O₃ 13 > 100 ppb were much less significant, with RR = 1.03 (CI 0.81-1.32). 14 15 C.2.4. Relationship of AHSMOG to Six Cities and ACS Study Findings

16 The results of the recent AHSMOG mortality studies (Abbey et al., 1999) are compared below with the earlier Six Cities Study (Dockery et al., 1993) and ACS Study (Pope et al., 1995). 17 18 Tables C-14, C-15, and C-16 compare the estimated RR for total, cardiopulmonary, and lung 19 cancer mortality, respectively, among the studies. The PM indices used are the mean PM_{10} concentration for the Six Cities and AHSMOG studies (increment 50 μ g/m³), and the mean PM₂₅ 20 and SO₄ concentrations (increments 25 and 15 μ g/m³ respectively) for the ACS study. The 21 22 comparisons for the Six Cities and ACS studies have been translated from published RR for the 23 most polluted vs. least polluted city for PM₁₀, PM₂₅, and SO₄. Results are shown by sex and 24 smoking status. The AHSMOG subjects are classified as 'non-smokers', although some former 25 smokers are included. The ACS study combines past and current smokers into an 'ever smoker' 26 category, although long-term past smokers are at much lower risk than current smokers. The 27 number of subjects in these studies varies greatly (6,338 AHSMOG subjects, 8,111 Six Cities 28 Study subjects; compared to 295,223 subjects in the 50 fine particle cities and 552,138 subjectsin the 151 sulfate cities of the ACS study), and may partially account for differences among their 29 30 results.

Table C-14 shows relative risks for total mortality at comparable standard increments. RR is generally highest for the Six Cities Study. The AHSMOG Study found a much smaller RR for women than did the other studies, whereas the effect for males was similar to non-smokers in the ACS Study and marginally significant. RR among the three studies varied substantially with sex and smoking categories. Six of the 16 independent analyses showed significant positive RR (LCL ≥ 1.0), but subsetting the data allowed less power to detect effects than the whole data sets would

- 1 have allowed. Neither of the AHSMOG RR were significant using the mean as the PM_{10} index,
- 2 but another PM_{10} index (exceedances over 100 μ g/m³) was significant for males.

3 Table C-15 shows relative risks for cardiopulmonary mortality at comparable standard 4 increments. RR is highest for the Six Cities Study, which did not report separate effects by sex 5 and smoking status. The AHSMOG Study found a much smaller cardiopulmonary RR for women 6 than did the other studies. However, the RR for male non-smokers was much more similar to the 7 ACS results than for female non-smokers. RR for the AHSMOG endpoint CRC ('contributing 8 respiratory causes') was more similar to the ACS findings for women, but higher in men, although 9 the confidence intervals are very wide. Seven of 13 of the independent analyses showed 10 significant positive RR (LCL > 1.0). The AHSMOG cardiopulmonary RRs using mean PM_{10} were not significant for either males or females. However, the 100 μ g/m³ exceedance index for 11 12 males was nearly so.

13 Table C-16 shows relative risks for lung cancer mortality at comparable standard 14 increments for PM-related variables. The lung cancer mortality RR estimates were highest for 15 males in the AHSMOG study, and statistically significant. The AHSMOG study also found a larger RR for women than did the other studies. The only other statistically significant finding for 16 17 lung cancer mortality was for past and current male smokers in the ACS 151-city sulfate study. 18 The overall pattern of results for lung cancer, then, is a somewhat conflicting set of findings 19 across the three prospective cohort studies assessed here, providing only somewhat suggestive 20 evidence at best for possible ambient PM relationship to increased lung cancer risk.

21 There is no obvious statistically significant relationship between PM effect sizes, gender, 22 and smoking status across these studies. The AHSMOG studies show no statistically significant 23 relationships between PM₁₀ and total mortality or cardiovascular mortality for either sex, and only 24 for male lung cancer incidence and lung cancer deaths in a predominantly non-smoking sample. 25 The ACS results, in contrast, show similar and significant associations with total mortality for both "never smokers" and "ever smokers", although the ACS cohort may include a substantial 26 27 number of long-term former smokers with much lower risk than current smokers. The Six Cities 28 Study cohort shows the strongest evidence of a higher PM effect in current smokers than in non-29 smokers, with female former smokers having a higher risk than male former smokers. This study 30 suggests that smoking status is "effect modifier" for ambient PM, just as smoking may be a health 31 effect modifier for ambient ozone (Cassino et al., 1999).

It is interesting to note, in relation to the above discussion, that a comparison of the Six-Cities Study non-smoker RRs with the Six-Cities results in Table C-14 for smokers indicates that larger and more significant effects of ambient PM pollution are found for smokers than non-smokers. This suggests that smoking is an effect modifier that increases the adverse effects of ambient pollution. This trend is consistent with air pollution effect causality, as smokers represent a compromised population, logically more likely to be adversely affected by air
pollution. This may also explain why the reported AHSMOG study RRs are generally not
significant, in contrast with the overall Six-Cities Study results (but consistent with the Six-Cities
nonsmoker results), as there are no identified smokers among the AHSMOG study group to
"drive up" the overall significance of the air pollution effect. This again indicates that more years
of follow-up may be required to see any statistically significant total mortality effects in both the
AHSMOG and Six-Cities studies' non-smoking populations.

8

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C.2.5. Studies by Particulate Matter Size-Fraction and Composition

10 Particulate matter mass varies widely over time and from place to place in size and chemical composition, and this likely affects the toxicity of that mass. The semi-individual cohort 11 12 studies assessed here investigated the relative roles of various PM components in the air pollution 13 association with mortality. As shown in Table C-17, the Harvard Six-Cities study (Dockery et al., 14 1993) results indicated that the PM_{2.5} and SO₄ RR associations (as indicated by their respective 15 95% CI's and t-statistics) were stronger than those for the coarser mass components. However, 16 the effects of sulfate and non-sulfate PM_{2.5} are indicated to be quite similar. Acid aerosol (H⁺) 17 exposure was also considered by Dockery et al. (1993), but only less than one year of 18 measurements collected near the end of the follow-up period were available in most cities, so the 19 Six-Cities results were much less conclusive for the acidic component of PM than for these other 20 PM metrics (that, in contrast, were measured over many years during the study). The Six-Cities 21 Study also yielded total mortality RR estimates for the reported range across those cities of PM₂₅ 22 and SO₄ concentrations that, although not statistically different, were roughly double analogous 23 RRs for the TSP-PM₁₅ and $PM_{15-2.5}$ mass components.

Table C-18 presents comparative $PM_{2.5}$ and SO_4 results from the ACS study that indicate that, although the RR differences were not statistically significant across pollutants, the SO_4 RRs were in every case more strongly significant than those for the $PM_{2.5}$ across the various mortality cause classifications considered, especially for lung cancer (SO_4 t=2.92 vs. t=0.38 for $PM_{2.5}$).

28 The most recent AHSMOG study analysis (Abbey et al., 1999) employed PM₁₀ as its PM 29 mass index, finding some significant associations with total and by-cause mortality, even after 30 controlling for potentially confounding factors (including other pollutants). This analysis also 31 considered SO₄ as a PM index for all health outcomes studied except lung cancer, but SO₄ was 32 not as strongly associated as PM₁₀ with mortality, and was not found to be statistically significant 33 for any mortality category. The significant mortality associations found for PM₁₀ contrasts with 34 previously published AHSMOG study PM analyses that found weaker mortality associations with 35 TSP (Abbey et al., 1991). Although the longer follow-up time in this new analysis may have also 36 contributed, the greater strength of association by PM_{10} vs. TSP is consistent with the Harvard

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- Six-City study results presented in Table C-17, as well as with the Özkaynak and Thurston (1987)
 cross-sectional comparisons of mortality associations with the various PM fractions.
- Single-pollutant results about PM components are informative, however, as shown in Table C-19 for total mortality, and in Table C-20 for cardiopulmonary causes. The t-statistics are compared for studies where appropriate: mean PM_{10} , $PM_{10-2.5}$, $PM_{2.5}$, and sulfate for the Six Cities (Dockery et al., 1993); mean $PM_{2.5}$ and sulfate for ACS (Pope et al., 1995); mean PM_{10} and sulfate, and PM_{10} exceedances of 100 μ g/m³ for AHSMOG (Abbey et al., 1999).
- 8 Estimates for Six Cities parameters were calculated in two ways: (1) mortality RR for 9 most versus least polluted city in (Table 3, Dockery et al., 1993) adjusted to standard increments; (2) ecological regression fits in (Table 12-18, U.S. Environmental Protection Agency, 1996). The 10 11 eastern and mid-western Six Cities suggest a strong and highly significant relationship for fine 12 particles and sulfates, a slightly weaker but still highly significant relationship to PM₁₀, and a marginal relationship to PM_{10-2.5}. The ACS study looked at a broader spatial representation of 13 14 cities, and found a stronger statistically significant relationship to PM_{2.5} than to sulfate (no other 15 pollutants were examined).
- Overall, the prospective cohort studies conducted to-date collectively confirm crosssectional study indications that, as opposed to the more coarse mass fractions, the fine mass
 component of PM (and sometimes including its acidic sulfate constituent) are strongly correlated
 with mortality.
- The credibility of the above findings of increased risk of mortality being associated with chronic, long-term exposures to fine particles is enhanced by analogous findings of increased risk of respiratory symptoms and lung function decrements being associated with long-term exposures to fine particles, as illustrated in Figure C-4. That figure graphically depicts results from the study reported on by Razienne et al. (1996), which demonstrate strong positive relationships between decrements in children's lung function and long-term exposure to fine particles (indexed by $PM_{2.1}$), but not to inhalable thoracic coarse particles ($PM_{10-2.1}$).
- 28 C.2.6. Conclusions

27

A review of the prospective cohort studies summarized in the previous PM AQCD (U.S. Environmental Protection Agency, 1996) indicates that past epidemiologic studies of chronic PM exposures collectively indicate increases in mortality to be associated with long-term exposure to airborne particles of ambient origins. The PM effect size estimates for total mortality from these studies also indicate that a substantial portion of these deaths reflected cumulative PM impacts above and beyond those exerted by acute exposure events.

1 The new AHSMOG study (Abbey et al., 1999) provides all-cause mortality RR estimates 2 for adult males that are quantitatively and qualitatively consistent with prior semi-individual 3 prospective cohort studies, especially the similarly designed 6-Cities study. Extensive new 4 by-gender, by-cause, and multiple pollutant sensitivity analyses, as well as a more comprehensive analyses of numerous potentially uncontrolled factors in this study (such as of the effects of 5 6 variations in the time spent outdoors) provide important new evidence that is largely supportive of 7 the mortality associations with PM of ambient origins previously reported by the Six-Cities and 8 ACS studies.

With regard to the role of various PM constituents in the PM-mortality association, cross sectional studies have generally found that the fine particle component, as indicated either by
 PM_{2.5} or sulfates, was the PM constituent most consistently associated with mortality.
 In addition, the Six-Cities prospective semi-individual study also indicates that the fine mass
 components of PM are more strongly associated with the mortality effects of PM than the coarse
 PM components.

15 The recent analyses of the long-term AHSMOG study provide some evidence indicative of 16 health effects being associated with ambient PM₁₀ exposure for which a substantially greater level 17 of individualized ambient PM₁₀ information is available, but also demonstrates some differences 18 with the earlier Six Cities and ACS studies (Dockery et al., 1993; Pope et al., 1995). Statistically 19 significant increases in lung cancer incidence (Beeson et al., 1998) and statistically significant 20 increases in lung cancer deaths and deaths associated with any contributing respiratory causes 21 (Abbey et al., 1999) were found in AHSMOG males, but not females. The results were generally 22 robust to different confounder specifications, population subsets, and inclusion of co-pollutants, 23 and were larger for and more significant for PM exceedance indices (number of days per year with PM_{10} greater than a cut point, typically 100 μ g/m³) than with the mean PM_{10} concentration. 24 25 However, PM₁₀ was estimated from TSP rather than measured in the earlier part of the AHSMOG 26 study and, therefore, the AHSMOG results may not be as credible as those from the other two prospective cohort studies where direct PM₁₀, PM₂₅, or SO₄ measurements data were used. 27 Using the same mean PM₁₀ increment of 50 μ g/m³, total mortality attributable to long-term 28 29 ambient PM₁₀ RR was similar to that of the ACS study for PM_{2.5} for male nonsmokers (1.24) and 30 smaller than that for the Six Cities study (1.57), albeit only significant for the ACS study 31 (Table C-13). The AHSMOG RR for females (Table 6-31) is smaller and non-significant (0.88), 32 whereas the ACS RR for female non-smokers is significant and only somewhat smaller than the 33 male RR (1.22 in the 50-city PM_{2.5} study, 1.15 in the 151-city SO₄ study) and 1.28 in the

34 Six Cities.

The AHSMOG findings for cardiopulmonary mortality attributable to long-term ambient
 PM₁₀ are positive for males, but not statistically significant, whereas the ACS findings are

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1 significant for female nonsmokers in both studies and in male nonsmokers for the 151-city study 2 (Table C-14). However, the male RR in AHSMOG (1.22 for cardiopulmonary deaths, 1.54 for 3 CRC deaths) is similar to that of ACS male non-smokers (1.24 for the 50-city study, 1.21 for the 4 151-city study) and smaller than that for all Six Cities subjects (1.74, includes smokers and 5 non-smokers). The ACS female non-smokers have RR of 1.58 and 1.32 respectively, both 6 significant, compared to 0.84 in AHSMOG. 7 Lung cancer mortality attributable to long-term ambient PM_{10} is not significant for females 8 in any of the studies, nor for male nonsmokers in ACS, but was reported to be statistically 9 significant for male nonmokers in AHSMOG and male smokers in ACS 151-city. Lung cancer 10 mortality attributable to long-term ambient PM_{2.5} was not significant for either gender in the ACS and Six Cities studies. Thus, the available overall evidence, from the three prospective cohort 11 12 studies of PM effects assessed here, definitely is not conclusive and can, at best, be viewed as 13 indicative of possible ambient PM associations with increased risk of lung cancer or associated 14 mortality.

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	RR (± CI) Only PM	RR (± CI) Other Pollutants	Reported PM ₁₀ Levels Mean
Study Location	in Model	in Model	(Min/Max) [†]
Increased Total Acute Morta	ality		
Six Cities ^a			
Portage, WI	1.04 (0.98, 1.09)	—	18 (±11.7)
Boston, MA	1.06 (1.04, 1.09)		24 (±12.8)
Topeka, KS	0.98 (0.90, 1.05)	—	27 (±16.1)
St. Louis, MO	1.03 (1.00, 1.05)	—	31 (±16.2)
Kingston/Knoxville, TN	1.05 (1.00, 1.09)	—	32 (±14.5)
Steubenville, OH	1.05 (1.00, 1.08)	—	46 (±32.3)
St. Louis, MO ^c	1.08 (1.01, 1.12)	1.06 (0.98, 1.15)	28 (1/97)
Kingston, TN ^c	1.09 (0.94, 1.25)	1.09 (0.94, 1.26	30 (4/67)
Chicago, IL ^h	1.04 (1.00, 1.08)		37 (4/365)
Chicago, IL ^g	1.03 (1.02, 1.04)	1.02 (1.01, 1.04)	38 (NR/128)
Utah Valley, UT ^b	1.08 (1.05, 1.11)	1.19 (0.96, 1.47)	47 (11/297)
Birmingham, AL ^d	1.05 (1.01, 1.10)	—	48 (21, 80)
Los Angeles, CA ^f	1.03 (1.00, 1.055)	1.02 (0.99, 1.036)	58(15/177)
Increased Hospital Admission	ons (for Elderly > 65 yrs.)		
Respiratory Disease			
Toronto, CAN ⁱ	1.23 (1.02, 1.43) [‡]	1.12 (0.88, 1.36) [‡]	30-39*
Tacoma, WA ^j	1.10 (1.03, 1.17)	1.11 (1.02, 1.20)	37 (14, 67)
New Haven, CT ^j	1.06 (1.00, 1.13)	1.07 (1.01, 1.14)	41 (19, 67)
Cleveland, OH ^k	1.06 (1.00, 1.11)	_	43 (19, 72)
Spokane, WA ¹	1.08 (1.04, 1.14)	_	46 (16, 83)
COPD			
Minneapolis, MN ⁿ	1.25 (1.10, 1.44)	_	36 (18, 58)
Birmingham, AL ^m	1.13 (1.04, 1.22)	—	45 (19, 77)
Spokane, WA ¹	1.17 (1.08, 1.27)	_	46 (16, 83)
Detroit, MI ^o	1.10 (1.02, 1.17)		48 (22, 82)

Table C-1. Effect estimates per 50 μ g/m ³ increase in 24-h PM ₁₀ concentration	ons from
U.S. and Canadian studies	

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Study Location	RR (± CI) Only PM in Model	RR (± CI) Other Pollutants in Model	Reported PM ₁₀ Levels Mean (Min/Max) [†]
Pneumonia			
Minneapolis, MN ⁿ	1.08 (1.01, 1.15)	_	36 (18,58)
Birmingham, AL ^m	1.09 (1.03, 1.15)	_	45 (19, 77)
Spokane, WA ¹	1.06 (0.98, 1.13)	_	46 (16, 83)
Detroit, MI°		1.06 (1.02, 1.10)	48 (22, 82)
Ischemic HD			
Detroit, MI ^p	1.02 (1.01, 1.03)	1.02 (1.00, 1.03)	48 (22, 82)
Increased Respiratory Sy	rmptoms		
Lower Respiratory			
Six Cities ^q	2.03 (1.36, 3.04)	Similar RR	30 (13,53)
Utah Valley, UT ^r	1.28 (1.06, 1.56) ^t		46 (11/195)
	$1.01 (0.81, 1.27)^{\pi}$		
Utah Valley, UT ^s	1.27 (1.08, 1.49)	—	76 (7/251)
Cough			
Denver, CO ^x	1.09 (0.57, 2.10)	—	22 (0.5/73)
Six Cities ^q	1.51 (1.12, 2.05)	Similar RR	30 (13, 53)
Utah Valley, UT ^s	1.29 (1.12, 1.48)	—	76 (7/251)
Decrease in Lung Function	<u>on</u>		
Utah Valley, UT ^r	55 (24, 86)**	—	46 (11/195)
Utah Valley, UT ^s	30 (10, 50)**	—	76 (7/251)
Utah Valley, UT ^w	29 (7,51)***		55 (1,181)

Table C-1. Effect estimates per 50 μ g/m³ increase in 24-h PM₁₀ concentrations from U.S. and Canadian studies (continued)

References:

^a Schwartz et al. (1996a).	¹ Schwartz (1996).	^x Ostro et al. (1991)
^b Pope et al. (1992, 1994)/O ₃ .	^m Schwartz (1994e).	[†] Min/Max 24-h PM ₁₀ in parentheses unless noted
^c Dockery et al. (1992)/O ₃ .	ⁿ Schwartz (1994f).	otherwise as standard deviation (± S.D), 10 and
^d Schwartz (1993).	°Schwartz (1994d).	90 percentile (10, 90). $NR = not$ reported.
^f Kinney et al. (1995)/O ₃ , CO.	^p Schwartz and Morris (1995)/O ₃ , CO, SO ₂ .	^T Children.
^g Ito and Thurston (1996)/O ₃ .	^q Schwartz et al. (1994).	^π Asthmatic children and adults.
^h Styer et al. (1995).	^r Pope et al. (1991).	*Means of several cities.
ⁱ Thurston et al. (1994)/O ₃ .	^s Pope and Dockery (1992).	**PEFR decrease in ml/sec.
^j Schwartz (1995)/SO ₂ .	^t Schwartz (1994g)	****FEV ₁ decrease.
^k Schwartz et al. (1996b).	^w Pope and Kanner (1993).	[‡] RR refers to total population, not just>65 years.

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Acute Mortality	Indicator	RR (± CI) per 25 µg/m ³ PM Increase	Reported PM Levels Mean (Min/Max) [†]
Six City ^a			
Portage, WI	PM _{2.5}	1.030 (0.993, 1.071)	11.2 (±7.8)
Topeka, KS	PM _{2.5}	1.020 (0.951, 1.092)	12.2 (±7.4)
Boston, MA	PM _{2.5}	1.056 (1.038, 1.0711)	15.7 (±9.2)
St. Louis, MO	PM _{2.5}	1.028 (1.010, 1.043)	18.7 (±10.5)
Kingston/Knoxville, TN	PM _{2.5}	1.035 (1.005, 1.066)	20.8 (±9.6)
Steubenville, OH	PM _{2.5}	1.025 (0.998, 1.053)	29.6 (±21.9)
Increased Hospitalization	n		
Ontario, CAN ^b	$\mathrm{SO}_4^=$	1.03 (1.02, 1.04)	R = 3.1 - 8.2
Ontario, CAN ^c	$\mathbf{SO}_4^=$ \mathbf{O}_3	1.03 (1.02, 1.04) 1.03 (1.02, 1.05)	R = 2.0-7.7
NYC/Buffalo, NY ^d	$\mathrm{SO}_4^=$	1.05 (1.01, 1.10)	NR
Toronto ^d	$\mathrm{H^{+}}\ (\mathrm{Nmol/m^{3}})$ $\mathrm{SO_{4}^{=}}$ $\mathrm{PM_{2.5}}$	1.16 (1.03, 1.30) [*] 1.12 (1.00, 1.24) 1.15 (1.02, 1.78)	28.8 (NR/391) 7.6 (NR, 48.7) 18.6 (NR, 66.0)
Increased Respiratory S	ymptoms		
Southern California ^e	$\mathrm{SO}_4^=$	1.48 (1.14, 1.91)	R = 2-37
Six Cities ^f (Cough)	$\begin{array}{c} PM_{2.5}\\ PM_{2.5} \ Sulfur\\ H^+ \end{array}$	1.19 (1.01, 1.42)** 1.23 (0.95, 1.59)** 1.06 (0.87, 1.29)**	18.0 (7.2, 37)*** 2.5 (3.1, 61)*** 18.1 (0.8, 5.9)***
Six Cities ^f (Lower Resp. Symp.)	$\begin{array}{c} PM_{2.5}\\ PM_{2.5} \ Sulfur\\ H^+ \end{array}$	1.44 (1.15-1.82)** 1.82 (1.28-2.59)** 1.05 (0.25-1.30)**	18.0 (7.2, 37)*** 2.5 (0.8, 5.9)*** 18.1 (3.1, 61)***

Table C-2. Effect estimates per variable increments in 24-h concentrations of fine particle indicators $(PM_{2.5}, SO_4^=, H^+)$ from U.S. and Canadian studies

Acute Mortality	Indicator	RR (± CI) per 25 μg/m ³ r PM Increase	Reported PM Levels Mean (Min/Max) [†]		
Decreased Lung Function					
Uniontown, PA ^g	PM _{2.5}	PEFR 23.1 (-0.3, 36.9) (per 25 $\mu g/m^3$)	25/88 (NR/88)		
References: ^a Schwartz et al. (1996a) ^b Burnett et al. (1994) ^c Burnett et al. (1995) O_3 ^d Thurston et al. (1992, 1994) ^e Ostro et al (1993) ^f Schwartz et al. (1994) ^g Neas et al. (1995)		[†] Min/Max 24-h PM indicator level shown in parentheses unless otherwise noted as (± S.D.), 10 and 90 percentile (10,90) or R = range of values from min-max, no mean value reporte [*] Change per 100 nmoles/m ³ ^{**} Change per 20 μ g/m ³ for PM _{2.5} ; per 5 μ g/m ³ for PM _{2.5} sulfur; per 25 nmoles/m ³ for H ⁺ . ^{***} 50th percentile value (10,90 percentile)			

Table C-2. Effect estimates per variable increments in 24-h concentrations of fine particle indicators ($PM_{2.5}$, $SO_4^=$, H^+) from U.S. and Canadian studies (continued)

Type of Health Effect & Location Indicator		Change in Health Indicator per Increment in PM ^a	Range of City PM Levels Means (µg/m ³)
Increased total chronic m	nortality in adults	Relative Risk (95% CI)	
Six City ^b	PM _{15/10}	1.42 (1.16-2.01)	18-47
	PM _{2.5}	1.31 (1.11-1.68)	11-30
	$\mathbf{SO}_4^=$	1.46 (1.16-2.16)	5-13
ACS Study ^c (151 U.S. SMSA)	PM _{2.5}	1.17 (1.09-1.26)	9-34*
	$SO_4^=$	1.10 (1.06-1.16)	4-24
Increased bronchitis in c	hildren	Odds Ratio (95% CI)	
Six City ^d	PM _{15/10}	3.26 (1.13, 10.28)	20-59
Six City ^e	TSP	2.80 (1.17, 7.03)	39-114
24 City ^f	$\mathrm{H}^{\scriptscriptstyle +}$	2.65 (1.22, 5.74)	6.2-41.0
24 City ^f	$\mathbf{SO}_4^=$	3.02 (1.28, 7.03)	18.1-67.3
24 City ^f	PM _{2.1}	1.97 (0.85, 4.51)	9.1-17.3
24 City ^f	PM_{10}	3.29 (0.81, 13.62)	22.0-28.6
Southern California ^g	$SO_4^=$	1.39 (0.99, 1.92)	_
Decreased lung function	in children		
Six City ^{d,h}	PM _{15/10}	NS Changes	20-59
Six City ^e	TSP	NS Changes	39-114
24 City ^{i,j}	H^+ (52 nmoles/m ³)	3.45% (-4.87, -2.01) FVC	—
24 City ⁱ	PM _{2.1} (15 μg/m ³)	3.21% (-4.98, -1.41) FVC	—
24 City ⁱ	${ m SO}_4^{=}(7\mu{ m g/m^3})$	3.06% (-4.50, -1.60) FVC	—
24 City ⁱ	$PM_{10} (17 \ \mu g/m^3)$	2.42% (-4.30,0.51) FVC	_

Table C-3. Effect estimates per increments ^a in annual average levels of fine particle	;
indicators from U.S. and Canadian studies	

^aEstimates calculated annual-average PM increments assume: a 100 μ g/m³ increase for TSP; a 50 μ g/m³ increase for PM₁₀ and PM₁₅; a 25 μ g/m³ increase for PM_{2.5}; and a 15 μ g/m³ increase for SO⁼₄, except where noted otherwise; a 100 nmole/m³ increase for H⁺.

^b Dockery et al. (1993)	^g Abbey et al. (1995a,b,c)
^c Pope et al. (1995)	^h NS Changes = No significant changes.
^d Dockery et al. (1989)	ⁱ Raizenne et al. (1996)
^e Ware et al. (1986)	^j Pollutant data same as for Dockery et al. (1996)
$f \mathbf{D}_{a} = \mathbf{I}_{a} = \mathbf{I}_{a} + \mathbf{I}_{a$	

^fDockery et al. (1996)

*Range of annual median values for subset of 50 cities.

Table C-4. Prospective cohort mortality studies

Source	Health Outcome	Population	Time Period/ No. Units	PM Indicators	PM Mean (µg/m³)	PM Range/ (Std. Dev.)	Sites Per City	Total Deaths	Model Type	PM Lag Structure	Other Pollutants	Other Factors	$\begin{array}{l} \textbf{Relative} \\ \textbf{Risk}^{a} \text{ at} \\ \textbf{SO}_{4} = \textbf{15}, \\ \textbf{PM}_{15} = \textbf{50}, \\ \textbf{PM}_{2.5} = \textbf{25} \end{array}$	RR. Confidence Interval	Elasticity
Abbey et al. (1991)	Total mortality from disease	Calif. 7th Day Adventist	1977-82 Defined by air monitoring sites	24 h TSP >200	102	25-175 (annual avg)	NA	845	Cox proportional hazards	10 yrs	none	age, sex, race, smoking, education, airway disease	0.99 TSP ^a	(0.87-1.13) ^a	NS^{b}
Dockery et al. (1993)	Total mortality	White adult volunteers in 6 U.S. cities ^c	1974-91	$\begin{array}{c} PM_{15} \\ PM_{2.5} \\ SO_4 \end{array}$	29.9 18 7.6	18-47 11-30 5-13	1	1429	Cox proportional hazards	none	none	age, sex, smoking, education, body mass, occup. exposure hypertension ^d , diabetes ^d	1.31 PM _{2.5}	(1.16-2.01) (1.11-1.68) (1.16-2.16)	0.25 0.22 0.23
Pope et al. (1995)	Total mortality	American Cancer Society,	1982-89 PM _{2.5} 50 cities SO ₄ 151 cities	PM _{2.5}	18.2	9-34	1	20,765	Cox proportional hazard	none	none	age, sex, race, smoking, education, body	1.17 PM _{2.5}	(1.09-1.26)	0.117
. ,		adult volunteers in U.S.		SO_4	11 ^e	4-24	1	38,963				mass, occup. exposure, alcohol consumption, passive smoking, climate	1.10 SO ₄	(1.06-1.16)	0.077

^aFor 1,000 h/yr > 200 μg/m³. ^bNS = non significant, confidence limits not shown. ^cPortage, WI; Topeka, KS; Watertown, MA; Harrisman-Kingston, TN; St. Louis, MO; Steubenville, OH. ^dUsed in other regression analyses not shown in this table. ^eValue may be affected by filter artifacts.

Source: PM CD (U.S. EPA, 1996a).

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		-	А	djusted Risl	ks
Risk Factor	PM_{2.5} Data (µ g / m ³)	Crude Risk	All ^a	Men ^a	Women ^a
Residence	_				
Portage	11.0 (1980-7)3 ^b	1.0°	1.0	1.0	1.0
Topeka	12.5 (1980-8)	0.90	1.01	1.04	0.97
Watertown	14.9 (1980-5)	1.16	1.07	0.94	1.22
Harriman	20.8 (1980-7)	1.16	1.17	1.21	1.07
St. Louis	19.0 (1980-6)	1.48	1.14	1.15	1.13
Steubenville	29.6 (1980-7)	1.51	1.26	1.29	1.23
Smoking Status	_				
Current			1.59	1.75	1.54
Previous			1.20	1.25	1.18
No high school education			1.19	1.22	1.13
Body mass index of 4.5			1.08	1.03	1.11

Table C-5. Relative mortality risks in six U.S. cities

^aAdjusted for age, smoking, education, and body mass.

^bPeriod of PM_{2.5}air monitoring.

^cBaseline annual crude death rate = 10.73 per thousand population.

Source: Dockery et al. (1993)

		Standard	Pollutant		
Species	Regr. Coeff.	Error	Range	Rel. Risk	95% CIs (n=6)
PM ₁₅	0.0085	(0.0026)	28.3	1.27	(1.04-1.56)
PM _{2.5}	0.0127	(0.0034)	18.6	1.27	(1.06-1.51)
SO_4^{2-}	0.0297	(0.0081)	8.5	1.29	(1.06-1.56)
TSP	0.0037	(0.0014)	55.8	1.22	(0.99-1.53)
TSP-PM ₁₅	0.0042	(0.0032)	27.5	1.12	(0.88-1.43)
PM ₁₅ -PM _{2.5}	0.0178	(0.0098)	9.7	1.19	(0.91-1.55)
PM _{2.5} -SO ₄	0.0255	(0.0029)	8.4	1.24	(1.16-1.32)
PM ₁₅ -SO ₄	0.0121	(0.0034)	18.1	1.24	(1.05-1.48)
SO_2	0.0093	(0.0032)	19.8	1.20	(1.01-1.43)
NO ₂	0.0126	(0.0046)	15.8	1.22	(1.00-1.49)
1970 TSP	0.0014	(0.00044)	154.0	1.25	(1.03-1.50)

Table C-6. Estimated relative risks of mortality in six U.S. cities associated with a range of air pollutants

Source: U.S. EPA (1996a) recalculations based on results of Dockery et al. (1993).

PM Covariate Model	RR	LCL	UCL
BASE (age, sex)	1.069	0.978	1.168
BASE + pack-years	1.096	1.000	1.201
BASE + pack-years + body-mass-index cats.	1.122	1.022	1.233
BASE + pack-years + body-mass-index cats.+ exercise cats.	1.122	1.017	1.239
STANDARD (age, pack-y., y. lived with smoker, occup., educ., BMI)	1.122	1.017	1.239
STANDARD w. PM_{10} (100) over last 4 years only	1.102	1.001	1.214
STANDARD, subset for former smokers	1.155	0.937	1.424
STANDARD, subset for never smokers	1.116	0.999	1.246
STANDARD, subset for low anti-oxidant vitamin intake	1.175	1.008	1.370
STANDARD, subset for high anti-oxidant vitamin intake	1.055	0.917	1.214
STANDARD, subset for < 4 h/wk outdoors	1.048	0.896	1.227
STANDARD, subset for 4-16 h/wk outdoors	1.122	0.928	1.358
STANDARD, subset for 16+ h/wk outdoors	1.207	1.015	1.436
STANDARD, subset for reported respiratory symptoms	1.321	1.079	1.616

Table C-7. Relative risk of mortality from contributing nonmalignant respiratory causes, for 30 days per year with $PM_{10}>100~\mu g/m^3$

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

			Females			Males	
Pollution Index	Pollution Incr.	RR	LCL	UCL	RR	LCL	UCL
PM ₁₀ >100, d/yr	30 days/yr	1.069	0.936	1.220	1.188	1.030	1.370
PM ₁₀ mean	$50 \ \mu g/m^3$	1.219	0.739	2.011	1.537	0.879	2.688
SO ₄ mean	$15 \ \mu g/m^3$	1.105	0.396	3.086	1.219	0.411	3.619
O ₃ >100 ppb, h/yr	551 h/yr (IQR)	1.01	0.77	1.33	1.20	0.88	1.64

 Table C-8. Relative risk of mortality from contributing nonmalignant respiratory causes, by sex and air pollutant, with alternative covariate model

LCL = Lower 95% Confidence Limit UCL = Upper 95% Confidence Limit.

			Females			Males	
Pollution Index	Pollution Incr.	RR	LCL	UCL	RR	LCL	UCL
PM ₁₀ >100, d/yr	30 days/yr	0.958	0.899	1.021	1.082	1.008	1.162
PM ₁₀ mean	$50 \ \mu g/m^3$	0.879	0.713	1.085	1.242	0.955	1.616
SO ₄ mean	$15 \ \mu g/m^3$	0.732	0.484	1.105	1.279	0.774	2.116
O ₃ >100 ppb, h/yr	551 h/yr (IQR)	0.90	0.80	1.02	1.140	0.98	1.32
SO ₂ mean	3.72 (IQR)	1.00	0.91	1.10	1.05	0.94	1.18

 Table C-9. Relative risk of mortality from all nonexternal causes, by sex and air pollutant, for an alternative covariate model

LCL = Lower 95% Confidence Limit UCL = Upper 95% Confidence Limit.

		Females				Males	
Pollution Index	Pollution Incr.	RR	LCL	UCL	RR	LCL	UCL
PM ₁₀ >100, d/yr	30 days/yr	0.929	0.857	1.007	1.062	0.971	1.162
PM ₁₀ mean	$50 \ \mu g/m^3$	0.841	0.639	1.107	1.219	0.862	1.616
SO ₄ mean	$15 \ \mu g/m^3$	0.857	0.498	1.475	1.279	0.002	1018
O ₃ >100 ppb, h/yr	551 h/yr (IQR)	0.88	0.76	1.02	1.06	0.87	1.29
O ₃ mean	10 ppb	0.975	0.865	1.099	1.066	0.920	1.236
SO_2 mean	3.72 (IQR)	1.02	0.90	1.15	1.01	0.86	1.18

 Table C-10. Relative risk of mortality from cardiopulmonary causes, by sex and air pollutant, for an alternative covariate model

LCL = Lower 95% Confidence Limit UCL = Upper 95% Confidence Limit.

			Females			Males		
Pollution Index	Pollution Incr.	Smoking Category	RR	LCL	UCL	RR	LCL	UCL
PM ₁₀ >100, d/yr	30 days/yr	All ^a	1.05 5	0.65 7	1.69 5	1.831	1.28 1	2.617
PM ₁₀ mean	$50 \ \mu g/m^3$	All	1.80 8	0.34 3	9.51 9	12.38 5	2.55 2	60.107
NO ₂ mean	19.78 (IQR)	All	2.81	1.15	6.89	1.82	0.93	3.57
O ₃ >100 ppb, h/yr	551 h/yr (IQR)	All	1.39	0.53	3.67	4.19	1.81	9.69
		never smoker				6.94	1.12	43.08
		past smoker				4.25	1.50	12.07
O ₃ mean	10 ppb	All	0.80 5	0.43 6	1.48 6	1.853	0.99 4	3.453
SO ₂ mean	3.72 (IQR)	All	3.01	1.88	4.84	1.99	1.24	3.20
		never smokers	2.99	1.66	5.40			

Table C-11. Relative risk of mortality from lung cancer, by sex and air pollutant, for an alternative covariate model

^aAll = both never smokers and past smokers.

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

Pollution Index	Pollution Incr.	Covariate Model or Sub-Group	RR	LCL	UCL
PM ₁₀ >40 µg/m ³	139 d/y (IQR)	standard	4.50	1.31	15.44
$PM_{10} > 50 \ \mu g/m^3$	149 d/y (IQR)	standard	4.96	1.54	16.00
$PM_{10} > 60 \ \mu g/m^3$	132 d/y (IQR)	standard	4.72	1.69	13.18
$PM_{10}\!\!>\!\!80 \ \mu g/m^3$	78 d/y (IQR)	standard	3.43	1.71	6.88
$PM_{10}\!\!>\!\!100\;\mu g/m^3$	30 d/y	standard	2.127	1.454	3.112
PM ₁₀ mean	$50 \ \mu g/m^3$	standard	31.147	3.978	243.85
SO_2 mean	3.7 ppb	standard	2.66	1.62	4.39
NO ₂ mean	2.0 ppb	standard	1.45	0.67	3.14
O ₃ >60 ppb	935 h/y	standard	2.14	0.82	5.62
O ₃ >80 ppb	756 h/y	standard	2.96	1.09	8.04
O ₃ >100 ppb	556 h/y	standard	3.56	1.35	9.42
O ₃ >120 ppb	367 h/y	standard	3.75	1.55	9.90
O ₃ >150 ppb	185 h/y	standard	3.61	1.78	7.35
O ₃ mean	2.1 ppb	standard	2.23	0.79	6.34
$PM_{10} > 100 \ \mu g/m^3$	30 d/y	never smokers	2.102	1.325	3.335
O ₃ >100 ppb	556 h/y	never smokers	4.48	1.25	16.04
O ₃ >100 ppb	556 h/y	past smokers	2.15	0.42	10.89
$PM_{10} > 100 \ \mu g/m^3$	30 d/y	high population density	2.865	1.794	4.574
O ₃ >100 ppb	556 h/y	high population density	10.18	2.44	42.45
SO_2 mean	3.7 ppb	high population density	3.22	1.87	5.54
PM ₁₀ mean	$50 \ \mu g/m^3$	> 80% data from monitors within20 miles of residence	9.256	1.135	75.516
SO ₂ mean	3.7 ppb	> 80% data from monitors within 20 miles of residence	2.18	0.92	5.20

 Table C-12. Relative risk of lung cancer incidence in males, by air pollutant, for

 Adventist health study

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

Source: Beeson et al. (1998).

	nu vonise neurin seuring							
Pollution Index	Pollution Incr.	Covariate Model or Sub-Group	RR	LCL	UCL			
$PM_{10} > 50 \ \mu g/m^3$	149 d/y (IQR)	standard	1.21	0.55	2.66			
$PM_{10} > 60 \ \mu g/m^3$	132 d/y (IQR)	standard	1.25	0.57	2.71			
SO_2 mean	3.7 ppb	standard	2.14	1.36	3.37			
O ₃ >100 ppb	556 h/y	standard	0.94	0.41	2.16			
$PM_{10} > 100 \ \mu g/m^3$	30 d/y	high population density	1.089	0.726	1.633			
SO_2 mean	3.7 ppb	high population density	2.11	1.32	3.38			
PM ₁₀ mean	$50 \ \mu g/m^3$	> 80% data from monitors within 20 miles	2.425	0.310	19.004			
SO ₂ mean	3.7 ppb	> 80% data from monitors within 20 miles	2.52	1.19	5.33			

Table C-13. Relative risk of lung cancer incidence in females, by air pollutant, for Adventist health study

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

Source: Beeson et al. (1998).

Sex	Smoking Status	Study	PM Index	PM Inc.	RR	LCL	UCL
F	NON-SMOKER	Six Cities	PM ₁₀	50	1.280	0.704	2.345
		ACS	PM _{2.5}	25	1.215	1.020	1.440
			\mathbf{SO}_4	15	1.147	1.045	1.261
		AHSMOG	PM_{10}	50	0.879	0.713	1.085
	PAST	Six Cities	\mathbf{PM}_{10}	50	1.999	0.704	5.632
	PAST + CURRENT	ACS	PM _{2.5}	25	1.102	0.898	1.338
			\mathbf{SO}_4	15	1.104	0.977	1.240
	CURRENT	Six Cities	\mathbf{PM}_{10}	50	1.442	0.719	3.166
М	NON-SMOKER	Six Cities	PM_{10}	50	1.568	0.674	3.678
		ACS	PM _{2.5}	25	1.245	1.000	1.554
			\mathbf{SO}_4	15	1.104	0.977	1.247
		AHSMOG	PM_{10}	50	1.242	0.955	1.616
	PAST	Six Cities	PM_{10}	50	1.611	0.930	2.825
	PAST + CURRENT	ACS	PM _{2.5}	25	1.164	1.051	1.297
			SO_4	15	1.104	1.037	1.176
	CURRENT	Six Cities	PM_{10}	50	1.858	1.090	3.166

Table C-14. Relative risk (RR) of total mortality in three prospective cohort studies, by sex and smoking status

LCL = Lower 95% Confidence Limit.

UCL = Upper 95% Confidence Limit.

Sources: Dockery et al. (1993); Pope et al. (1995); Abbey et al. (1999).

Sex	Smoking Status	Study	PM Index	PM Inc.	RR	LCL	UCL
F	NON-SMOKERS	ACS	PM _{2.5}	25	1.585	1.235	2.039
			SO_4	15	1.316	1.147	1.518
		AHSMOG	PM_{10}	50	0.841	0.639	1.107
		AHSMOG - CRC	PM_{10}	50	1.219	0.739	2.011
	PAST + CURRENT	ACS	PM _{2.5}	25	1.276	0.918	1.760
			SO_4	15	1.219	1.008	1.465
Μ	NON-SMOKERS	ACS	PM _{2.5}	25	1.245	0.929	1.668
			SO_4	15	1.205	1.023	1.412
		AHSMOG	PM_{10}	50	1.219	0.862	1.616
		AHSMOG - CRC	PM_{10}	50	1.537	0.879	2.688
	PAST + CURRENT	ACS	PM _{2.5}	25	1.235	1.061	1.440
			SO_4	15	1.126	1.037	1.233
F+M	ALL	Six Cities	PM_{10}	50	1.744	1.202	2.501

Table C-15. Relative risk (RR) of cardiopulmonary mortality in three prospective cohort studies, by sex and smoking status

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

Sources: Dockery et al. (1993); Pope et al. (1995); Abbey et al. (1999).

Sex	Smoking Status	Study	PM Index	PM Inc.	RR	LCL	UCL
F	NON-SMOKERS	ACS	PM _{2.5}	25	0.644	0.203	2.091
			\mathbf{SO}_4	15	1.432	0.731	2.800
		AHSMOG	\mathbf{PM}_{10}	50	1.808	0.343	9.519
	PAST + CURRENT	ACS	PM _{2.5}	25	0.949	0.563	1.595
			SO_4	15	1.074	0.781	1.479
М	NON-SMOKERS	ACS	PM _{2.5}	25	0.483	0.086	2.714
			SO_4	15	1.261	0.501	3.190
		AHSMOG	PM_{10}	50	12.385	2.552	60.107
	PAST + CURRENT	ACS	PM _{2.5}	25	1.123	0.827	1.533
			SO_4	15	1.316	1.104	1.577
F+M	ALL	Six Cities	PM_{10}	50	1.744	0.689	4.390
		ACS	PM _{2.5}	25	1.031	0.796	1.338
			SO ₄	15	1.261	1.082	1.465

Table C-16. Relative risk (RR) of lung cancer mortality in three prospective cohort studies, by sex and smoking status

LCL = Lower 95% Confidence Limit. UCL = Upper 95% Confidence Limit.

Sources: Dockery et al. (1993); Pope et al. (1995); Abbey et al. (1999).

PM Species	Concentration Range (µg/m ³)	Relative Risk Estimate	RR 95% CI	Relative Risk t-Statistic
$SO_4 =$	8.5	1.29	(1.06-1.56)	3.67
$PM_{2.5} - SO_4 =$	8.4	1.24	(1.16-1.32)	8.79
PM _{2.5}	18.6	1.27	(1.06-1.51)	3.73
PM _{15-2.5}	9.7	1.19	(0.91-1.55)	1.81
TSP-PM ₁₅	27.5	1.12	(0.88-1.43)	1.31

Table C-17. Comparison of estimated relative risks (**RR**) for all-cause mortality in six U.S. cities associated with the reported inter-city range of concentrations of various PM metrics

Source: Dockery et al. (1993); U.S. Environmental Protection Agency (1996).

Mortality Cause	$SO_4^{=}$ (Range = 19.9 µg/m ³)			$PM_{2.5}$ (Range = 24.5 μ g/m ³)		
	Relative Risk			Relative Risk	RR 95% CI	RR t-Statistic
All Cause	1.15	(1.09-1.22)	4.85	1.17	(1.09-1.26)	4.24
Cardiopulmonary	1.26	(1.15-1.37)	5.18	1.31	(1.17-1.46)	4.79
Lung Cancer	1.35	(1.11-1.66)	2.92	1.03	(0.80-1.33)	0.38

Table C-18. Comparison of reported $SO_4^=$ and $PM_{2.5}$ relative risks (RR) for various mortality causes in the ACS study

Source: Pope et al. (1995).

PM Index	Study	Subgroup	Relative Risk	t Statistic
PM ₁₀ (50 μg/m ³)	Six Cities	All	1.504 ^a ; 1.530 ^b	2.94 ^a ; 3.27 ^b
		Male Nonsmoker	1.280^{a}	0.81ª
	AHSMOG	Male Nonsmoker	1.242	1.616
PM _{2.5} (25 μg/m ³)	Six Cities	All	1.364 ^a ; 1.379 ^b	2.94 ^a ; 3.73 ^b
		Male Nonsmoker	1.207 ^a	0.81ª
	ACS (50 cities)	All	1.174	4.35
		Male Nonsmoker	1.245	1.96 0
$SO_4 = (15 \ \mu g/m^3)$	Six Cities	All	1.504 ^a ; 1.567 ^b	2.94 ^a ; 3.67 ^b
		Male Nonsmoker	1.359	0.81ª
	ACS (151 cities)	All	1.111	5.107
		Male Nonsmoker	1.104	1.586
	AHSMOG	Male Nonsmoker	1.279	0.960
Days/y with PM ₁₀ >100 (30 days)	AHSMOG	Male Nonsmoker	1.082	2.183
$PM_{10-2.5} (25 \ \mu g/m^3)$	Six Cities	All	1.814 ^a ; 1.560 ^b	2.94 ^{a,c} ; 1.816 ^b
		Male Nonsmoker	1.434ª	0.81ª

Table C-19. Comparison of total mortality relative risk (RR) estimates and T-statistics
for PM components in three prospective cohort studies

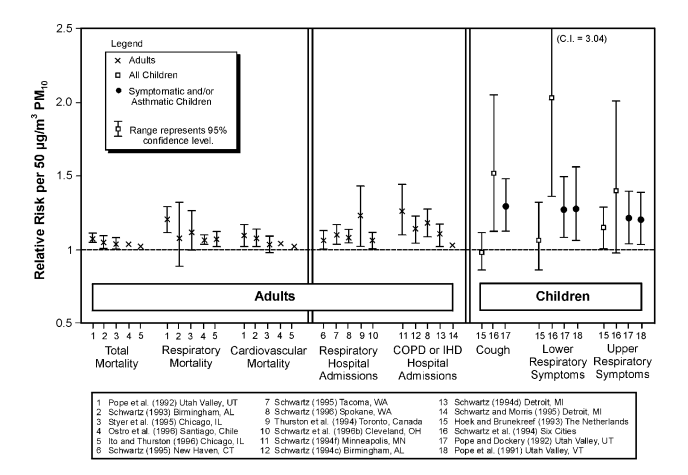
^aMethod 1 compares Portage vs. Steubenville (Table 3, Dockery et al., 1993). ^bMethod 2 is based on ecologic regression models (Table 12-18, U.S. Environmental Protection Agency, 1996). ^cMethod 1 not recommended for PM10-2.5 analysis due to high concentration in Topeka.

PM Index	Study	Subgroup	Relative Risk	t Statistic
PM ₁₀ (50 μg/m ³)	Six Cities	All	1.744ª	2.94 ^a
	AHSMOG	Male Nonsmoker	1.219	1.120
		Male Non CRC	1.537	2.369
PM _{2.5} (25 μg/m ³)	Six Cities	All	1.527ª	2.94 ^a
	ACS (50 cities)	All	1.317	4.699
		Male	1.245	3.061
		Male Nonsmoker	1.245	1.466
$SO_4 = (15 \ \mu g/m^3)$	Six Cities	All	1.743ª	2.94 ^a
	ACS (151 cities)	All	1.190	5.470
		Male	1.147	3.412
		Male Nonsmoker	1.205	2.233
	AHSMOG	Male Nonsmoker	1.279	0.072
		Male Non CRC	1.219	0.357
Days/y with PM ₁₀ >100 (30 days)	AHSMOG	Male Nonsmoker	1.082	1.310
		Male Non CRC	1.188	2.370
$PM_{10-2.5} (25 \ \mu g/m^3)$	Six Cities	All	2.251 ^a	2.94 ^{a,b}

Table C-20. Comparison of cardiopulmonary mortality relative risk (RR)estimates and T-statistics for PM components in three prospective cohort studies("Male Non. - CRC" identifies subjects who died of any contributing nonmalignantrespiratory cause in the AHSMOG study)

^aMethod 1 compares Portage vs. Steubenville (Table 3, Dockery et al., 1993).

^bMethod 1 not recommended for PM10-2.5 analysis due to high concentration in Topeka.



- Figure C-1. Relative risk (RR) estimates for increased mortality and morbidity endpoints associated with 50 μ g/m³ increments in PM₁₀ concentrations as derived from studies cited by numbers listed above each given type of health endpoint. Note the consistency of RR elevations across studies for given endpoint and coherence of RR estimates across endpoints, e.g., higher RR values for symptoms versus hospital admissions and cause-specific mortality.
- Source: PM Staff Paper (1996b). See U.S. EPA (1996b) for full reference citations for each study identified in figure.

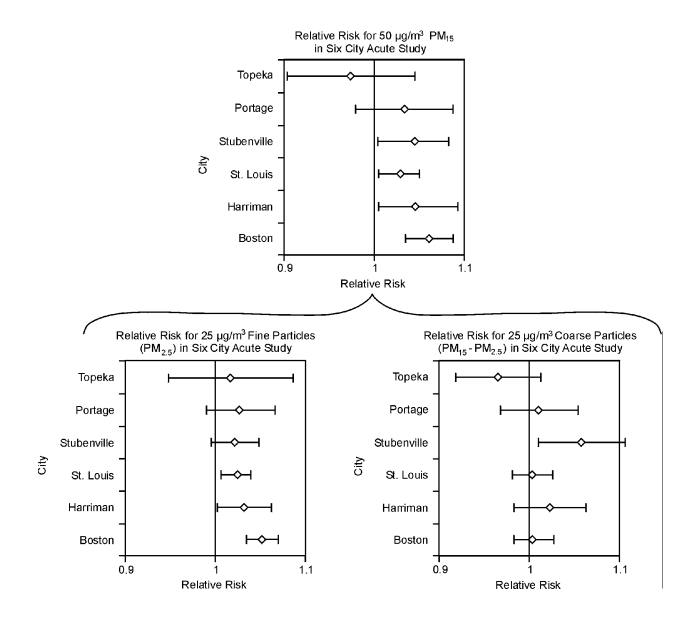


Figure C-2. Relative risks of acute mortality in Harvard Six Cities Study, for inhalable thoracic particles (PM_{15}/PM_{10}) , fine particles $(PM_{2.5})$, and coarse fraction particles $(PM_{15}-PM_{2.5})$. Note that the coarse fraction effects are smaller and statistically non-significant (i.e., lower 95% confidence intervals do not exceed relative risk of 1.0), except in Steubenville where there is high correlation between fine and coarse particles $(R^2 = 0.69)$.

Source: PM CD (U.S. EPA, 1996a) graphical depiction of results from Schwartz et al. (1996).

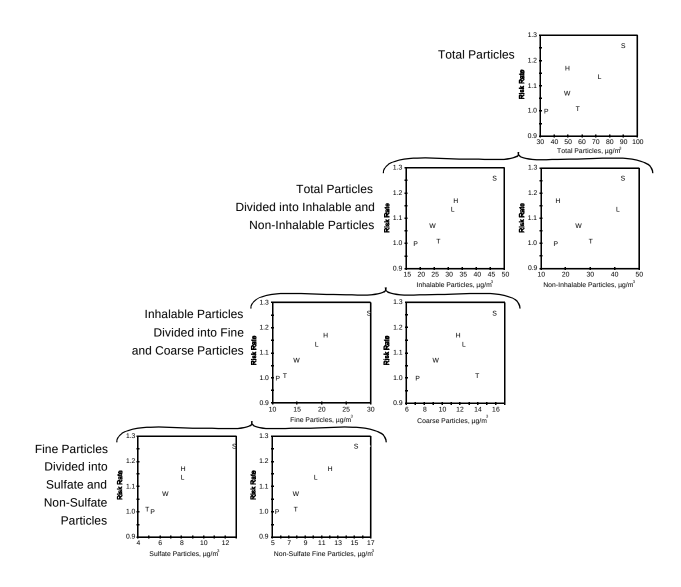


Figure C-3. Adjusted relative risks for mortality are plotted against each of seven longterm average particle indices in the Harvard Six City Study, from largest range (total suspended particles, upper right) through sulfate and nonsulfate fine particle concentrations (lower left). Note that a relatively strong linear relationship is seen for fine particles, and for its sulfate and non-sulfate components. Topeka, which has a substantial coarse particle component of inhalable (thoracic) particle mass, stands apart from the linear relationship between relative risk and inhalable particle concentration.

Source: U.S. EPA (1996a) replotting of results from Dockery et al. (1993).

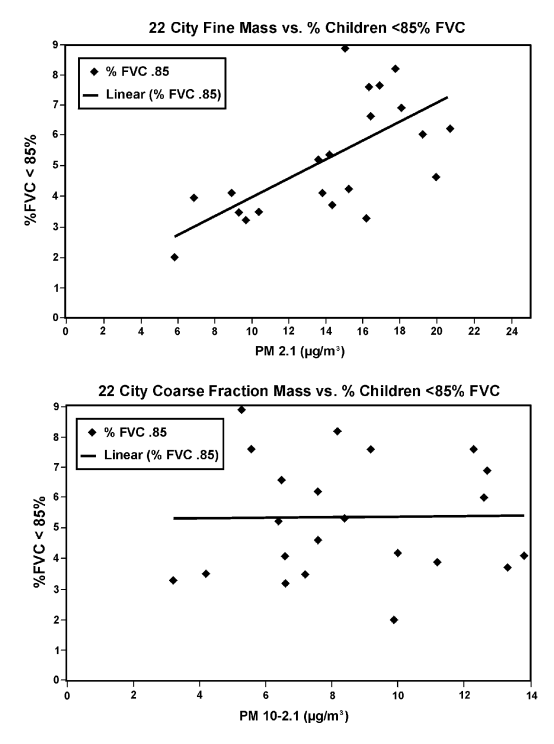


Figure C-4. Percent of children with <85% normal FVC versus annual-average fine $(PM_{2.1})$ particle concentrations and coarse fraction $(PM_{10-2.1})$ levels for 22 North American cities. Note much stronger relationship of fine particles to lung function decrements (top panel) versus for coarse fraction particles (bottom panel).

Source: PM Staff Paper (1996b) graphical depiction of results from Razienne et al. (1996).

C.3. REFERENCES

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