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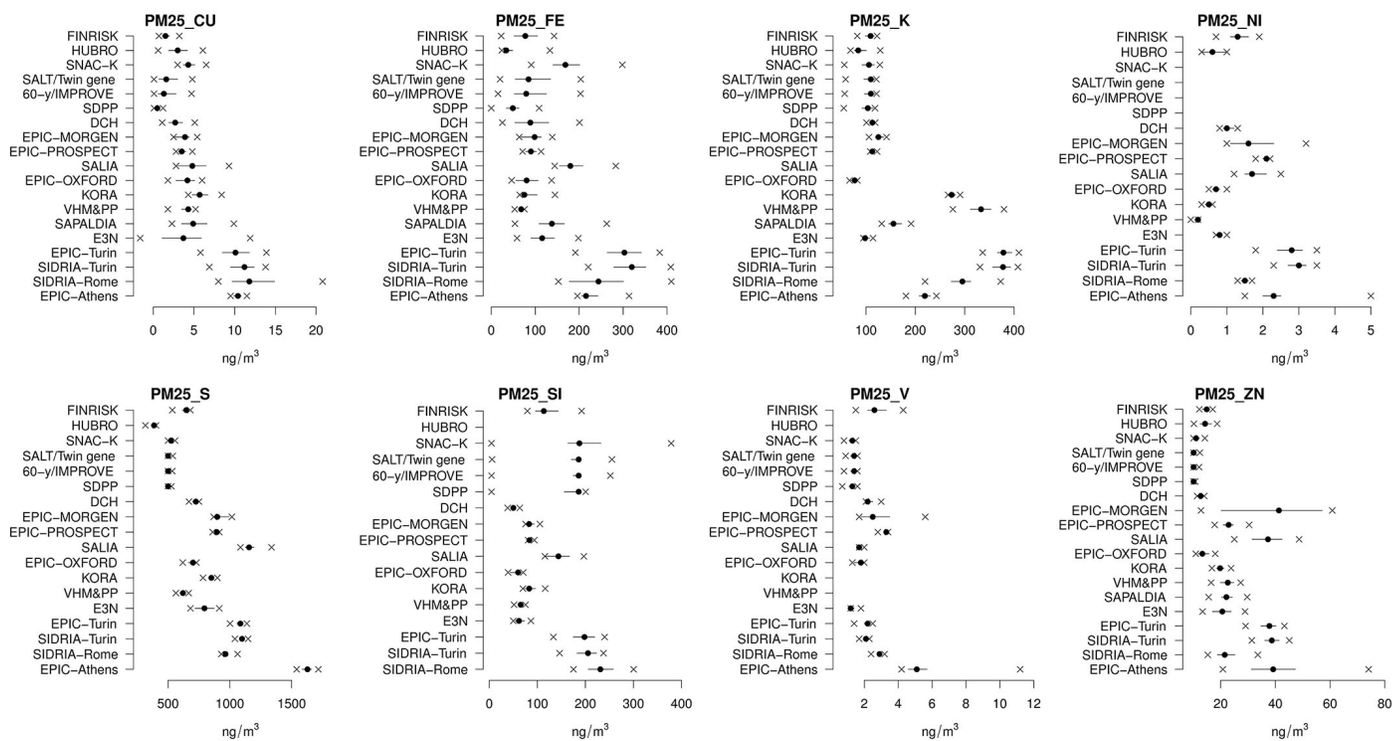
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The Advance Publication of this article contained the wrong version of Figure 2. The correct version is shown below. *EHP* regrets the error.



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Abstract

Background: Studies have shown associations between mortality and long-term exposure to particulate matter air pollution. Few cohort studies have estimated the effects of the elemental composition of particulate matter on mortality.

Objectives: Our aim was to study the association between natural cause mortality and long-term exposure to elemental components of particulate matter.

Methods: Mortality and confounder data from 19 European cohort studies were used. Residential exposure to eight a priori selected components of particulate matter (PM) was characterized following a strictly standardized protocol. Annual average concentrations of Copper (Cu), Iron (Fe), Potassium (K), Nickel (Ni), Sulfur (S), Silicon (Si), Vanadium (V) and Zinc (Zn) within PM size fractions $<2.5\mu\text{m}$ ($\text{PM}_{2.5}$) and $<10\mu\text{m}$ (PM_{10}) were estimated using land-use regression models. Cohort-specific statistical analyses of the associations between mortality and air pollution were conducted using Cox proportional hazards models using a common protocol followed by meta-analysis.

Results: The total study population consisted of 291,816 participants, of which 25,466 died from a natural cause during follow-up (average time of follow-up 14.3 years). Hazard ratios were positive for almost all elements and statistically significant for $\text{PM}_{2.5}$ S (1.14; 95% CI: 1.06, 1.23 per 200 ng/m^3). In a two-pollutant model, the association with $\text{PM}_{2.5}$ S was robust to adjustment for $\text{PM}_{2.5}$ mass, whereas the association with $\text{PM}_{2.5}$ mass was reduced.

Conclusions: Long-term exposure to $\text{PM}_{2.5}$ S was associated with natural cause mortality. This association was robust to adjustment for other pollutants and $\text{PM}_{2.5}$.

Introduction

Studies have shown associations between long-term exposure to particulate matter air pollution and mortality, with exposure characterized as the mass concentration of particles smaller than 10 μm (PM_{10}) or 2.5 μm ($\text{PM}_{2.5}$) (Brook et al. 2010; Brunekreef and Holgate 2002). Although these studies have identified associations between exposure to particulate matter mass and mortality, there is still uncertainty as to which particle components are the most harmful. In addition, particulate matter effect estimates for long-term studies on mortality have differed among studies, and an explanation for this might be differences in the chemical composition of particulate matter (Hoek et al. 2013).

Particulate matter is a heterogeneous mixture varying spatially and temporally in chemical composition related to the sources from which it originates (Kelly and Fussell 2012; Stanek et al. 2011). Components for which associations with a range of health endpoints have been reported in epidemiological and / or toxicological studies include (transition) metals, elemental carbon, inorganic secondary aerosols (sulfate, nitrate), and organic components, but the evidence is not consistent (Kelly and Fussell 2012; Stanek et al. 2011).

Most studies that have assessed mortality in association with exposure to elemental components have been short-term exposure studies and their results have varied considerably (Kelly and Fussell 2012; Stanek et al. 2011). Few studies have investigated mortality in relation to long-term exposure to particle components. A lack of spatially resolved elemental composition measurement data and exposure models for elemental composition partly explains this (De Hoogh et al. 2013). The U.S. Six City and American Cancer Society cohort studies have suggested an association between long-term exposure to sulfate and mortality (Dockery et al. 1993; Krewski et al. 2000; Pope et al. 1995; Pope et al. 2002), but no other particle composition

parameters have been evaluated in these studies. A cohort study among California Teachers found no statistically significant associations between all-cause mortality and long-term exposures to PM_{2.5} and several of its constituents, including elemental carbon, organic carbon (OC), sulfates, nitrates, iron, potassium, silicon, and zinc, although statistically significant associations were reported for more specific outcomes, especially ischemic heart disease mortality (Ostro et al. 2011).

In the framework of the multi-center ESCAPE (European Study of Cohorts for Air Pollution Effects) and TRANSPHORM (Transport related Air Pollution and Health impacts – Integrated Methodologies for Assessing Particulate Matter) projects, we added standardized exposure assessment for air pollution to mortality data from 19 ongoing cohort studies across Europe. Associations of particle mass (PM_{2.5}, PM₁₀, PM_{coarse}, and PM_{2.5} absorbance) and nitrogen oxides (NO₂ and NO_x) with natural cause mortality in the same cohorts have been reported previously (Beelen et al. 2014). We found a statistically significant elevated hazard ratio for PM_{2.5} of 1.07 (95% CI: 1.02, 1.13) per 5 µg/m³. In this paper we report associations with particle elemental composition in 19 European cohorts to assess whether specific components are associated with natural cause mortality. A second aim was to assess whether the previously reported association with PM_{2.5} mass was explained by specific elements. Associations of particle composition and cardiovascular mortality have been published separately (Wang et al. 2014).

Methods

As described earlier, the association between natural cause mortality and particle components was analyzed in each cohort separately following the analysis protocol of the ESCAPE study (Beelen et al. 2014). A common STATA script was used which was explained in a training workshop for all local analysts. Cohort-specific results were sent to the coordinating institute

(IRAS, Utrecht University) for central evaluation. Cohort-specific effect estimates were combined by random-effects meta-analysis. Pooling of the cohort data was not possible due to data transfer and privacy issues.

Study populations

Nineteen cohorts from 12 countries across Europe were selected (Table 1, and Supplemental Material “Description of each cohort and study area” and Figure 1). The study areas of most cohorts consisted of a large city with surrounding smaller rural communities. Some cohorts included large regions of the country such as EPIC-MORGEN in the Netherlands, and the VHM&PP cohort in Austria. All included cohort studies were approved by the institutional medical ethics committees and undertaken in accordance with the Declaration of Helsinki. Each cohort study followed the rules for ethics and data protection set up in the country in which they were based. All participants gave consent according to national rules.

Mortality outcome definition

In all cohorts, follow-up was based upon linkage to mortality registries. Natural cause mortality was defined on the basis of the underlying cause of death recorded on death certificates as ICD-9 codes: 001-779 and ICD-10 codes: A00-R99.

Exposure assessment

Particle composition concentrations at the baseline residential addresses of study participants were estimated by land use regression models following a standardized procedure described elsewhere (Beelen et al. 2013; De Hoogh et al. 2013; Eeftens et al. 2012a). Measurements of particles with aerodynamic diameter $<2.5\mu\text{m}$ ($\text{PM}_{2.5}$) and $<10\mu\text{m}$ (PM_{10}) were performed at 20 sites in each of the study areas. Within each study area, each of the 20 sites was measured during

three two-week periods (during summer, winter, and an intermediate season) within one year. The total measurement period over all study areas was between October 2008 and May 2011. PM filters were weighed before and after each measurement centrally at IRAS, Utrecht University and were then sent to Cooper Environmental Services (Portland, OR, USA) to detect elements. All filters were analyzed for elemental composition using X-Ray Fluorescence (XRF) (De Hoogh et al. 2013). The three two-week measurements were averaged, adjusting for temporal trends using data from a background monitoring site with continuous data (Cyrus et al. 2012; De Hoogh et al. 2013; Eeftens et al. 2012b).

In ESCAPE we *a priori* selected eight of the 48 measured elements for further epidemiological evaluation based upon evidence of health effects (toxicity), representation of major anthropogenic sources, a high percentage of detected samples (>75%), and good precision of measurements (coefficient of variation < 10% for all elements, except Ni and V due to low concentration levels). We selected Cu, Fe, and Zn mainly for (nontailpipe) traffic emissions; S for long-range transport; Ni and V for mixed oil burning/industry; Si for crustal material; and K for biomass burning (Viana et al. 2008). Elements may have multiple sources, so they do not necessarily represent single sources.

Predictor variables for nearby traffic intensity, population/household density, and land use were derived from Geographic Information Systems (GIS), and were evaluated to explain spatial variation of annual average concentrations using land use regression modeling. If values of predictor variables for the cohort addresses were outside the range of values for the monitoring sites, values were truncated to the minimum and maximum values at the monitoring sites.

Truncation was performed to prevent unrealistic predictions (e.g. related to too small distance to roads in GIS) and because we did not want to extrapolate the derived model beyond the range for

which it was developed. Truncation has been shown to improve predictions at independent sites (Wang et al. 2012).

The results of the land use regression models were then used to estimate ambient particle composition concentration at the participants' baseline addresses. A detailed description of the land use regression models for each of the 8 elements is presented in Supplemental Material, Tables S1-S9.

Statistical analyses

Cohort specific analyses

Cox proportional hazards models were used for the cohort specific analyses following the analysis protocol in the ESCAPE study (Beelen et al. 2014). Age was used as the time scale because of evidence of better adjustment for potential confounding by age (Thiébaud and Bénichou 2004). Censoring occurred at the time of death for non-natural causes, emigration, loss to follow-up for other reasons, or at end of follow-up, whichever came first. Air pollution exposure was analyzed as a linear time-invariant variable. Potential confounders were available from questionnaires at baseline. We specified three confounder models with increasing levels of adjustment *a priori*. Confounder models were selected based upon previous cohort studies of air pollution and mortality and availability of data in a majority of the cohorts. The specific variables included as model covariates are listed for each cohort in Supplemental Material, Tables S10–S28. Model 1 included only age (time axis), gender, and calendar time (year(s) of enrollment, continuous for baseline periods of 5 years or less). Model 2 added the following individual level variables (as available for the individual cohorts): smoking status (never/former/current), smoking intensity, smoking duration, environmental tobacco smoke, fruit intake, vegetables intake, alcohol consumption (linear and squared term), body mass index

(BMI) (linear and squared term), educational level (low, medium, high), occupational class (white/blue collar classification), employment status, and marital status. Model 3 added area-level socio-economic status (SES) variables, including mean income, percentage of people with a low income, unemployment rate, and educational level or deprivation index, which were defined for most of the cohorts at the neighborhood or municipality level (see Supplemental Material, Tables S10–S28 for details).

Model 3 was selected as the main confounder model. Only subjects with complete information for Model 3 variables were included in the analyses.

Two-pollutant models were conducted for each element by adjusting for particle mass ($PM_{2.5}$, PM_{10} , PM_{coarse}), $PM_{2.5}$ absorbance, NO_2 , NO_x , and other elements in separate models. As two pollutants may reflect the same source, two-pollutant models representing the independent effect of two pollutants may be difficult to interpret. Therefore, each two-pollutant model was restricted to data from studies for which the correlation between the two pollutants was ≤ 0.7 .

In sensitivity analyses, we added prevalent hypertension and physical activity to Model 3, and additionally adjusted for the classical cardiovascular risk factors prevalent diabetes and cholesterol level. Extended confounder models were used in sensitivity analyses because some potential effects of air pollution might be mediated (e.g. hypertension) or affected (physical activity) by these factors.

All cohort-specific analyses were done in STATA versions 10-12 (StataCorp, College Station, TX, USA).

Meta-analysis

Meta-analyses of cohort-specific effect estimates were conducted using the DerSimonian-Laird method with random effects (DerSimonian and Laird 1986). To keep exposure contrasts broadly comparable among pollutants, hazard ratios (HR) and 95% confidence intervals (CIs) were estimated for fixed increments corresponding to the mean difference between the 10th and 90th percentiles of measured pollutant concentrations across all study areas. Heterogeneity among cohorts was quantified by the I^2 statistic and tested by the X^2 test from Cochran's Q statistic (Higgins and Thompson 2002).

We tested whether effect estimates differed for cohorts for which the land use regression model cross-validation explained variance was smaller or larger than 50% by computing the X^2 test of heterogeneity. In addition, we tested whether effect estimates differed by region of Europe (North: Sweden, Norway, Finland, Denmark; West and Middle: United Kingdom, the Netherlands, Germany, France, Austria, and Switzerland; South: Italy and Greece). We did not perform effect modification analyses for individual-level variables as this paper focuses on differences in effect estimates related to elemental composition. Only sex was an effect modifier for the association between $PM_{2.5}$ and natural mortality in the same cohorts (Beelen et al. 2014).

All tests were two-sided and p values of <0.05 were deemed statistically significant.

All meta-analyses were conducted in STATA, version 12.1 (StataCorp, College Station, TX, USA).

Results

Characteristics of the study population

The total study population consisted of 291,816 participants contributing 4,168,461 person-years at risk (average time of follow-up 14.3 years), of which 25,466 died from a natural cause during follow-up (Table 1). Cohorts were recruited mostly in the 1990s. Cohorts differed in the number of participants, the mean baseline age, and the availability of specific covariate data (Table 2, and Supplemental Material Tables S10–S28). Age, gender, smoking status (current, former, or never smoker), and an area-level SES variable were available for all cohorts. Smoking intensity (average cigarettes/day) and duration (years of smoking) were available as continuous variables for all cohorts with the exception of the VHM&PP (Vorarlberg state) and E3N (Paris and surrounding rural areas) cohorts, for which only smoking status was available. VHM&PP had data on occupation and employment status, but not on education. On average, we had complete Model 3 covariate information for more than 90% of cohort participants.

Air pollution exposure

Substantial variations of estimated annual mean concentrations at participant addresses were observed within and between the majority of cohorts and elements (see Figure 2 and Supplemental Material, Figure S1 for $PM_{2.5}$ and PM_{10} elemental composition concentrations, respectively). The largest within-cohort contrasts were found for Cu, Fe, Si, and Zn, with the largest contrasts generally found in South European study areas. The main exception was Si where the largest within-area contrast was found for the North European study areas (Supplemental Material, Figure S1). The smallest within-cohort contrasts were found for S. Higher concentrations of most elements were observed in Southern study areas. Estimated annual mean S in $PM_{2.5}$ concentrations, for example, show a steady increasing north– south

gradient with averages from 635 ng/m³ for FINRISK, Finland to 1626 ng/m³ for EPIC-Athens, Greece. Correlations between elements and particle mass varied considerably among elements and cohorts; average correlations between elements and mass (in the same PM size fraction) were ~0.5, with a range from ~0.3 to ~0.7 (Supplemental Material, Table S29), indicating that associations with individual elements could be estimated after adjusting for PM mass in the majority of cohorts. .

Good land use regression exposure models were developed for Cu, Fe, and Zn in both fractions (PM₁₀ and PM_{2.5}), as indicated by average cross-validation explained variances (R²) between 55% and 81%, although R² values varied between areas (see Supplemental Material, Tables S1–S9). Traffic variables were the dominant predictors, reflecting nontailpipe emissions (De Hoogh et al. 2013). In general, models for the other elements performed moderately well, with average cross-validation R² values between ~50% and ~60%. However, for PM_{2.5} S the average cross-validation R² was 30% (range 2 to 67%, Supplemental Material, Table S6), consistent with the relatively low spatial variation of sulfur concentrations.

Single pollutant results

Positive HRs were estimated for almost all exposures, with a statistically significant association for PM_{2.5} S (HR = 1.14; 95% CI: 1.06, 1.23 per 200 ng/m³) (Table 3, Figure 3, and Supplemental Material, Figures S2–S15). Borderline statistically significant associations ($p > 0.05$ and ≤ 0.10) were found for PM_{2.5} Si (HR = 1.09; 95% CI: 0.99, 1.09 per 100 ng/m³), PM₁₀ Ni (HR = 1.09; 95% CI: 1.00, 1.19 per 2 ng/m³) and PM₁₀ K (HR = 1.03; 95% CI: 1.00, 1.06 per 100 ng/m³). The evidence for an association was smaller for Zn and V. Estimates did not support associations of mortality with the non-tailpipe traffic pollutants Cu and Fe. In general, HRs based on confounder Model 1 (adjusted for calendar year and gender only) were the highest, while HRs

moved closer to the null after adjustment for individual level confounders (Model 2). Sensitivity analyses showed that especially smoking variables were responsible for this decrease (Beelen et al. 2014). In contrast, additional adjustment for area-level socio-economic status variables (Model 3) had relatively little influence on HRs (Table 3). Cohort specific HRs for PM_{2.5} S were above 1 for all cohorts, except for SDPP and KORA (Figure 3). There was no statistical evidence of heterogeneity among the individual cohort effect estimates for PM_{2.5} S ($I^2 = 0$, $p = 0.94$). Average correlation between PM_{2.5} S and PM₁₀ S over the different cohorts was 0.56 with a range of 0.18-1.00 (data not shown). The HR for PM₁₀ S was also positive (HR = 1.09; 95% CI: 0.99, 1.19 per 200 ng/m³), although not statistically significant (Figure 3).

For the other elements there was more heterogeneity among individual cohort effect estimates, although for most elements heterogeneity was low ($I^2 < 25\%$) to moderate ($I^2 25\text{--}50\%$) and not statistically significant (Table 3 and Supplemental Material, Figures S2–S15).

Two-pollutant results

Results from the two-pollutant models suggested that the associations of elements were generally robust to adjustment for other elements and pollutants (see Supplemental Material, Figures S16 and S17). We also investigated whether the previously reported association between natural cause mortality and PM_{2.5} mass (Beelen et al. 2014) was robust to adjustment for PM_{2.5} S. The median correlation between PM_{2.5} and PM_{2.5} S over the cohorts was 0.53 (range 0.26 – 0.86) (Supplemental Material, Table S29). The combined effect estimate for PM_{2.5} S from the two-pollutant model adjusted for PM_{2.5} did not differ from the single-pollutant model estimate (Table 4). However, the HR for PM_{2.5} was closer to the null and statistically non-significant when adjusted for PM_{2.5} S (HR = 1.07; 95% CI: 1.02, 1.13 vs HR = 1.02; 95% CI: 0.96, 1.09 per 5 µg/m³). In addition, Table 4 shows the two-pollutant model results for PM_{2.5} Si, PM₁₀ K and

PM₁₀ Ni as the single-pollutant associations for these elements were borderline statistically significant. After adjustment for PM_{2.5} S, associations with PM₁₀ Ni (HR = 1.09; 95% CI: 0.98, 1.22 vs HR = 1.06; 95% CI: 0.95, 1.18 per 2 ng/m³) were slightly reduced (Table 4).

Sensitivity analyses

Additional adjustment for hypertension and physical activity, and for diabetes and cholesterol, had little effect on combined HRs compared with Model 3 HRs (see Supplemental Material, Table S30).

Because the VHM&PP cohort had a weight of ~47% in the pooled PM_{2.5} S analyses (Figure 3), we conducted a sensitivity analyses without this cohort. Confidence intervals became slightly wider, but PM_{2.5} S HR remained similar after exclusion of the VHM&PP cohort: (HR = 1.12; 95% CI: 1.01, 1.24 compared with HR = 1.14; 95% CI: 1.06, 1.23 before exclusion). Effect estimates for all elements were similar for the cohorts for which the land use regression model cross-validation explained variance was < 50% or >50% (for example, for PM_{2.5} S, HR = 1.12; 95% CI: 1.01, 1.25; N=14 and HR = 1.16; 95% CI: 1.05, 1.28; N=4, respectively) (p = 0.65). PM_{2.5} S effect estimates were also not statistically different between the cohorts in different regions: 1.17 (95% CI: 0.94, 1.45) for North (N=7); 1.13 (95% CI: 1.04, 1.23) for West and Middle (N=7); and 1.27 (95% CI: 0.92, 1.75) for South (N=4) (p = 0.78). For the other elements also no significant differences were found between effect estimates based on validation R² or region (data not shown).

Discussion

Long-term exposure to PM_{2.5} S was positively associated with natural cause mortality, with no indication of heterogeneity among individual cohort effect estimates.

The association between $PM_{2.5}$ S and mortality was robust to adjustment for co-pollutants including $PM_{2.5}$ mass. The $PM_{2.5}$ mass effect estimate was reduced and became statistically non-significant when adjusted for $PM_{2.5}$ S.

Comparison of S mortality associations with previous studies

Only a few studies have estimated associations of mortality with long-term exposures to particle components. Sulfate has received the most attention in epidemiological studies. Elemental sulfur is assumed to be present as a marker for sulfate. Several cohort studies suggested an association between long-term exposure to sulfate and mortality. An association between sulfate and mortality was reported in the Six Cities study (Dockery et al. 1993). The adjusted HR comparing the cities with the highest and lowest sulfate concentrations (a contrast of $8 \mu\text{g}/\text{m}^3$) was 1.26 (95% CI: 1.08; 1.47), corresponding to a HR of 1.03 (95% CI: 1.01; 1.05) per $1 \mu\text{g}/\text{m}^3$. Within the initial ACS study the adjusted HR of all-cause mortality for areas with the highest and lowest concentrations of sulfate ($19.9 \mu\text{g}/\text{m}^3$ contrast) was 1.15 (95% CI: 1.09, 1.22) (Pope et al. 1995), resulting in a HR of 1.01 (95% CI: 1.00; 1.01) per $1 \mu\text{g}/\text{m}^3$. Pope et al. (2002) investigated additional years of follow-up in the ACS Study and estimated an HR for sulfate and natural mortality of about 1.01 (95% CI: 1.00; 1.01) per $1 \mu\text{g}/\text{m}^3$ (Pope et al. 2002). A recent analysis of the ACS cohort reported that sulfate, elemental carbon, and ozone all had positive and statistically significant associations with all-cause mortality, but sulfate had the most robust association (HR = 1.01 (95% CI: 1.00; 1.01) per $1 \mu\text{g}/\text{m}^3$) (Smith et al. 2009). In the recent National Particle Component Toxicity (NPACT) initiative, a similar risk for the association between sulfur exposure and all-cause mortality (HR 1.09 per $200 \text{ ng}/\text{m}^3$) was estimated using ACS cohort data (Lippmann et al. 2013). Within the NPACT initiative also data from the Women's Health Initiative–Observational Study (WHI-OS) cohort were used to study the

association with cardiovascular mortality and (fatal and non-fatal) cardiovascular events (Vedal et al. 2013). Long-term exposure to air pollutant concentrations was estimated with a national exposure spatial model. No association was found with all cardiovascular deaths and sulfur (HR 1.01; 95%: 0.92, 1.12 per $0.25 \mu\text{g}/\text{m}^3$), but the association with cardiovascular events was statistically significant (HR 1.09; 95%: 1.05, 1.14 per $0.25 \mu\text{g}/\text{m}^3$). A cohort study of ~45,000 active and former female public school professionals in the California Teachers Study investigated the association between mortality and long-term exposures to $\text{PM}_{2.5}$ and several of its constituents, including elemental carbon, organic carbon, sulfates, nitrates, iron, potassium, silicon, and zinc (Ostro et al. 2011). Participants whose residential addresses were within 8 or 30 km of a monitor collecting $\text{PM}_{2.5}$ constituent data were included in the analyses. No statistically significant associations between all-cause mortality and $\text{PM}_{2.5}$ mass or any of its measured constituents were reported. The HR for sulfate was 1.06 (95% CI: 0.97, 1.16) for an interquartile range contrast of $2.2 \mu\text{g}/\text{m}^3$, corresponding to a HR of 1.03 per $1 \mu\text{g}/\text{m}^3$. However, the HR for sulfate and ischemic heart disease mortality was 1.48 (95% CI: 1.20, 1.82) for an interquartile range contrast of $2.2 \mu\text{g}/\text{m}^3$.

The estimated effect of $\text{PM}_{2.5}$ S on natural cause mortality in our study population (HR 1.14 per $0.2 \mu\text{g}/\text{m}^3$ S) corresponds to a HR of 1.24 (95% CI: 1.10, 1.41) per $1 \mu\text{g}/\text{m}^3$ sulfate, assuming all S is present as sulfate (sulfate to S ratio of 3). Our effect estimate is thus much larger than the estimate from the US cohort studies that investigated total mortality. A major difference between our study and these US studies is that our study was based upon contrasts within study areas, whereas the US studies focused on between-area contrasts. Sulfate is mostly formed in the atmosphere by oxidation of gaseous sulfur dioxide (SO_2) emissions (U.S. Environmental Protection Agency 2004). Sulfate is concentrated in fine particles that can be transported over

long distances, resulting in a high regional background with typically small spatial variation within metropolitan areas (U.S. Environmental Protection Agency 2004). Most of our study areas comprised a major city and smaller surrounding communities, with some cohorts covering a larger area (e.g., the Vorarlberg region). Consistently, the exposure contrast in our study was much smaller than in the US studies, both for the S measurements (De Hoogh et al. 2013), and cohort exposures. Measured urban background PM_{2.5} S concentrations were on average 9% higher than regional background concentrations. Concentrations at traffic sites were only 2% higher than at urban background sites. Predictor variables in the land use regression models for PM_{2.5} S included especially traffic at various scales, population or address density, and urban green space (see Supplemental Material, Tables S1–S9). Presumably because of the small measured within-study area contrasts, the average cross-validation R² was 30% for PM_{2.5} S, with a range of 7 – 70%. Because land use regression models were developed for each study area separately, we could not exploit between-study area variations in PM_{2.5} S that would have improved the model performance. In the ESCAPE study, which focuses on within-area contrasts in pollution, these models reflect a combination of variation in primary sulfate emissions and secondary sulfate formation (De Hoogh et al. 2013). Depending on meteorological conditions, SO₂ to sulfate conversion rates of 1-5% per hour have been estimated (U.S. Environmental Protection Agency 2004), implying that some conversion already occurs at scales of 10 - 50 km (a typical wind speed is 10 km/hour). A study in Berlin, Germany documented measurable sulfate formation within 50km of the source (Lammel et al. 2005).

PM_{2.5} mass also was associated with mortality in the three US studies (Dockery et al. 1993; Pope et al. 1995; Pope et al. 2002). However, sulfate concentrations were highly correlated with PM_{2.5} mass concentrations in the US studies, and thus associations between mortality and sulfate may

be difficult to distinguish from associations between mortality and PM_{2.5} mass. The median correlation between estimated PM_{2.5} and PM_{2.5} S over the 19 cohorts in our study was 0.53 (range 0.26 – 0.86), which made it possible to estimate mutually-adjusted associations with PM_{2.5} S and PM_{2.5} mass. The lower correlation in our study probably reflects the finer spatial resolution at which concentrations were estimated. The median correlation of measured within-area contrast in PM_{2.5} and S was very similar (0.6) to the median correlation within cohorts, suggesting that the moderate model R² values for S did not artificially induce the low correlation. Another study that reported evidence of effects of sulfur on mortality was an intervention study in Hong Kong that studied the effects of limiting the sulfur content of fuel oils used in both power plants and vehicles (Hedley et al. 2002). Initial findings indicated a decrease in sulfur dioxide that was associated with prompt and persistent reductions in mortality, suggesting that higher mortality prior to the limitation may have been related to sulfate and/or sulfur dioxide. Subsequent analysis, however, revealed that the reduction in sulfur dioxide was highly correlated with reductions in both vanadium and nickel derived from residual oil emissions (Hedley et al. 2006). In our study correlations between elements were smaller, suggesting that the association between PM_{2.5} S and mortality is not explained by exposure to other elements such as V and Ni. This is also supported by the robust HRs for PM_{2.5} S after adjustment for co-pollutants. However, we cannot rule out the possibility that the association with PM_{2.5} S may be due to other correlated PM components.

Interpretation of S associations

Toxicological studies have provided little support for a causal effect of sulfate, despite fairly consistent associations in epidemiological studies (Kelly and Fussell 2012). Sulfate may indirectly affect health e.g. by solubilizing metals and thereby increasing their bioavailability,

and by catalyzing the formation of secondary organic PM (Kelly and Fussell 2012). We identified associations with small-scale spatial variations in S and we speculate that this may reflect an influence of primary combustion from S containing fuels and serve as a marker of within-city air pollution differences, i.e., between city centers and surrounding areas.

Associations with other elements

None of the other elements evaluated in our analysis were significantly associated with mortality, though HRs were positive for almost all elements. There was greater heterogeneity among individual cohort effect estimates for elements other than $PM_{2.5}$ S, though for most elements the heterogeneity was not statistically significant. There was little evidence of associations with Cu and Fe, which were mainly selected as markers of (non-tailpipe) traffic emissions. Source apportionment studies conducted elsewhere have reported that Fe is mostly associated with road dust and brake abrasion, while Cu is associated with tire and brake abrasion (reviewed by Viana et al. 2008). Our land use regression models had the best fit for these elements because traffic predictors were available and traffic sites were overrepresented in the measurement campaign. Therefore, we believe that the lack of an association in our study is unlikely to be due to exposure measurement error. In our previous analysis of the same set of cohorts we estimated non-significant positive HRs for NO_2 (1.01; 95% CI: 0.99, 1.03 per $10 \mu g/m^3$), NO_x (1.02; 95% CI: 1.00, 1.04 per $20 \mu g/m^3$) and $PM_{2.5}$ absorbance (1.02; 95% CI: 0.97, 1.07 per $10^{-5}/m$), pollutants affected by tailpipe emissions (Beelen et al. 2014).

In single pollutant models we found borderline statistically significant positive associations between natural cause mortality and Si in $PM_{2.5}$, but not Si in PM_{10} , despite substantially higher Si concentrations in the coarse fraction. Source apportionment studies suggest that Si is primarily associated with crustal material in resuspended soil and road dust (Viana et al. 2008). In our

previous analyses we did not find an association between mortality and coarse particles (Beelen et al. 2014).

Source apportionment studies suggest that both V and Ni are linked to crude oil and derived mainly from shipping emissions, and that K is linked to biomass burning (Viana et al. 2008). In single pollutant models we found borderline statistically significant associations for Ni and K in PM₁₀. General industry and port land use were the only predictor variables available for Ni and V in our exposure models. A specific predictor variable for wood smoke was not available (De Hoogh et al. 2013). The lack of more specific predictors in the V, Ni, and K exposure models may have limited our ability to detect element-specific mortality associations for these PM components.

Strengths and limitations

Our study has several strengths: large sample size, broad European coverage, adjustment for a wide range of potential (individual) confounders, and multiple elements with a high percentage of detected samples (>75%) and good precision of measurements in all 19 cohorts (coefficient of variation < 10% for all elements, except Ni and V due to low concentration levels). An advantage compared with previous long-term studies of elemental composition that compared between-city variation and ignored within-city variation, is that we could estimate spatial contrasts at much smaller spatial scales using land use regression models that were developed in a standardized way for all 19 cohorts.

We used data from measurements in 2008–2011 to develop land use regression models that were applied to addresses at baseline, mostly in the mid-1990s. Emissions of sulfur in Europe have been reduced following a series of control measures during the last two decades (Fowler et al.

2007). However, recent studies in the Netherlands, Rome, the UK, and Vancouver have reported that the spatial contrast of nitrogen dioxide air pollution has been stable over a period of 10 years or longer (Cesaroni et al. 2012; Eeftens et al. 2011; Gulliver et al. 2013; Wang et al. 2013). In addition, spatial models for black smoke and sulfur dioxide in the UK provided reasonable predictions, even going back to the 1960s, with a correlation between 1962 and 1991 concentrations of 0.53 for black smoke and 0.26 for sulfur dioxide (Gulliver et al. 2011). However, we cannot rule out the possibility that spatial contrasts for specific components may have been less stable over time.

We did not account for residential mobility during follow-up in the current analyses. In our previous analysis of natural cause mortality in association with particulate matter and nitrogen oxides in the same cohorts, HRs for participants who moved during follow-up did not differ significantly from HRs for the complete study population, though they were slightly higher (Beelen et al. 2014).

We investigated 8 *a priori* selected elements in both the PM_{2.5} and PM₁₀ fractions, so there might be some spurious associations due to multiple comparisons. In addition, correlated elements may act as surrogates for elements that are the actual causes of increased mortality. Although for almost all elements HRs were positive, the association with PM_{2.5} S clearly was the strongest. In addition, the PM_{2.5} S mortality associations were robust to adjustment for other elements, as well as particle mass. In addition, cohort-specific PM_{2.5} S HRs were almost all above 1 (Figure 3) and there was no significant heterogeneity among cohort-specific PM_{2.5} S HRs (Table 3), indicating consistency among the cohort results. The strength of the association, its consistency among cohorts, and its robustness to adjustment decrease the likelihood that the association is a spurious finding.

Differences in the accuracy of exposure estimates could bias effect estimates and standard errors for individual elements. When the measurements of two elements are correlated, part of the association between mortality and the element with more measurement error could be shifted to the estimate of association with the element with less measurement error. Accuracy of exposure estimates may depend on both the precision of the measurements as well as the performance of the exposure models. The eight selected elements were detected in large majority (>75%) of the samples. Measurement precision was best for S, Cu, and Fe but poorer for Ni and V, especially in study areas with low concentration levels (De Hoogh et al. 2013).

Conclusion

In conclusion, long-term exposure to PM_{2.5} S was associated with natural cause mortality. This association was robust to adjustment by other pollutants, including particle mass.

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Table 1. Description of the included cohort studies.

Cohort ^a	N total ^b	N NM ^c	Mean age (years) at baseline \pm SD	Baseline period	Total follow-up time in person-years (mean follow-up)	Study area description
FINRISK, Finland	10,224	602	47.9 \pm 13.2	1992; 1997; 2002; 2007	108,434 (10.6)	Greater Helsinki Area and Turku city and its rural surroundings
HUBRO, Norway	18,102	1182	48.3 \pm 15.2	2000-2001	173,798 (9.6)	City of Oslo
SNAC-K, Sweden	2401	395	70.3 \pm 8.1	2001-2004	15,568 (6.5)	City of Stockholm
SALT/Twin gene, Sweden	5473	581	58.0 \pm 9.9	1998-2002	47,767 (8.7)	Stockholm County
60-y/IMPROVE, Sweden	3612	303	60.4 \pm 0.1	1997-1999	40,612 (11.2)	Stockholm County
SDPP, Sweden	7408	248	47.1 \pm 5.0	1992-1998	102,831 (13.9)	Stockholm County
DCH, Denmark	35,458	3770	56.7 \pm 4.4	1993-1997	469,571 (13.2)	City of Copenhagen and surrounding areas
EPIC-MORGEN, Netherlands	16,446	795	43.9 \pm 10.9	1993-1997	217,722 (13.2)	Cities of Amsterdam, Maastricht and Doetinchem and surrounding rural areas
EPIC-PROSPECT, Netherlands	15,670	1269	57.7 \pm 6.0	1993-1997	202,809 (12.9)	City of Utrecht and surrounding rural areas
SALIA, Germany	4352	618	54.5 \pm 0.6	1985-1987; 1990-1994	81,093 (18.6)	Areas in the cities of Dortmund, Duisburg, Essen, Gelsenkirchen and Herne situated in the Ruhr Area and adjacent towns Borken and Dülmen
EPIC-Oxford, UK	8598	443	45.0 \pm 13.1	1993-2001	110,097 (12.6)	Urban and rural areas in a buffer of 10 km around London-Oxford area
KORA, Germany	8399	673	49.5 \pm 13.8	1994-1995; 1999-2001	88,592 (10.5)	City of Augsburg and two adjacent rural counties
VHM&PP, Austria	117,824	13,081	41.9 \pm 14.9	1985-2005	2,039,328 (17.3)	State of Vorarlberg, excluding high mountain areas (> 600m) and areas within 300m of state border
SAPALDIA, Switzerland	1250	65	42.0 \pm 11.9	1991	20,294 (16.2)	City of Lugano
E3N, France	10,915	516	53.0 \pm 6.8	1993-1996	147,021 (13.5)	City of Paris and surrounding rural areas
EPIC-Turin, Italy	7261	302	50.4 \pm 7.5	1993-1998	97,549 (13.4)	City of Turin
SIDRIA-Turin, Italy	5054	129	44.2 \pm 6.2	1999	55,667 (11.0)	City of Turin
SIDRIA-Rome, Italy	9177	239	44.3 \pm 6.0	1999	102,856 (11.2)	City of Rome
EPIC-Athens, Greece	4192	255	49.4 \pm 11.7	1994-1999	46,852 (11.2)	Greater Athens Area

^aOrder of cohorts is North to South gradient, ^bTotal study population: Number of observations with complete data for all Model 3 (main model) covariates.

^cNumber of deaths from natural-cause mortality.

Table 2. Population characteristics of the included cohort studies at baseline.

Cohort ^a	% women	% never smokers	Cigarettes / day ^b	Years of smoking ^b	BMI (kg/m ²) ^b	Fruit intake ^c	Alcohol intake ^d	% married / living with partner	% low educational level	% employed / self-employed
FINRISK, Finland	54%	45%	3.8 ± 7.8	8.6 ± 12.2	26.4 ± 4.6	66%	0.9 ± 1.3	70%	31%	69%
HUBRO, Norway	56%	46%	6.8 ± 8.4	11.6 ± 14.4	25.7 ± 4.1	40%	51%	50%	18%	73%
SNAC-K, Sweden	60%	44%	7.1 ± 9.5	9.8 ± 15.2	26.0 ± 4.1	NA	22%	54%	21%	29%
SALT/Twin gene, Sweden	56%	39%	8.5 ± 9.7	16.7 ± 17.3	28.6 ± 4.1	NA	NA	68%	22%	NA
60-y/IMPROVE, Sweden	53%	41%	8.0 ± 9.1	15.2 ± 16.4	26.8 ± 4.2	64%	8.9 ± 9.7	72%	28%	51%
SDPP, Sweden	62%	37%	8.5 ± 8.8	12.3 ± 12.4	25.6 ± 4.0	92%	1.3 ± 1.9	84%	26%	92%
DCH, Denmark	54%	36%	6.3 ± 10.4	18.7 ± 17.1	26.0 ± 4.1	183.2 ± 151.2	21.7 ± 22.8	69%	30%	80%
EPIC-MORGEN, Netherlands	54%	35%	10.4 ± 11.1	14.3 ± 13.7	25.2 ± 4.0	171.9 ± 129.2	12.7 ± 18.0	68%	12%	NA
EPIC-PROSPECT, Netherlands	100%	45%	5.7 ± 7.4	15.2 ± 16.5	25.5 ± 4.1	231.6 ± 139.2	9.0 ± 12.4	77%	22%	NA
SALIA, Germany	100%	75%	2.6 ± 6.6	4.4 ± 10.5	NA	NA	NA	NA	29%	NA
EPIC-Oxford, UK	75%	60%	5.5 ± 8.8	7.3 ± 11.5	24.3 ± 4.3	253.6 ± 216.5	10.0 ± 12.3	67%	34%	77%
KORA, Germany	51%	44%	9.2 ± 13.3	12.0 ± 14.2	27.2 ± 4.6	60%	16.3 ± 22.3	76%	13%	58%
VHM&PP, Austria	56%	70%	NA	NA	24.8 ± 4.3	NA	NA	68%	NA	69%
SAPALDIA, Switzerland	56%	45%	11.1 ± 14.4	11.1 ± 13.0	23.8 ± 3.9	NA	NA	58%	11%	81%
E3N, France	100%	49%	NA	NA	22.8 ± 3.3	236.2 ± 162.5	12.4 ± 15.4	NA	5%	NA
EPIC-Turin, Italy	48%	43%	7.2 ± 8.2	17.6 ± 16.3	25.3 ± 3.8	318.2 ± 182.2	18.1 ± 20.3	86%	44%	NA
SIDRIA-Turin, Italy	52%	38%	9.3 ± 10.2	11.3 ± 10.6	NA	NA	NA	95%	18%	72%
SIDRIA-Rome, Italy	53%	35%	10.1 ± 10.5	11.7 ± 10.4	NA	NA	NA	100%	45%	NA
EPIC-Athens, Greece	55%	40%	1.7 ± 15.0	10.8 ± 13.1	27.5 ± 4.5	402.6 ± 258.2	9.2 ± 14.5	78%	24%	67%

^aOrder of cohorts is North to South gradient. ^bMean ± SD. ^cMean ± SD (g/day) or percentage reporting daily fruit consumption. For SDPP it is percentage daily/weekly fruit consumption. ^dMean ± SD (g/day) or percentage reporting daily alcohol consumption. For FINRISK it is number of glasses of alcoholic drink during last week. For SDPP it number of glasses of alcoholic drinks per day. For HUBRO it is the percentage reporting weekly alcohol consumption. NA is not available or available with large number of missings (e.g. BMI in SALIA and smoking variables in E3N).

A detailed description of each cohort can be found in Supplemental Material, Tables S10–S28.

Table 3. Association between natural cause mortality and exposure to elemental composition of PM: Results from random-effects meta-analyses (HRs and 95%-CIs) (using main confounder models 1, 2 and 3).^a

Exposure	Number of cohorts	Model 1 ^b	Model 2 ^b	Model 3 ^b	p-value model 3	I ² (p-value) ^c
PM _{2.5} Cu	19	1.08 (1.00, 1.17)	1.00 (0.94, 1.06)	0.98 (0.92, 1.04)	0.54	16.4 (0.25)
PM ₁₀ Cu	19	1.07 (1.00, 1.15)	1.02 (0.95, 1.08)	1.01 (0.95, 1.07)	0.83	43.5 (0.02)
PM _{2.5} Fe	19	1.12 (1.05, 1.18)	1.04 (0.99, 1.10)	1.03 (0.98, 1.09)	0.20	10.1 (0.33)
PM ₁₀ Fe	19	1.08 (1.02, 1.15)	1.03 (0.97, 1.09)	1.02 (0.97, 1.08)	0.44	43.9 (0.02)
PM _{2.5} Zn	19	1.07 (1.00, 1.15)	1.04 (1.00, 1.08)	1.03 (0.99, 1.08)	0.17	21.4 (0.19)
PM ₁₀ Zn	19	1.09 (1.01, 1.17)	1.04 (1.00, 1.09)	1.04 (0.99, 1.09)	0.18	31.5 (0.09)
PM _{2.5} S	18 ^d	1.29 (1.11, 1.50)	1.16 (1.08, 1.25)	1.14 (1.06, 1.23)	0.003	0 (0.94)
PM ₁₀ S	18 ^d	1.23 (1.07, 1.42)	1.09 (1.00, 1.19)	1.09 (0.99, 1.19)	0.11	29.8 (0.11)
PM _{2.5} Ni	14 ^e	1.12 (1.02, 1.22)	1.05 (0.97, 1.15)	1.05 (0.97, 1.13)	0.27	20.3 (0.23)
PM ₁₀ Ni	17 ^f	1.22 (1.05, 1.41)	1.09 (1.00, 1.19)	1.09 (1.00, 1.19)	0.08	30.3 (0.12)
PM _{2.5} V	15 ^g	1.22 (1.03, 1.44)	1.07 (0.95, 1.20)	1.07 (0.93, 1.23)	0.35	32.5 (0.11)
PM ₁₀ V	18 ^d	1.07 (0.93, 1.24)	1.04 (0.96, 1.12)	1.03 (0.95, 1.12)	0.46	5.7 (0.39)
PM _{2.5} Si	16 ^h	1.18 (1.03, 1.34)	1.10 (0.99, 1.21)	1.09 (0.99, 1.09)	0.10	31.6 (0.11)
PM ₁₀ Si	18 ^d	1.13 (1.00, 1.28)	1.04 (0.97, 1.11)	1.03 (0.97, 1.11)	0.37	47.6 (0.01)
PM _{2.5} K	18 ⁱ	1.06 (0.98, 1.14)	1.05 (0.99, 1.11)	1.07 (0.99, 1.15)	0.12	28.6 (0.13)
PM ₁₀ K	18 ^j	1.05 (0.99, 1.12)	1.03 (1.00, 1.06)	1.03 (1.00, 1.06)	0.08	0 (0.74)

^aHRs are presented for the following increments: 5 ng/m³ PM_{2.5} Cu, 20 ng/m³ PM₁₀ Cu, 100 ng/m³ PM_{2.5} Fe, 500 ng/m³ PM₁₀ Fe, 10 ng/m³ PM_{2.5} Zn, 20 ng/m³ PM₁₀ Zn, 200 ng/m³ PM_{2.5} S, 200 ng/m³ PM₁₀ S, 1 ng/m³ PM_{2.5} Ni, 2 ng/m³ PM₁₀ Ni, 2 ng/m³ PM_{2.5} V, 3 ng/m³ PM₁₀ V, 100 ng/m³ PM_{2.5} Si, 500 ng/m³ PM₁₀ Si, 50 ng/m³ PM_{2.5} K, and 100 ng/m³ PM₁₀ K. ^bModel 1: adjusted for gender and calendar time; Model 2: as in Model 1 also adjusting for smoking status, smoking intensity, smoking duration, environmental tobacco smoke, fruit intake, vegetables intake, alcohol consumption, body mass index, educational level, occupational class, employment status, marital status; and Model 3: as in Model 2 also adjusting for area-level socio-economic status. ^cI² and Cochran's test for heterogeneity for model 3. ^dNo modeled air pollution estimates available for SAPALDIA. ^eNo modeled air pollution estimates available for SNAC-K, SALT/Twin gene, 60-yr/IMPROVE, SDPP. ^fNo modeled air pollution estimates available for HUBRO, SAPALDIA. ^gNo modeled air pollution estimates available for HUBRO, KORA, VHM&PP, SAPALDIA. ^hNo modeled air pollution estimates available for HUBRO, SAPALDIA, EPIC-Athens. ⁱNo modeled air pollution estimates available for SALIA. ^jNo modeled air pollution estimates available for HUBRO.

Table 4. Results from random-effects meta-analyses from single pollutant and two-pollutant models for association with natural cause mortality (using main model 3) (HRs and 95%-CIs).^a

Exposure	Adjusted for	Single pollutant	Two-pollutant
PM _{2.5} S ^b	PM _{2.5}	1.15 (1.06, 1.24)	1.13 (1.03, 1.24)
PM _{2.5} S ^c	PM ₁₀ Ni	1.14 (1.04, 1.25)	1.14 (1.04, 1.25)
PM _{2.5} S ^d	PM _{2.5} Si	1.14 (1.05, 1.23)	1.13 (1.04, 1.22)
PM _{2.5} S ^e	PM ₁₀ K	1.16 (1.06, 1.27)	1.15 (1.05, 1.26)
PM _{2.5} ^b	PM _{2.5} S	1.07 (1.02, 1.13)	1.02 (0.96, 1.09)
PM ₁₀ Ni ^c	PM _{2.5} S	1.09 (0.98, 1.22)	1.06 (0.95, 1.18)
PM _{2.5} Si ^d	PM _{2.5} S	1.09 (0.98, 1.21)	1.08 (0.97, 1.20)
PM ₁₀ K ^e	PM _{2.5} S	1.03 (0.99, 1.08)	1.02 (0.98, 1.06)

^aLimited to studies for which correlation between 2 pollutants was < 0.7. HRs are presented for the following increments: 200 ng/m³ PM_{2.5} S, 5 µg/m³ for PM_{2.5}, 2 ng/m³ PM₁₀ Ni, 100 ng/m³ PM_{2.5} Si, 100 ng/m³ PM₁₀ K. ^bFINRISK and SAPALDIA not included. ^cHUBRO, SALIA and SAPALDIA not included. ^dHUBRO, SAPALDIA and EPIC-Athens not included. ^eFINRISK, HURBO and SIDRIA-Rome not included.

Figure Legends

Figure 1. Cohort locations in which elements were measured.

Figure 2. Description of estimated annual mean PM_{2.5} elemental composition concentrations (ng/μg³) at participant addresses in each cohort. The solid circle and bars shows the median and 25%, 75% percentile of elemental composition concentrations; the x shows the 5% and 95% percentile values.

Figure 3. Adjusted hazard ratio (HR) between natural cause mortality and (A) a 200-ng/m³ increment in PM_{2.5} S and (B) a 200-ng/m³ increment in PM₁₀ S (using main model 3): Results from cohort-specific analyses and from random-effects meta-analyses.

Figure 1.

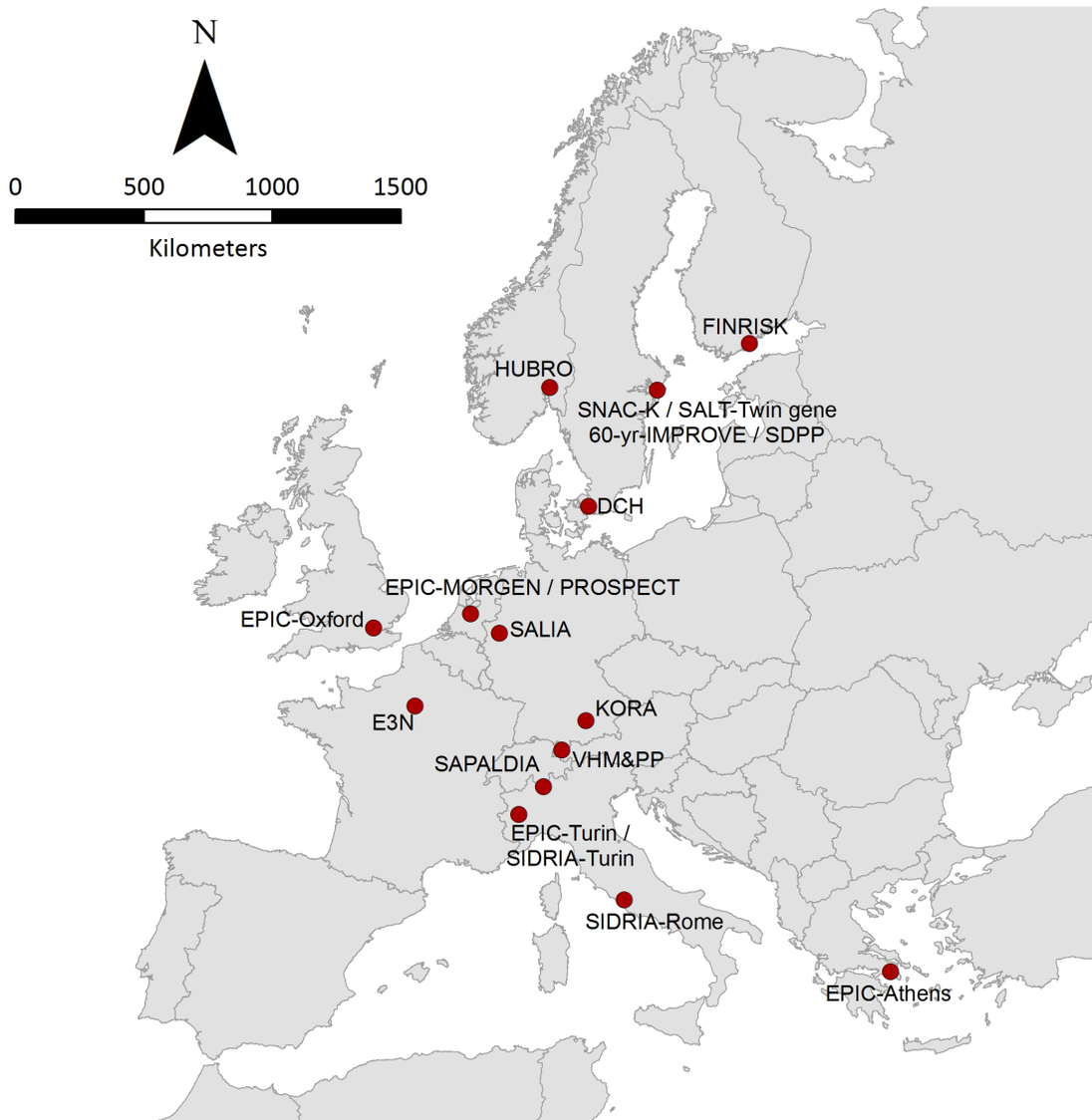


Figure 2.

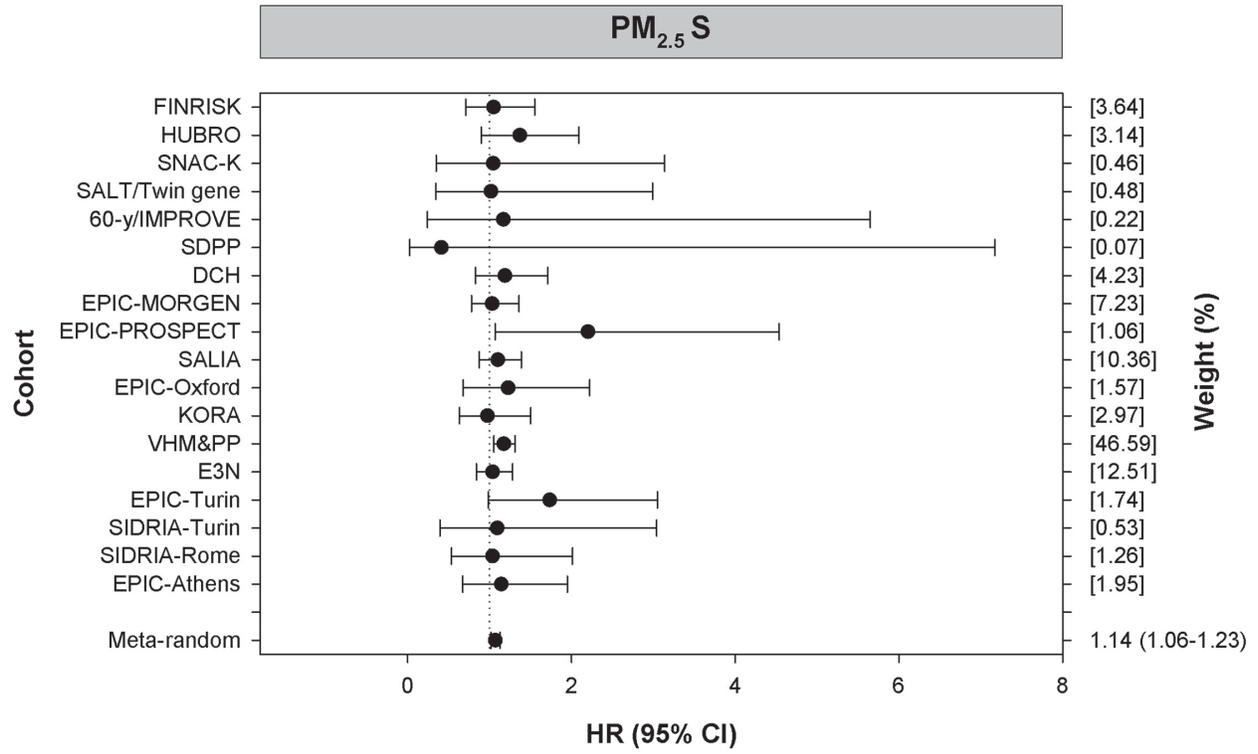


Figure 3A.

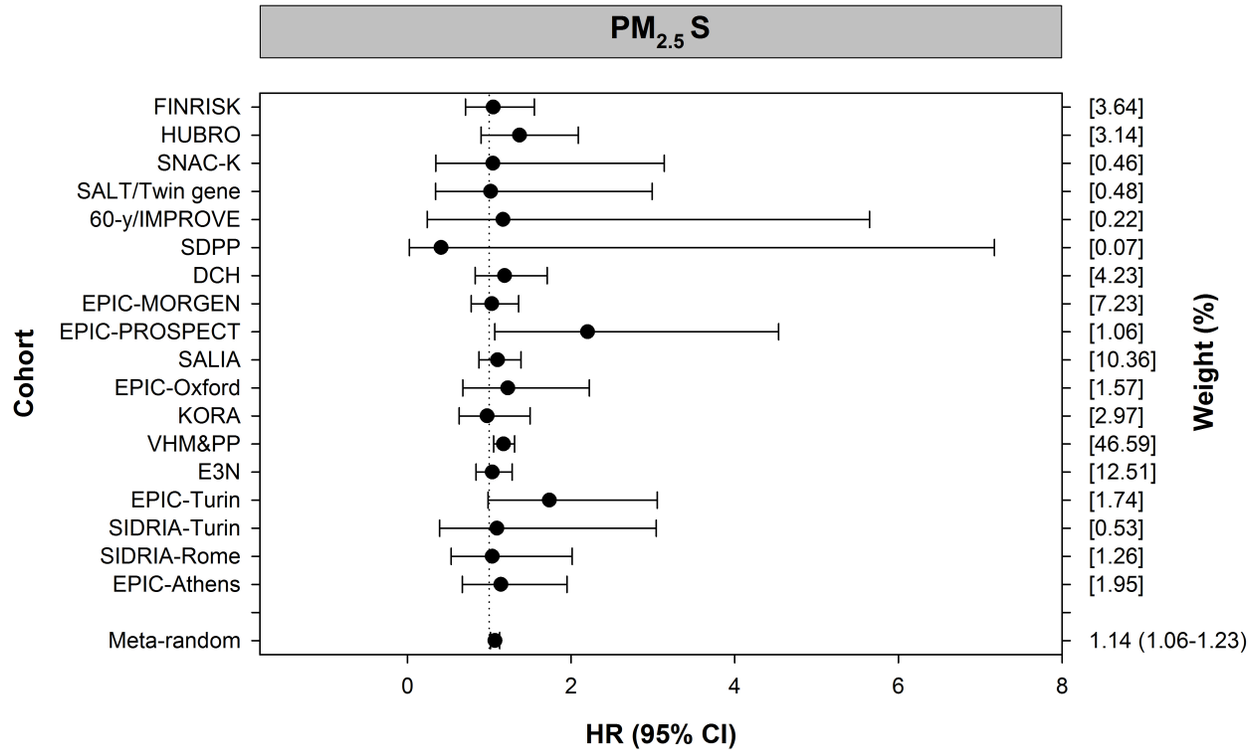


Figure 3B.

