Ambient air pollution and low birthweight: a European cohort study (ESCAPE)


Summary
Background Ambient air pollution has been associated with restricted fetal growth, which is linked with adverse respiratory health in childhood. We assessed the effect of maternal exposure to low concentrations of ambient air pollution on birthweight.

Methods We pooled data from 14 population-based mother–child cohort studies in 12 European countries. Overall, the study population included 74 178 women who had singleton deliveries between Feb 11, 1994, and June 2, 2011, and for whom information about infant birthweight, gestational age, and sex was available. The primary outcome of interest was low birthweight at term (weight <2500 g at birth after 37 weeks of gestation). Mean concentrations of particulate matter with an aerodynamic diameter of less than 2.5 µm (PM2.5), less than 10 µm (PM10), and between 2.5 µm and 10 µm during pregnancy were estimated at maternal home addresses with temporally adjusted land-use regression models, as was PM10 absorbance and concentrations of nitrogen dioxide (NO2) and nitrogen oxides. We also investigated traffic density on the nearest road and total traffic load. We calculated pooled effect estimates with random-effects models.

Findings A 5 µg/m³ increase in concentration of PM2.5 during pregnancy was associated with an increased risk of low birthweight at term (adjusted odds ratio [OR] 1.18, 95% CI 1.06–1.33). An increased risk was also recorded for pregnancy concentrations lower than the present European Union annual PM2.5 limit of 25 µg/m³ (OR for 5 µg/m³ increase in participants exposed to concentrations of less than 20 µg/m³ 1.41, 95% CI 1.20–1.65). PM10 (OR for 10 µg/m³ increase 1.16, 95% CI 1.00–1.35), NO2 (OR for 10 µg/m³ increase 1.09, 1.00–1.19), and traffic density on nearest street (OR for increase of 5000 vehicles per day 1.06, 1.01–1.11) were also associated with increased risk of low birthweight at term. The population attributable risk estimated for a reduction in PM2.5 concentration to 10 µg/m³ during pregnancy corresponded to a decrease of 22% (95% CI 8–33%) in cases of low birthweight at term.

Interpretation Exposure to ambient air pollutants and traffic during pregnancy is associated with restricted fetal growth. A substantial proportion of cases of low birthweight at term could be prevented in Europe if urban air pollution was reduced.

Funding The European Union.

Introduction
Air pollution with ambient particulate matter is one of the most important controllable health threats.1 Maternal exposure to air pollution during pregnancy could increase the risk of preterm birth (<37 weeks of gestation), low birthweight (<2500 g), congenital malformations, and other adverse health effects.2,3

Infants with low birthweight are at greater risk of mortality and morbidity than are infants with higher birthweight.4,5 Low birthweight has been associated with wheezing and asthma in childhood,6,7 and with decreased lung function in adults,8 although findings are not consistent. Infants with low birthweight could have accelerated weight gain in the first 3 months of infancy, which has been associated with asthma symptoms in children aged up to 4 years.9 In addition to active and passive smoking,10 atmospheric pollution exposure is a highly prevalent and controllable potential risk factor for low birthweight.11,12

Meta-analyses13–15 have shown heterogeneity of effects of air pollution across studies, but have suggested that particulate matter with an aerodynamic diameter of less than 2.5 µm (PM2.5) is most consistently associated with low birthweight. Exposure assessment in many previous studies of the effects of air pollution on fetal growth relied on routine air pollution monitoring stations, which do not capture within-city exposure contrasts adequately, possibly resulting in misclassification of exposure and...
possibly reduced risk estimates.\(^3\)–\(^6\,\(^8\)\) In studies in the past 7 years, methods such as land-use regression (LUR) have been applied to improve spatial resolution.\(^2\)–\(^8\,\(^16\)\) The largest body of evidence comes from a series of studies in the Los Angeles basin (CA, USA), where air pollution sources and mixtures could be different from those in smaller urban areas.\(^2\)–\(^8\) Apart from a few studies (usually based on birth registers\(^17\)\(^,\)\(^18\)), many had small sample sizes or poor control of confounders.\(^2\)–\(^8\,\(^16\)\) The individual excess risk of low birthweight reported in these studies was low, but the large proportion of exposed women in the general population warrants an estimation of the attributable risk at the population level.

In this study, we investigated the associations of low exposure to air pollutants with low birthweight at term, birthweight, and head circumference in a population from urban areas. We postulated that increased maternal exposure to ambient air pollution during pregnancy would be associated with intrauterine growth restriction. Additionally, we estimated the proportion of cases of low birthweight at term that were attributable to air pollution.

**Methods**

**Study population**

This study was part of the European Study of Cohorts for Air Pollution Effects (ESCAPE), in which the association between exposure to outdoor air pollution and health is being investigated with prospective cohort studies.\(^19\)\(^,\)\(^20\) We pooled data from 14 European mother–child cohort studies in which birthweight was not part of inclusion criteria: MoBa (Norway); BAMSE (four centres; Sweden); DNBC (Denmark); KANC (Lithuania); BiB (England); ABCD; GENERATION R, and PIAMA (three centres; Netherlands); DUIUSB (Germany); EDEN (two centres; France); APREG (Hungary); Gaspi (Italy); INMA (five centres; Spain); and RHEA (Greece; figure 1). Eligibility criteria were applied in each cohort (appendix p 4). Overall, the study population included 74178 women who lived in the ESCAPE study areas;\(^20\) had singleton deliveries between Feb 11, 1994, and June 2, 2011; and for whom information about home addresses during pregnancy, infant birthweight, gestational age, and sex was available.

Approval was obtained from the ethics committees in every site. All participating women provided informed consent—written or oral, or both, depending on the cohort—for themselves and their children.

**Procedures**

To ensure comparability of the information about maternal and child characteristics from the cohorts, variable definitions were standardised, and quality control was done centrally before data were pooled. We excluded women for whom more than 25% of values for LUR estimates or daily monitoring of air pollution were missing for the whole pregnancy exposure period and for each trimester.

The primary outcome of interest was low birthweight at term (ie, weight <2500 g at birth after 37 weeks of gestation). Other outcomes of interest were term birthweight and head circumference at birth. Information about gestational age, birthweight, head circumference, sex, and mode of delivery was obtained from birth records and questionnaires (appendix p 4).

Because of financial reasons, sampling of particulate matter was not done everywhere. Therefore, data for particulate matter are missing for the BiB and EDEN cohorts, and for four of the five INMA centres. That some data would be missing was known at the start of the project, and was considered during planning. Furthermore, because of scarce data for nitrogen dioxide (NO\(_2\)) and nitrogen oxides (NO\(_x\)) from the centralised routine air monitoring networks in Heraklion, Greece, it was not possible to back extrapolate NO\(_2\) and NO\(_x\) to the RHEA cohort.

Annual mean concentrations of PM\(_{2·5}\) and particulate matter with an aerodynamic diameter of less than 10 μm (PM\(_{10}\)) of between 2·5 μm and 10 μm (PM\(_{2·5-10}\); coarse particulate matter), PM\(_{2·5}\) absorbance (a measure of black carbon), NO\(_2\), and NO\(_x\) were estimated at the maternal home addresses with LUR models.\(^20\)\(^,\)\(^21\) We examined PM\(_{2·5}\) absorbance to establish its role independently from that of PM\(_{2·5}\). PM\(_{2·5}\) absorbance provides a measure of particulate matter attributes distinct from PM\(_{2·5}\). Since 2000, research into air pollution has frequently addressed the LUR estimates to the periods corresponding to each area that were selected to represent spatial variation of air pollution in the residential areas of the participants, allowing development of the LUR models for each pollutant in each study area. However, measurements for the EDEN Nancy cohort were taken in 2002, and for the EDEN Poitiers cohort in 2005. The prediction of LUR models varied between centres and between pollutants.\(^20\) Depending on the area, the LUR models explained 60–88% of the variability in the annual average concentrations of PM\(_{2·5}\), 50–75% of PM\(_{10}\) variability, 87–92% of variability in PM\(_{2·5}\) absorbance, 58–90% of NO\(_2\) variability, and 63–88% of NO\(_x\) variability (cross-validation R\(^2\)).\(^20\) Data from routine monitoring stations were used to temporally adjust the LUR estimates to the periods corresponding to each individual pregnancy and trimester of pregnancy.\(^20\)

In addition to LUR-estimated pollutants, we investigated traffic intensity on the nearest road (vehicles per day) and total traffic load (sum of the lengths of each road segment multiplied by the traffic
intensity [vehicles per day] on all major roads within 100 m of the residence. We accounted for changes of home address during pregnancy when the date of moving and the new address were available (appendix p 2), except for traffic density, which we analysed only for women who did not change home address during pregnancy.

Detailed information about individual characteristics was obtained during the pregnancy through interviews and self-administered questionnaires in most cohorts (appendix p 4). We selected adjustment variables a priori (appendix p 19):^2^ gestational age (not rounded; continuous and quadratic terms), sex, parity (0, 1, 2, and more), maternal height, weight before pregnancy (broken stick model with a knot at 60 kg), mean number of cigarettes smoked per day during second trimester of gestation, maternal age, maternal education (cohort-specific definitions of low, middle, and high), and season of conception (January–March, April–June, July–September, or October–December). We further adjusted models with traffic indicators for background NO\textsubscript{2} concentration. Because air pollution can affect early intrauterine growth, gestational age should preferably be defined from the last menstrual period rather than from early measures of fetal growth. Therefore, gestational age was estimated as the interval between the start of the last menstrual period and delivery when possible (62% of births). Ultrasound-based estimation (16%) was used only when date of last menstrual period was unavailable. When estimates based on last menstrual period or ultrasound were not possible, we used obstetrician estimates (22%). In cohorts for which trimester-specific data for maternal smoking were available, the association with birthweight (and thus presumably the potential for smoking were available, the association with birthweight at term as the percentage of cases that would be averted within the population if PM\textsubscript{2.5} pregnancy concentrations were reduced to 10 µg/m\textsuperscript{3}—the WHO yearly air quality guideline value. We checked that the risk ratio associated with exposure was equivalent to the OR, because low birthweight at term is a rare outcome.

We estimated the population attributable risk of low birthweight at term as the percentage of cases that would be averted within the population if PM\textsubscript{2.5} concentrations were reduced to 10 µg/m\textsuperscript{3}—the WHO yearly air quality guideline value. We estimated the population attributable risk of low birthweight at term as the percentage of cases that would be averted within the population if PM\textsubscript{2.5} concentrations were reduced to 10 µg/m\textsuperscript{3}—the WHO yearly air quality guideline value.

### Statistical analysis

We used logistic regression models to produce odds ratios (ORs) and their 95% CIs for associations between exposure to air pollution and low birthweight at term. We used linear regression models for birthweight and birth head circumference. We restricted analyses of birthweight to term births, because residual plots showed deviation from the models assumptions when preterm deliveries were included (appendix p 20). We analysed birth head circumference for all births. We did pooled analyses using mixed models, including a random effect for centre. Each variable specific to exposure and exposure window was introduced to models with all other potential confounders. As an example of the analysis, we restricted analyses to areas where exposure models had the highest predictive value (defined as a cross-validation R\textsuperscript{2} of more than 0·6). Third, we restricted analyses to one pregnancy per woman, excluding pregnancies other than the first in women with more than one pregnancy during the study period. Fourth, we restricted analyses of birth head circumference to term deliveries. Fifth, we made further adjustment for whether the mother had been born in or had maternal exposure to second-hand smoke during pregnancy only available in a subpopulation). Finally, we did analyses stratified by sex, parity (0, 1, and more), maternal active smoking (no or yes), and maternal education (low, middle, high) to examine potential effect measure modification.

We estimated the population attributable risk of low birthweight at term as the percentage of cases that would be averted within the population if PM\textsubscript{2.5} pregnancy concentrations were reduced to 10 µg/m\textsuperscript{3}—the WHO yearly air quality guideline value. We assessed the impacts of air pollution on birthweight at term as the percentage of cases that would be averted within the population if PM\textsubscript{2.5} pregnancy concentrations were reduced to 10 µg/m\textsuperscript{3}—the WHO yearly air quality guideline value.

Department of Public Health and Preventive Medicine, University of the Basque Country, Bilbao, Spain (A Lurchiund PhD); Department of Environmental Health, Harvard School of Public Health, Boston, MA, USA (I Lepeloe); Department of Community Medicine, Medical Faculty, University of Oslo, Oslo, Norway (Jof P Nafstad); Florence Nightingale School of Nursing and Midwifery, King’s College London, London, UK (P Paterakis); Department of Pulmonology, University Medical Center Groningen, University of Groningen, Groningen, Netherlands (Prof D Postma PhD); Danish Cancer Society Research Centre, Copenhagen, Denmark (O Raaschou-Nielsen PhD, M Sørensen PhD); National Institute of Environmental Health, Budapest, Hungary (P Rudas MD, M Yard MD); Pompeu Fabra University, Barcelona, Spain (Prof J Sunyer); Institute of Epidemiology,
We also calculated the number of cases of low birthweight at term that could be avoided if pregnancy exposure to PM$_{2.5}$ was reduced to 10 µg/m$^3$ in all study areas. We used the formula $N_{Am} = n_m \times (RR_{m-10} - 1) / RR_{m-10}^2$, where $RR_{m-10}$ is the adjusted relative risk (RR) of low birthweight at term comparing women exposed to $m$ and those exposed to 10 µg/m$^3$, and $n_m$ is the number of cases of low birthweight at term in the population with an exposure level $m$. For comparison, we also calculated the population attributable risk corresponding to maternal active smoking.

We used Stata SE (version 12.1) and chose an α of 5% (two tailed).

### Role of the funding source

The sponsors of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

### Results

Mothers were predominately born in the country of their cohort, were non-smokers, and had a mean age of 30 years (table 1, appendix p 8). About 85% of women for whom information about change of address was available did not change address during pregnancy (table 1). Mean gestational age was almost 40 weeks (table 1). 3627 (4.9%) of the 74178 births were preterm. 3087 infants (4.2%) had low birthweight.
PM$_{2.5}$ and PM$_{10}$ concentrations were lower in northern areas than in central and southern areas (figure 2, appendix p 21). The correlations between concentrations of different pollutants during pregnancy were modest to high (appendix p 10).

Risk of low birthweight at term was significantly associated with mean pregnancy exposure to PM$_{10}$, PM$_{2.5}$, NO$_x$ (with pooled data from all areas), and traffic density (table 2). The OR for the association between NO$_x$ and low birthweight at term only in areas where PM$_{2.5}$ concentrations were also available decreased when adjusted for PM$_{2.5}$ concentrations (table 2). The association between PM$_{2.5}$ and low birthweight at term was hardly changed by adjustment for NO$_x$ (table 2). The association between low birthweight at term and PM$_{2.5}$ remained significant when we restricted the population to women exposed to mean PM$_{2.5}$ concentrations of less than 25, 20, or 15 μg/m$^3$ during pregnancy (table 3).

The association between PM$_{2.5}$ and low birthweight at term was stronger for women who gave birth to boys than those who had girls, for women who smoked than in those who did not, and in women of low or medium education than in those of high education (table 4). However, these differences were not significant ($p_{interaction} > 0.1$ for all). After restriction to the 25 313 women for whom gestational age was estimated both with ultrasound and from last menstrual period, the association was slightly stronger with the definition based on last menstrual period (adjusted OR 1·15, 95% CI 0·87–1·52) than when the ultrasound-based definition was used (1·10, 0·97–1·13).

Trimester-specific exposures were highly correlated, which restricted identification of crucial windows of exposure and susceptibility. Generally, the point estimates of ORs were fairly similar in most exposure windows (appendix p 11). The combined effect estimates from random-effect meta-analyses were generally similar to those from the adjusted pooled analysis, and we recorded no important heterogeneity between centres in meta-analyses (appendix p 11).

The population attributable risk estimated for a reduction in PM$_{2.5}$ concentration to 10 μg/m$^3$ during pregnancy corresponded to a decrease of 22% (95% CI 8–33%) in cases of low birthweight at term—ie, 145 (57–223) fewer cases of low birthweight at term in a total of 50 151 term babies. By comparison, the adjusted RR for the association between low birthweight at term and maternal active smoking was 2·26 (1·89–2·69), and the proportion of cases of low birthweight at term attributable to maternal active smoking in the population for whom PM$_{2.5}$ data were available was 14% (10–17%). Because ambient air pollution exposure affects more women than active smoking does (45 430 [91%] of 50 151 women exposed to PM$_{2.5}$, concentration of more than 10 μg/m$^3$ vs 6 237 [12%] who actively smoked), and had a weaker individual effect, the two population attributable risks were similar in size.

All pollutants seemed to reduce birthweight for term births, with the exception of traffic density, but none of the associations were significant (table 5). We noted no strong sign of between-centre heterogeneity for PM$_{10}$, PM$_{2.5}$, PM$_{10}$ absorbance, and traffic indicators in association with birthweight (appendix p 12). Significant reductions in term birthweight were recorded in both unadjusted and adjusted models for PM$_{2.5}$ absorbance exposure during the third trimester and for full pregnancy exposure, and exposure to PM$_{2.5}$, PM$_{2.5}$, and PM$_{10}$ during the third trimester. Associations were also recorded during all exposure periods for NO$_x$ and NO$_x$ in unadjusted models (appendix p 12).

In adjusted models, concentrations of all pollutants and traffic density were associated with reductions in birth head circumference (table 5, appendix p 13). We recorded reductions in head circumference with each 5 μg/m$^3$ increase in PM$_{2.5}$, only in women exposed to PM$_{2.5}$ concentrations of less than 20 μg/m$^3$, but not in those exposed to concentrations of less than 15 μg/m$^3$ (appendix p 14). Adjusted random-effect meta-analyses of birth head circumference showed significant heterogeneity between centres for all pollutants, but not for the traffic indicators (appendix p 13). Overall, in terms of estimated reduction in both birthweight and birth head circumference, the combined effect estimates from the random-effect meta-analyses were smaller than those from the pooled analyses (appendix p 13).

We noted significantly stronger associations with PM$_{2.5}$ concentrations in women who had not moved than in the whole study population for birthweight (change –13 g [–24 to –1] vs –9 g [95% CI –18 to 0]) and birth head circumference (change –0·17 cm [–0·23 to –0·11] vs

![Figure 2: PM$_{2.5}$ concentrations in the ambient air during pregnancy by cohort and overall](image-url)
Further adjusted for the indicated pollutants. Traffic density on nearest street and traffic load models were further adjusted for pollutants on low birthweight at term estimated in pooled analyses using logistic regression with random effect of data for all adjustment variables and indicated exposure are available. Effect of pregnancy mean exposure to air pollutants on low birthweight at term estimated in pooled analyses using logistic regression with random effect of centre adjusted for gestational age (weeks and [weeks/²], sex, parity (0, 1, 2, 3, or more), maternal height, weight before pregnancy (broken stick model with a knot at 60 kg), maternal active smoking during second trimester (cigarettes per day), maternal age, maternal education (low, middle, high), and season of conception (January-March, April-June, July-September, October-December). Participating women exposed to <15 µg/m³ are from M0BA, BARKSE, DNBC, KANC, PIAMA Sabadell, and RHEA cohorts. Participating women exposed to <20 µg/m³ are also from ABCD, GENERATION R, DUSBURG, APRIG, and GASHP cohorts. Participating women exposed to <25 µg/m³ are also from the PIAMA cohort. Number of women who gave birth at term for whom data for all adjustment variables and an exposure estimate for PM ≤2.5 were available for about a third of the cohorts. Information about alcohol consumption during pregnancy was qualitatively unchanged after further adjustments for alcohol intake.

### Discussion

We have shown that ambient air pollutants—particularly PM<sub>2.5</sub>—and traffic density are associated with increases in risk of low birthweight at term and reductions in birthweight and birth head circumference (panel). Our findings suggest that in-utero exposure to ambient air pollution in European urban areas could explain a substantial proportion of cases of low birthweight at term. We used data from a large population with fairly low exposure levels, and increased risks were recorded in the population exposed to levels below the present European Union annual limit of 25 µg/m³.

The estimated 18% increase in risk of low birthweight at term associated with a 5 µg/m³ increase in exposure to PM<sub>2.5</sub>, in our study (RR 1.18) is larger than that reported in meta-analyses (appendix p 17). We recalculated the OR for low birthweight at term from Dadvand and colleagues' study<sup>5</sup> for a 5 µg/m³ increase: 1.02 (95% CI 0.99–1.04), after adjustment for centre-specific covariates. The recalculated OR for Steib and colleagues' analysis<sup>5</sup> was 1.02 (0.99–1.06). The recalculated decrease in birthweight for this study<sup>6</sup> was 12 g (23 to 1), compared with 7 g in our study. The studies included in the meta-analyses<sup>4,7</sup> were

### Table 2: Associations of ambient air pollution during pregnancy with low birthweight at term

<table>
<thead>
<tr>
<th>Total number with term birth*</th>
<th>Number with low birthweight at term</th>
<th>Odds ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposed to &lt;15 µg/m³</td>
<td>21575</td>
<td>1.23 (1.20–1.25)</td>
</tr>
<tr>
<td>Exposed to &lt;20 µg/m³</td>
<td>38017</td>
<td>1.41 (1.40–1.43)</td>
</tr>
<tr>
<td>Exposed to &lt;25 µg/m³</td>
<td>47737</td>
<td>1.21 (1.20–1.23)</td>
</tr>
<tr>
<td>None</td>
<td>50151</td>
<td>1.18 (1.16–1.20)</td>
</tr>
</tbody>
</table>

All effect estimates correspond to an increase of 5 µg/m³ in personal exposure level for PM<sub>2.5</sub> and PM<sub>2.5–10</sub> 20 µg/m³ per m for PM<sub>2.5</sub>, absorbance, 20 µg/m³ per m for NO<sub>x</sub>, 5000 vehicles per day for traffic density, and 40 000 vehicles per day for traffic load. PM<sub>2.5</sub>–10=particulate matter with aerodynamic diameter 2·5–10 µm. PM<sub>2.5</sub>–10=particulate matter with aerodynamic diameter <2·5 µm. NO<sub>x</sub>=nitrogen oxides.

### Table 3: Associations of PM<sub>2.5</sub> concentrations during pregnancy with low birthweight at term, restricted to concentrations less than specified values

<table>
<thead>
<tr>
<th>Total number with term birth*</th>
<th>Number with low birthweight at term</th>
<th>Odds ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PM&lt;sub&gt;2.5&lt;/sub&gt; adjusted for PM&lt;sub&gt;2.5–10&lt;/sub&gt;</td>
<td>48995</td>
<td>1.01 (0.97–1.04)</td>
</tr>
<tr>
<td>PM&lt;sub&gt;2.5&lt;/sub&gt; adjusted for NO&lt;sub&gt;x&lt;/sub&gt;</td>
<td>49285</td>
<td>1.01 (0.97–1.05)</td>
</tr>
<tr>
<td>PM&lt;sub&gt;2.5&lt;/sub&gt; absorbance adjusted for PM&lt;sub&gt;2.5–10&lt;/sub&gt;</td>
<td>49931</td>
<td>1.00 (0.96–1.05)</td>
</tr>
<tr>
<td>PM&lt;sub&gt;2.5&lt;/sub&gt; absorbance adjusted for NO&lt;sub&gt;x&lt;/sub&gt;</td>
<td>49285</td>
<td>1.00 (0.96–1.05)</td>
</tr>
</tbody>
</table>

All effect estimates correspond to an increase of 5 µg/m³ in personal exposure level for PM<sub>2.5</sub> and PM<sub>2.5–10</sub>, 20 µg/m³ per m for PM<sub>2.5</sub>, absorbance, 20 µg/m³ per m for NO<sub>x</sub>, 5000 vehicles per day for traffic density, and 40 000 vehicles per day for traffic load. PM<sub>2.5</sub>–10=particulate matter with aerodynamic diameter 2·5–10 µm. PM<sub>2.5</sub>–10=particulate matter with aerodynamic diameter <2·5 µm. NO<sub>x</sub>=nitrogen oxides. Number of women who gave birth at term for whom data for all adjustment variables and an exposure estimate for PM<sub>2.5</sub> during the whole pregnancy are available. PM=particulate matter with aerodynamic diameter <2·5 µm.

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-0.08 cm [−0.12 to −0.03]; appendix p 15). Results of the other sensitivity analyses were similar to those of the main analyses (appendix p 15). Maternal active smoking was associated with a significant reduction in birth head circumference when compared with mothers who had not smoked (difference −0.31 cm, 95% CI −0.36 to −0.27; appendix p 16). Maternal exposure to second-hand smoke was not associated with a reduction in head circumference (difference 0.0005 cm, 95% CI −0.04 to 0.04; appendix p 16).

Interaction tests suggested no strong evidence of effect measure modification on head circumference between maternal active smoking (binary coding) and maternal exposure to PM<sub>2.5</sub> (p=0.50; appendix p 16). For PM<sub>2.5</sub>, absorbance and NO<sub>x</sub>, effect measure modification with maternal smoking suggested a larger reduction in birth head circumference in children of smokers than in children of non-smokers (appendix p 15). Information about alcohol consumption during pregnancy was available for about a third of the cohorts (appendix p 17). Few women (<1%) consumed at least one glass of alcohol per day. The reported effect estimates of PM<sub>2.5</sub> were qualitatively unchanged after further adjustments for alcohol intake (appendix p 17).
heterogeneous in design—eg, the exposure assessment varied from proximity-based or monitoring-based to model-based—and the assessment of potential confounders varied. Part of the heterogeneity noted could also be explained by the different chemical composition of particulate matter in the various study areas.27

Low birthweight adjusted for gestational duration is the most frequently studied outcome;2–8,17 however, as in other studies,2–8,27 we restricted the analyses of low birthweight to term births as a way to distinguish between babies with low birthweight because of growth restriction and those with low birthweight because of early delivery.28 Exclusion of preterm births from our analyses of birthweight slightly reduced the sample size and might also have resulted in a reduced variability in birthweight compared with the entire study population.

The effects of PM$_{2.5}$ on low birthweight and birth head circumference have not been simultaneously assessed in any previous large studies.5–7,17–18 Head circumference is an important outcome because of the potential effect of air pollution on neurodevelopment,29 and because birth circumference has been associated with intelligence.30 The effect of PM$_{2.5}$ on head circumference has also not been assessed in previous studies.31–34 Adverse effects of other air pollutants on head circumference have been reported in two small studies,31–34 but not in others.35,36

The proportion of cases of low birthweight at term that is attributable to PM$_{2.5}$ exposure in our study population was similar to that attributable to maternal active smoking during pregnancy. The smaller individual effect of PM$_{2.5}$ on low birthweight at term compared with smoking was counterbalanced by the higher prevalence

| Study areas with the land-use regression models of the highest prediction | PM$_{2.5}$ 50 151  –7 (–17 to 2)  34 499  –0.08 (–0.12 to –0.03) PM$_{2.5–10}$ 48 995  –3 (–11 to 6)  33 301  –0.09 (–0.12 to –0.06) PM$_{10}$ 50 151  –8 (–19 to 3)  34 499  –0.13 (–0.18 to –0.09) PM$_{10}$ absorbance 50 835  –3 (–13 to 7)  35 119  –0.18 (–0.22 to –0.13) NO$_2$ 61 452  –1 (–6 to 4)  45 466  –0.08 (–0.10 to –0.07) NO$_X$ 60 254  –1 (–4 to 3)  44 207  –0.06 (–0.07 to –0.05) Traffic density on nearest street 59 030  2 (–1 to 5)  43 109  –0.02 (–0.03 to –0.01) Traffic load on major road 60 254  –1 (–4 to 2)  44 271  –0.02 (–0.05 to 0.00)

Data in parentheses are 95% CIs. Effect of pregnancy mean exposure to air pollutants estimated in pooled analyses using linear regression with random effect of centre adjusted for gestational age (weeks and [weeks]$^2$), sex, parity (0, 1, 2, or more), maternal height, weight before pregnancy (broken stick model with a knot at 60 kg), maternal active smoking during second trimester (cigarettes per day), maternal age, maternal education (low, middle, high), and season of conception (January–March, April–June, July–September, October–December). PM$_{10}$ concentrations not available for BIL and EDEN cohorts. *Number of women who gave birth at term for whom data for all adjustment variables and an exposure estimate for PM$_{10}$ during the whole pregnancy are available. PM$_{2.5}$–10=particulate matter with aerodynamic diameter <2·5 μm. PM$_{10}$=particulate matter with aerodynamic diameter 2·5–10 μm. PM$_{10}$=particulate matter with aerodynamic diameter <10 μm.

### Table 5: Associations between birthweight and head circumference with ambient air pollution during pregnancy

<table>
<thead>
<tr>
<th>Study area</th>
<th>Total number of participants</th>
<th>Change in birthweight (g)</th>
<th>Total number of participants</th>
<th>Change in birth head circumference (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Entire study population</td>
<td>50 151</td>
<td>–7 (–17 to 2)</td>
<td>34 499</td>
<td>–0.08 (–0.12 to –0.03)</td>
</tr>
<tr>
<td>Women who did not change address</td>
<td>33 927</td>
<td>–3 (–11 to 6)</td>
<td>33 301</td>
<td>–0.09 (–0.12 to –0.06)</td>
</tr>
<tr>
<td>Women who participated once</td>
<td>47 192</td>
<td>–8 (–19 to 3)</td>
<td>34 499</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Women for whom data for ethnic origin not missing</td>
<td>49 652</td>
<td>–3 (–13 to 7)</td>
<td>35 119</td>
<td>–0.18 (–0.22 to –0.13)</td>
</tr>
<tr>
<td>Additional adjustment for maternal ethnic origin</td>
<td>49 652</td>
<td>–1 (–6 to 4)</td>
<td>45 466</td>
<td>–0.08 (–0.10 to –0.07)</td>
</tr>
<tr>
<td>Women for whom data for second-hand smoke not missing</td>
<td>47 583</td>
<td>–1 (–4 to 3)</td>
<td>44 207</td>
<td>–0.06 (–0.07 to –0.05)</td>
</tr>
<tr>
<td>Additional adjustment for second-hand smoke</td>
<td>47 583</td>
<td>–1 (–7 to 2)</td>
<td>34 499</td>
<td>–0.08 (–0.12 to –0.03)</td>
</tr>
<tr>
<td>Boys</td>
<td>25 405</td>
<td>–3 (–17 to 2)</td>
<td>21 116</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Girls</td>
<td>24 746</td>
<td>–8 (–19 to 3)</td>
<td>23 020</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Primiparous women</td>
<td>23 229</td>
<td>–10 (–31 to 11)</td>
<td>23 020</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Multiparous women</td>
<td>26 922</td>
<td>–12 (–32 to 8)</td>
<td>23 020</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Non-smoking women</td>
<td>43 914</td>
<td>–1 (–6 to 4)</td>
<td>23 020</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Smoking women</td>
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<td>23 020</td>
<td>–0.13 (–0.18 to –0.09)</td>
</tr>
<tr>
<td>Women with high education</td>
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<tr>
<td>Women with middle education</td>
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<td>23 020</td>
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</tr>
<tr>
<td>Boys</td>
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<tr>
<td>Girls</td>
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<td>23 020</td>
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<td>Primiparous women</td>
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<tr>
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<tr>
<td>Non-smoking women</td>
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</tr>
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</tr>
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<tr>
<td>Women who participated once</td>
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<td>34 499</td>
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</tr>
<tr>
<td>Women for whom data for ethnic origin not missing</td>
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<td>–3 (–13 to 7)</td>
<td>35 119</td>
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<tr>
<td>Additional adjustment for maternal ethnic origin</td>
<td>49 652</td>
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</tr>
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</table>
of PM$_{2.5}$ exposure of more than 10 µg/m$^3$ during pregnancy than that of active smoking. In view of the widespread exposure of pregnant women worldwide to ambient air pollution at similar or even higher concentrations than those assessed in our study, further actions towards improving the ambient air quality in residential areas are recommended.

Ambient air contains a dynamic mixture of pollutants, and the mechanisms by which the airborne inhalable particles, metals, or polycyclic aromatic hydrocarbons could affect fetal growth are unknown. Hypotheses are that air pollutants could cause endocrine disruption, alter placental growth, decrease placental exchange of nutrients and gases, or cause oxidative stress—all of which could possibly lead to altered fetal growth.\(^2\)\(^3\)\(^5\)

In our study, PM$_{2.5}$, PM$_{10}$, NO$_x$, and traffic density were significantly associated with low birthweight at term. The effects of particulate matter were mostly explained by the fine (diameter <2.5 µm) rather than the coarse (2.5–10 µm) fraction. Within-city differences in pregnancy exposure are mostly driven by spatial (rather than temporal) variations. Our finding that within-centre exposure contrasts were related to the risk of low birthweight at term (as shown by the meta-analysis results) shows that local sources are important contributors to the toxic component of the inhaled ambient air and could restrict intrauterine growth.

**Panel: Research in context**

**Systematic review**

We searched PubMed for reports published in English before April 20, 2013, with the search terms “air pollution and birthweight”, “air pollution and fetal growth”, “air pollution and intrauterine restriction”, “air pollution and birth head circumference”, and “air pollution and birth outcomes”. We used three reviews and meta-analyses, last updated in January, 2011,\(^2\)\(^3\)\(^5\) as the basis of our systematic review. At the time our study began in 2008, some evidence of an association between air pollution and fetal growth restriction was available, but many previous studies had poor exposure assessment, small size of included cohorts, and no or little information about potential confounders. We identified no large studies from European populations exposed to low amounts of air pollution.

**Interpretation**

We have shown that fine particulate matter air pollution is associated with an increased risk of low birthweight at term and reduced newborn head circumference. Increased risks were recorded at exposure levels well below the present European Union limits. Our study had a large sample, broad European coverage, standardised fine-scale exposure assessment, and adjustment for a wide range of potential confounders. Urban particulate matter air pollution is widespread, and our results suggest that further reductions could reduce the number of cases of low birthweight at term in Europe.

Our study had three important strengths: the standardised, comprehensive exposure assessment; the harmonised and detailed information about potential confounders; and the large population spread throughout a large geographical area. Detailed information about individual characteristics (eg, maternal stature, parity, ethnic origin, education, and active and passive smoking during pregnancy) was prospectively obtained in a way that enabled us to reduce potential biases through adjustment in a large study population.

Parity was the most important confounder, and it was adjusted for a priori. Indeed, all effect estimates increased when models were not adjusted for parity (data not shown). Few women reported frequent alcohol consumption during pregnancy and the effect estimate of PM$_{2.5}$ varied little after further post-hoc adjustment for alcohol consumption of more than one drink per day in the cohorts for which data for alcohol consumption were available, which suggests that confounding by alcohol consumption was not a big concern in our study. Illegal drug use is likely to be infrequent in the pregnant women included in the cohorts, and we do not believe that it was a strong potential confounder. It is unlikely that prescribed drugs that are known to affect fetal growth would have biased our results, because participation of pregnant women with chronic diseases for which drugs are needed during pregnancy is low in cohort studies.

Most of the pregnant women who were included attended prenatal care, because most participants were recruited at the place and time of prenatal care, which is free in most European countries. After adjustment for active smoking, body-mass index, socioeconomic status, and other factors, our data did not suggest that maternal exposure to second-hand smoke during pregnancy confounded the reported results. To our knowledge, no studies have shown an effect of area-level deprivation on birthweight in which individual factors likely to mediate this effect (eg, active or passive smoking, body-mass index, and individual socioeconomic status) have been controlled for, as in our study. Deprivation score was available for only a few cohorts, and accounting for it hardly changed the adjusted effect of PM$_{2.5}$ concentrations on birthweight (data not shown). We adjusted our models for factors such as individual socioeconomic status defined by education and for several variables (eg, smoking, maternal weight, and maternal height) that possibly mediate the effect of deprivation on health.

The harmonisation of international data allowed us to do pooled analyses. We based our conclusions on the powerful pooled analyses, but we also presented results from the meta-analyses, because this approach has been used in most previous multicentre studies of air pollution in which pooling was not possible. Differences between both approaches are expected, if only because the meta-analysis relied on only within-centre exposure contrasts...
and the pooled analysis made use of within-centre and between-centre exposure contrasts. The point estimates for the effect of PM$_{2.5}$, pregnancy exposure on low birthweight at term were similar for the pooled analyses and the meta-analyses, although the point estimates were attenuated in the meta-analyses when compared with the pooled analysis for birthweight and head circumference. The analyses of low birthweight at term seem to be most robust to model change.

Several pollutants at individual home addresses were assessed in a harmonised manner throughout Europe through LUR modelling using ad-hoc measurement campaigns and detailed data for ambient air quality and road traffic. Thus, we could estimate small-scale spatial exposure contrasts during the pregnancy periods. Air pollution measurements to develop LUR models were taken in 2008–11, whereas pregnancies occurred in 1994–2011—a median of 5 years (range 1–14) earlier. All centres had long-term routine monitoring data that meant we could temporally adjust exposure during the exact pregnancy periods. This approach relied on the assumption that the spatial distribution of the determinants of air pollution (eg, traffic, land use, and household density) had not changed substantially. Spatial contrasts have been shown to be stable with time, but because we sometimes had to rely on a proxy pollutant to extrapolate back to another one, we recognise the potential for exposure misclassification. Assumptions about similar temporal trends for PM$_{1}$ and PM$_{10}$ were made to extrapolate back to PM$_{2.5}$. Our exposure assessment was limited to home address, and exposures elsewhere were not estimated, because detailed information about time-activity patterns or personal measures were not available. Incomplete information about residential mobility could introduce exposure misclassification. However, most women did not move during pregnancy, and analyses restricted to women who did not change home address during pregnancy gave similar results to those reported for the full study population. Confidence intervals were wider in an analysis restricted to non-movers, probably as a result of smaller sample size.

In conclusion, we recorded that exposure to ambient air pollution in pregnancy at levels currently reported in Europe is associated with reduced fetal growth. Our findings suggest that a substantial proportion of cases of low birthweight at term could be prevented in Europe if urban air pollution—particularly PM$_{2.5}$—were reduced.

**Contributors**

MP, JL, EP, MJN, GP, BB, MKog, and RS conceived and planned the study. MP analysed data, reviewed previous reports, interpreted findings, and wrote the first draft of the report. MP, LG-A, CB, and RS harmonised and pooled data and created the variables. LG-A, IA, RMJB, MC, AE, GH, KD, SEH, WN, OR-N, ES, MJN, BB, and RS contributed to the air pollution assessment. LG-A contributed to statistical analyses and did the back- extrapolation of exposure. A-MNA, FB, LC, MV, FF, FG, BG, BH, SEH, VVJ, CK, MKog, UK, PN, WN, DPor, DPos, PR, JS, MJV, TGMV, AW, JW, MKog, and RS designed the cohort study, obtained funding, managed the cohort, and collected data from the cohort’s participants. ADa, ADW, ME, AF-S, MFF, UG, Efvdh, RG, OG, UK, AL, JL, EP, DPor, DPos, MS, DT, MJV, TGMV, MW, and MJV collected data. MJN, MKog, and RS supervised the analysis and contributed to the writing of the report. All authors contributed to critical reading of the report, interpreted data, and approved the report for publication.

**Conflicts of interest**

We declare that we have no conflicts of interest.

**Acknowledgments**

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20 Eeftens M, Beelen R, de Hoogh K, et al. Development of land use regression models for PM(2.5), PM(2.5) absorbance, PM(10) and PM(coarse) in 20 European study areas; results of the ESCAPE project. Environ Sci Technol 2012; 46: 1195–250.


Effects of air pollution on fetal development—more than low birthweight?

There is increasing evidence that inhalation of air pollution not only directly damages the lung, but also has a wide range of effects on other organs.1 In The Lancet Respiratory Medicine, Marie Pedersen and colleagues address the question of whether maternal exposure to ambient air pollution, including fossil-fuel-derived particulate matter, during pregnancy adversely affects the fetus;2 using birthweight as a marker of fetal growth. They combined data from 14 population-based birth cohorts (74 178 mother–child pairs) across Europe, and noted an association between exposure to ambient air pollution and traffic during pregnancy and low birthweight at term (weight <2500 g at birth after 37 weeks of gestation).2 A 5 μg/m³ increase in exposure to particulate matter with an aerodynamic diameter of less than 2·5 μm (PM2·5) was associated with an increased risk of low birthweight at term (adjusted odds ratio 1·18, 95% CI 1·06–1·33).

This association was previously shown in an analysis of cohort data across nine countries.3 However, Pedersen and colleagues’ analysis2 is unique in that great effort has been made to ensure standardised protocols were used for exposure assessment and data analyses. Therefore, its findings are the most robust estimates as yet. Is a reduction in birthweight of a few grams—albeit sufficient to increase the proportion of infants born at term weighing less than 2·5 kg—clinically relevant? The reduction might be evenly spread across all fetal tissues, but also could reflect a clinically significant effect on susceptible fetal organs, such as the brain and lungs. Indeed, Pedersen and colleagues noted small inverse associations between head circumference and ambient air pollution during pregnancy;7 which in turn suggests a small reduction in brain volume.4 The strongest association was with PM10 absorbance (a measure of the carbonaceous fraction of particulate matter), with a change in head circumference of −0·18 cm (95% CI −0·22 to −0·13) for every increase of 10–5 per m.2

Although fetal lung development was not directly assessed in the study, research in animals suggests that maternal exposure to particulate matter alters fetal lung development. Mauad and colleagues5 reported that postnatal exposure of newborn mice to traffic-derived particulate matter did not reduce the ratio of lung surface to volume at ages 15 and 90 days, but they did record a significant reduction when exposed to particulate matter during both the prenatal and postnatal periods. The mechanism whereby deposition of particulate matter in the maternal lung affects the fetus is unclear, but could be via the release of pro-inflammatory mediators into the systemic circulation. The placenta could be an important target for these mediators; an inverse association between maternal exposure to particulate matter and placental weight has been reported.1 Additional evidence to support a causal link between maternal inhalation of air pollutants and fetal lung development has been provided by a pooled analysis6 from eight birth cohorts, including one of the cohorts that was part of Pedersen and colleagues’ analysis (BAMSE). This analysis showed that maternal smoking in pregnancy has an independent effect on the risk of wheeze and asthma at age 4–6 years. One of the strengths of Pedersen and colleagues’ assessment is that the reduction in birthweight has to be a result of a perturbation in fetal growth. By contrast, the independent association between maternal exposure to particulate matter and respiratory symptoms in infants and children is difficult to assess, because maternal and infant exposures to particulate matter are highly correlated.7 One way to overcome this difficulty would be to measure lung function soon after birth.8

Pedersen and colleagues’ study2 also raises a question about the effect of maternal exposure to particulate matter on birthweight and infant respiratory health in countries where women are exposed to high concentrations of particulate matter from biomass smoke. A study in Zimbabwe showed that babies born to mothers cooking with wood, dung, or straw were 175 g lighter (95% CI 50–300) than were those born to mothers using cleaner fuels. Therefore, maternal exposure to biomass smoke could be a major driver of infant respiratory morbidity and potentially mortality.

What are the policy implications of Pedersen and colleagues’ findings? Public awareness that air pollution is an important health issue is already high. In a 2013 European Union commission survey...
of 25 525 adults, more than half the participants thought that air quality had deteriorated in the past 10 years, and more than three-quarters thought that the European Union should propose additional measures to address problems related to air quality. Dissemination of Pedersen and colleagues’ results to the wider public could therefore further increase the pressure on policy makers to reduce exposure of urban populations to particulate matter. Difficult decisions still need to be made. The introduction of the low emission zone in London, UK, has had little effect on concentration of particulate matter, although the vehicle mix has been altered. UK policy makers have shied away from radical solutions to the issue, such as changing diesel-powered black cabs (which contribute 20% of London’s locally generated particulate matter) to cleaner petrol-powered alternatives.

Overall, maternal exposure to traffic-derived particulate matter probably increases vulnerability of their offspring to a wide range of respiratory disorders in both infancy and later life. Should pregnant women therefore act to reduce their own exposure to particulate matter? Small changes in behaviour—eg, avoidance of pollution hotspots such as those around road junctions—might significantly reduce individual exposure to traffic-derived particulate matter. However, until the determinants of individual exposure to particulate matter within cities are clearly defined, no specific advice to pregnant women should be given. In 2014, the effects of air pollution on fetal and infant respiratory health will be reviewed by a UK Royal College of Physicians Working Party on air quality and life effects. The challenge for this group and other academic committees is to present data for the effects of air pollution on fetal health so that they drive policy change, but do not increase the anxiety of individual women.

Jonathan Grigg
Blizard Institute, Queen Mary University of London, London E1 2AT, UK
j.grigg@qmul.ac.uk
I am co-chair of the Royal College of Physicians’ Working Party on air quality and life effects.