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Prenatal and early life exposure to traffic pollution and cardiometabolic health in childhood

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Summary

Background: Prenatal exposure to traffic pollution has been associated with faster infant weight gain, but implications for cardiometabolic health in later childhood are unknown.

Methods: Among 1418 children in Project Viva, a Boston-area pre-birth cohort, we assessed anthropometric and biochemical parameters of cardiometabolic health in early (median age 3.3 years) and mid- (median age 7.7 years) childhood. We used spatiotemporal models to estimate prenatal and early life residential PM_{2.5} and black carbon exposure as well as traffic density and roadway proximity. We performed linear regression analyses adjusted for sociodemographics.

Results: Children whose mothers lived close to a major roadway at the time of delivery had higher markers of adverse cardiometabolic risk in early and mid-childhood. For example, total fat mass was 2.1 kg (95%Cl: 0.8, 3.5) higher in mid-childhood for children of mothers who lived <50 m vs. \geq 200 m from a major roadway. Black carbon exposure and traffic density were generally not associated with cardiometabolic parameters, and PM_{2.5} exposure during the year prior was paradoxically associated with improved cardiometabolic profile.

Conclusions: Infants whose mothers lived close to a major roadway at the time of delivery may be at later risk for adverse cardiometabolic health.

Keywords: Adiposity, air pollution, cardiometabolic health, PM_{2.5}, pregnancy, traffic.

Abbreviations: BC, black carbon; BMI, body mass index; CI, confidence interval; HOMA-IR, homeostasis model assessment for insulin resistance; LMP, last menstrual period; PM_{2.5}, fine particulate matter; RAs, research assistants; SD, standard deviation; SES, socioeconomic status; SS, subscapular; TR, triceps; UFP, ultrafine particles.

Introduction

Childhood obesity is epidemic, recalcitrant to treatment, and associated with costly comorbidities, including adverse cardiometabolic health that tracks into adulthood (1). The prenatal and early life environment influences propensity for excess adiposity (2), and it is a priority to identify remediable early life environmental triggers.

Air pollution is one environmental exposure that may promote adiposity. After release from automobiles and power plants, gaseous and particulate air pollutants with an aerodynamic diameter less than $2.5 \,\mu m \,(PM_{2.5})$ enter the airways and may induce adiposity and dysmetabolism through endothelial dysfunction, inflammation and oxidative stress (3). In rodents, $PM_{2.5}$ exposure altered adipokine secretion and increased adipose inflammation, visceral adiposity and insulin resistance (4,5).

Despite a convincing rodent literature, there has been limited investigation of PM2.5 on cardiometabolic health in human studies. Prior cohorts have demonstrated an association between air pollution exposure and obesity in childhood (6-9) but included limited investigation of adipose distribution and no consideration of cardiometabolic biomarkers. Population-based studies in children (10-12) and adults (13) have linked air pollution exposure with insulin resistance but lacked consideration of prenatal exposures despite emerging evidence that in utero air pollution exposure may prime offspring for adiposity (8,14). Late prenatal exposure to traffic pollution was associated with faster infant weight gain in our prior analysis of the Boston-area Project Viva cohort (14), but whether these weight-promoting effects persist throughout childhood and whether exposure is also associated with adverse cardiometabolic health in childhood is unclear.

In the present analysis, our primary objective was to evaluate the extent to which late prenatal exposure to $PM_{2.5}$ and black carbon (BC) (a traffic-related $PM_{2.5}$ component), as well as residential traffic density and roadway proximity, were associated with anthropometric and biochemical markers of adiposity and insulin resistance in early and mid-childhood. We also evaluated postnatal, proximate pollution exposures. We hypothesized that air pollution exposure would be associated with an adverse cardiometabolic profile.

Methods

Study population and design

Participants were recruited to Project Viva, a prospective cohort study of prenatal exposures and offspring health, from 1999 to 2002 during their first prenatal visit to Atrius Health in Eastern Massachusetts (15). Of 2128 participants with a live singleton offspring, 1418 had data for at least one exposure and one outcome studied. We included a subset in each analysis based primarily on available outcome data (Figure S1). As compared with those without follow-up, mothers of children who attended early and mid-childhood visits were more likely to be nonsmokers, college

graduates and have higher birth weight-forgestational age infants (Table S1).

Mothers provided informed consent at enrollment and for their child at each in-person visit. Institutional Review Boards of participating institutions approved the study.

Air pollution exposures

Participants provided their residential address at enrollment (median 9.9 weeks gestation) and updated it at study visits at the end of the second trimester, soon after birth and during their child's infancy (median: 6 months of age), early childhood (median: 3.3 years of age) and mid-childhood (median: 7.7 years of age). Our estimates of residential BC and $PM_{2.5}$ exposure accounted for moves during exposure windows of interest.

We estimated daily BC exposure with a land-use regression model (mean 'out-of-sample' 10-fold cross-validation $R^2 = 0.73$) (16). We used aerosol optical depth data to estimate PM_{2.5} exposure at 10 km × 10 km spatial grid resolution (mean daily 'out-of-sample' 10-fold cross-validation $R^2 = 0.87$ for days with aerosol optical depth data and 0.85 for days without) (17). To obtain third trimester exposure estimates, we averaged daily exposures from the 188th day (i.e. - 27 weeks gestation) after the last menstrual period (LMP) to the day before birth. To obtain exposure estimates for the year prior to the health outcome measurement, we averaged daily exposures over 365 days prior to the in-person visit (anthropometric outcomes) or blood draw (biomarker outcomes). We assigned exposures to addresses where model predictions were available (Eastern Massachusetts for the BC model and New England for the PM_{2.5} model) for at least 90% of days in an exposure period. We also examined associations using our model for PM_{2.5} exposure at $1 \text{ km} \times 1 \text{ km}$ spatial grid resolution (18), available after 2003. Results using this model were similar, and because estimates were not available for prenatal time periods, we present all results using the $10 \text{ km} \times 10 \text{ km} \text{ PM}_{2.5}$ model.

We used the 2002 road inventory from the Massachusetts Executive Office of Transportation to calculate traffic density by multiplying annual average daily traffic (vehicles/day) by length of road (km) within 100 m of each participant's residential address. We used 2005 ESRI Street MapTM North America ArcGIS 10 Data and Maps to estimate home roadway proximity as distance to Census Feature Class Code A1 or A2 roads (i.e. highways).

Assessment of child anthropometry and cardiometabolic biomarkers

Research assistants (RAs) measured participants' weight in light clothing using an electronic scale (Tanita, Arlington Heights, IL, USA) and height without shoes using a stadiometer (Shorr Productions, Olney, MD, USA). We calculated age-specific and sex-specific body mass index (BMI) *z*-scores from CDC 2000 reference data. RAs used Holtain calipers (Cross-well, UK) to measure subscapular (SS) and triceps (TR) skinfold thicknesses, and we calculated the sum (SS + TR) of the skinfold thicknesses. RAs measured waist circumference underneath clothing using a nonstretchable measuring tape (Hoechstmass Balzer GmbH, Sulzbach, Germany). We measured total and truncal fat mass using a Hologic DXA scan (Bedford, MA, USA).

In early and mid-childhood, we measured plasma leptin and adiponectin concentrations, and in mid-childhood, plasma fasting glucose and insulin, as previously described (19). We calculated the homeostasis model assessment for insulin resistance (HOMA-IR) [(glucose (mg/dL) x insulin (mU/L))/405].

Covariates

We obtained mothers' age, race/ethnicity, education and smoking habits at study enrollment. We calculated pre-pregnancy BMI from self-reported weight and height. Women underwent a two-tiered glucose screening test during pregnancy, as previously described (20). We obtained infant sex, birth weight and date of delivery from the hospital medical record. We calculated length of gestation by LMP and birth weight-for-gestational age and sex *z*-score from a US national reference (21). We abstracted residential census tract median annual household income at the time of delivery from 2000 US Census data.

Statistical analyses

We used linear regression to evaluate associations of air pollution exposures with anthropometric and cardiometabolic biomarkers in early childhood (BMI *z*-score, waist circumference, sum of skinfold thickness, leptin and adiponectin) and mid-childhood (BMI *z*-score, total fat mass, truncal fat mass, leptin, adiponectin and HOMA-IR). For outcomes available at both time points, we examined each separately to accommodate potential differences in the association between the outcome and each confounder by developmental stage. Blood concentrations of leptin, adiponectin and HOMA-IR were not normally

distributed, so we In-transformed them for analyses. For ease of interpretation, we exponentiated the resulting regression coefficients, which we reported as a percent change.

We considered each exposure (BC, $PM_{2.5}$, traffic density and roadway proximity) at each time period in separate models. To account for the exponential spatial decay of traffic pollution (22), we *a priori* categorized residential proximity to major roadway as >200 m, 100 to <200 m, 50 to <100 m and <50 m, as we have done previously (14). We initially modeled BC, $PM_{2.5}$ and traffic density in quartiles, and because exposure–outcome relationships appeared linear, we reported continuous measures and expressed associations per interquartile range (IQR) increment.

We first fit unadjusted models, followed by full multivariable models for each exposure-outcome relationship. We included covariates potentially associated with air pollution exposure and/or childhood cardiometabolic health: maternal age (continuous), education (with or without college degree) and smoking habits (smoked during pregnancy, formerly smoked never smoked); child age (continuous), sex (dichotomous), and race/ethnicity (White, Black, Asian, Hispanic and others); and census tract median household income (continuous). To account for trends in air pollution and adiposity by season and over time, we also included season (continuous sine and cosine of date) and date (continuous) at the time of health outcome in multivariable models. We did not include personal household income, fetal growth or maternal glucose tolerance because inclusion did not appreciably change results. We substituted maternal for child race/ethnicity in 10% of participants missing data on child race/ethnicity. Ninety-eight percent of participants had complete covariate information for the multivariable models. We found no effect modification by child sex or maternal pre-pregnancy BMI, so we present all results without stratification or inclusion of an interaction term for these variables.

In secondary analyses, we examined associations of BC and PM_{2.5} exposure during other time periods [i.e. first trimester (date of LMP to 93rd day after LMP), second trimester (94th day after LMP to 187th day after LMP), and one week prior to health outcome assessment] with early and mid-childhood cardiometabolic health. To account for potential bias due to cohort attrition, we repeated key analyses of roadway proximity at delivery and cardiometabolic outcomes using inverse probability weighting. In addition, because roadway category sample sizes were small and because we occasionally identified non-monotonic associations, we also performed a penalized spline analysis using R Version 3.0.0 (R Foundation for Statistical Computing, Vienna, Austria) to evaluate potential nonlinearity across the range of roadway proximity. For all other analyses, we used SAS version 9.3 (SAS Institute, Cary NC, USA).

Results

Population characteristics

Mean (SD) maternal age was 32.1(5.2) years; 68% of mothers were college graduates, and 69% were nonsmokers. Sixty-four percent of children were White people. Details on early and mid-childhood cardiometabolic outcomes are presented in Table 1.

Third trimester mean (SD, range) BC concentration was 0.7 μ g/m³(0.2, 0.1–1.6). For context, the annual US urban average ranged 0.2–1.9 μ g/m³ from 2005–2007 (23). Third trimester mean(SD, range) PM_{2.5} concentration was 11.8 μ g/m³(1.6, 7.5–16.8), and the Environmental Protection Agency air quality standard for annual PM_{2.5} exposure was 15 μ g/m³ during 1999–2002. At the time of delivery, mean(SD, range) neighborhood traffic density was 1410 (1846, 0–30.860) vehicles/day × km of road within 100 m of residential address; most mothers (88%) lived >200 m from a major roadway, and 3% lived <50 m. Exposures were moderately correlated (Spearman correlation coefficients 0.10–0.64) (Tables S2 and S3).

Mothers with lower 3rd trimester BC exposure were more likely to be older, educated, nonsmokers and live in a census tract with higher median household income. Their children were more likely to be white, heavier at birth and younger at follow-up visits with lower leptin concentration in early childhood and lower total and truncal fat mass, leptin and HOMA-IR in mid-childhood (Table 1).

Air pollution exposure and early childhood cardiometabolic risk

Children whose mothers lived closest (<50 m vs. \geq 200 m) to a major roadway at the time of delivery had 0.3 kg/m² (95%CI: 0.0, 0.7) higher BMI, 1.7 cm (95%CI: 0.6, 2.8) larger waist circumference, 1.9 mm (95%CI: 0.6, 3.2) larger sum of skinfold thickness and 40.7% (95%CI: 5.2, 88.1) higher leptin concentration in early childhood. Children whose mothers lived intermediate distances from a major roadway at delivery (100–<200 m) also had higher BMI *z*-score and larger waist circumference in early childhood. Residential roadway proximity in early childhood was contemporaneously associated with

increased leptin concentration but not other cardiometabolic outcomes (Table 2).

For each IQR increment in neighborhood traffic density at the time of delivery, early childhood leptin concentration was 5.4% (95%Cl: 1.3, 9.7) higher. Traffic density was not associated with other early childhood cardiometabolic parameters. Prenatal and contemporaneous BC and $PM_{2.5}$ exposure were not associated with cardiometabolic risk in early childhood (Table 2).

Air pollution exposure and mid-childhood cardiometabolic risk

Children whose mothers lived closest (<50 vs. \geq 200m) to a major roadway at the time of delivery had 2.1 kg (95%CI: 0.8, 3.5) greater total fat mass, 0.9 kg (95%CI: 0.4, 1.5) greater truncal fat mass and 78.3% (95%CI: 18.5, 168.3) higher leptin concentration in mid-childhood. Children whose mothers lived intermediate distances from a major roadway at delivery (100–<200 m) had higher BMI *z*-score and higher total and truncal fat mass in mid-childhood. Residential roadway proximity at the time of the mid-childhood follow-up visit was not associated with cardiometabolic outcomes (Table 3).

In contrast to our *a priori* hypothesis, exposure to $PM_{2.5}$ during the year prior to the mid-childhood visit was associated with lower rather than higher BMI *z*-score, total and truncal fat mass, and HOMA-IR [e.g. truncal fat mass was 0.3 kg (95%Cl: -0.5, -0.0) lower for each IQR increment $PM_{2.5}$]. Also, for each IQR increment in neighborhood traffic density at the time of delivery, mid-childhood HOMA-IR was 5.7% (95%Cl: -10.1, -1.1) lower. Other air pollution exposure metrics were not associated with mid-childhood outcomes (Table 3).

Secondary analyses

When we considered associations of BC and $PM_{2.5}$ exposure during first and second trimesters and one week prior to the health outcome assessment, for each IQR increment in $PM_{2.5}$ exposure during the first trimester, adiponectin in early childhood was 5.8% lower (95%CI: -10.5, -1.0). Contrary to our *a priori* hypothesis, for each IQR increment in BC exposure during the week prior, HOMA-IR was 17.1% lower (95%CI: -27.6, -5.2) in mid-childhood, not higher. Other exposure-outcome relationships were null (data not shown).

In analyses with (versus without) inverse probability weighting, roadway proximity at delivery had stronger associations with early childhood outcomes and

| Table 1 | Characteristics | of study particip | oants overall and | d by third trimeste | r black carbon | (BC) expos | sure |
|---------|-----------------|-------------------|-------------------|---------------------|----------------|------------|------|
|---------|-----------------|-------------------|-------------------|---------------------|----------------|------------|------|

| | | Quartiles | s of third trimes | ter BC | |
|--|----------------------|----------------------------|-------------------|-------------------|-----------------------------|
| 3 rd trimester BC (μg/m ³), Mean (SD) | Total 0.69 (0.23) | Q1 (lowest) 0.40 (0.09) | Q2 0.60 (0.05) | Q3 0.76 (0.05) | Q4 (highest) 1.00 (0.14) |
| | Mean (SD) or % | | Mean (S | SD) or % | |
| Maternal characteristics | | | | | |
| Age at enrollment (years) | 32.1 (5.2) | 33.0 (4.3) | 32.7 (5.1) | 31.5 (5.6) | 31.2 (5.8) |
| Prepregnancy BMI (kg/m ²) | 24.8 (5.3) | 24.6 (5.2) | 24.8 (5.6) | 24.6 (5.0) | 25.3 (5.6) |
| College graduate (%) | 68 | 78 | 71 | 66 | 58 |
| Smoking habits (%) | | | | | |
| Never | 69 | 67 | 68 | 70 | 71 |
| Former | 20 | 24 | 21 | 18 | 18 |
| During pregnancy | 11 | 10 | 11 | 12 | 11 |
| Glucose tolerance (%) | | | | | |
| Normal | 83 | 82 | 82 | 85 | 85 |
| Failed GCT, normal OGTT | 9 | 11 | 10 | 6 | 7 |
| IGT | 3 | 3 | 3 | 3 | 4 |
| GDM | 5 | 5 | 5 | 6 | 4 |
| Neighborhood characteristics | | | | | |
| Median household income in | 57 763 | 70993 | 60 396 | 53 505 | 45 508 |
| census tract (\$)* | (21 656) | (20 006) | (20 4 1 1) | (20740) | (17 035) |
| Child characteristics in infancy | | | | | |
| Gestational age (weeks) | 39.5 (1.8) | 39.5 (1.7) | 39.4 (1.8) | 39.6 (1.6) | 39.5 (1.8) |
| Birth weight-for-gestational age z-score | 0.20 (0.97) | 0.32 (1.02) | 0.20 (0.93) | 0.19 (0.96) | 0.06 (0.94) |
| Sex (%) | 49 | 47 | 50 | 49 | 49 |
| Race/ethnicity (%) [†] | | | | | |
| White | 64 | 84 | 67 | 56 | 47 |
| Black | 17 | 5 | 14 | 24 | 25 |
| Hispanic | 6 | 1 | 6 | 5 | 11 |
| Asian | 5 | 3 | 5 | 6 | 4 |
| Other | 9 | 7 | 8 | 9 | 13 |
| Early childhood characteristics | | | | | |
| Age at early childhood visit | 3.3 (0.4) | 3.3 (0.3) | 3.3 (0.3) | 3.3 (0.4) | 3.3 (0.5) |
| BMI z-score | 0.5 (1.0) | 0.4 (1.0) | 0.5 (1.0) | 0.4 (1.1) | 0.5 (1.0) |
| Waist circumference (cm) | 51.4 (3.7) | 51.4 (3.5) | 51.4 (3.5) | 51.2 (3.8) | 51.4 (4.0) |
| Sum of skinfolds (mm) | 16.7 (4.3) | 16.7 (4.2) | 17.1 (4.2) | 16.6 (4.7) | 16.5 (4.2) |
| Leptin (ng/mL) | 2.0 (2.0) | 1.9 (2.1) | 1.8 (1.7) | 2.1 (2.3) | 2.0 (1.8) |
| Adiponectin (µg/mL) | 22.3 (5.6) | 22.8 (5.2) | 22.1 (5.6) | 22.4 (5.4) | 22.0 (5.9) |
| Mid-childhood characteristics | | | | | |
| Age at mid-childhood visit | 8.0 (0.9) | 7.8 (0.7) | 7.9 (0.9) | 8.0 (0.8) | 8.1 (1.0) |
| BMI z-score | 0.4 (1.0) | 0.4 (1.0) | 0.4 (1.0) | 0.3 (1.1) | 0.5 (1.0) |
| Total fat mass (kg) | 7.5 (3.9) | 7.0 (3.2) | 7.4 (3.9) | 7.4 (4.1) | 8.0 (4.2) |
| Truncal fat mass (kg) | 2.5 (1.7) | 2.3 (1.4) | 2.5 (1.8) | 2.5 (1.7) | 2.7 (1.8) |
| Leptin (ng/mL) | 6.1 (7.5) | 5.6 (6.6) | 5.4 (6.5) | 5.8 (7.5) | 7.6 (8.7) |
| Adiponectin (µg/mL) | 15.6 (8.8) | 15.8 (8.9) | 14.4 (8.8) | 16.2 (8.5) | 15.5 (8.9) |
| HOMA-IR | 1.9 (1.8) | 1.6 (1.4) | 1.8 (1.4) | 1.8 (1.6) | 2.3 (2.5) |

*Based on address at the time of delivery. [†]Maternal race/ethnicity is substituted in 10% of children whose race/ethnicity is missing. BMI, body mass index; GCT, glucose tolerance test; GDM, gestational diabetes mellitus; HOMA-IR, homeostasis model assessment for insulin resistance; IGT, impaired glucose tolerance; OGTT, oral glucose tolerance test; SD, standard deviation.

| Table 2 | Covariate-adjusted* associations of traffic-related air pollution in pregnancy and in early childhood with adiposity and cardiometabolic health in early childhooc |
|------------|--|
| (median: S | : 3.3 years of age) |

| | BMI z-score (z-units) | Waist circumference(cm) | Sum of skinfold thickness(mm) | Leptin (% change) | Adiponectin (% change) |
|---|--|---|---|---|---|
| Black carbon (BC) exposure (µg/m ³) Third trimester | -0.0 (-0.1, 0.1) | 0.0 (-0.3, 0.4) | -0.0 (-0.4. 0.4) | 7.4 (-1.7, 17.3) | -2.1 (-6.2. 2.3) |
| Year prior to early childhood visit | -0.0 (-0.1, 0.1) | -0.1 (-0.4, 0.3) | -0.1 (-0.4, 0.3) | 3.0 (-4.9, 11.6) | -0.6 (-4.5, 3.4) |
| Fine particulate (PM $_{2.5}$) exposure (μ g/m 3) | | | | | |
| Third trimester | 0.0 (-0.1, 0.1) | 0.2 (-0.1, 0.5) | 0.3 (-0.1, 0.6) | -5.4 (-12.4, 2.1) | -3.1 (-6.7, 0.6) |
| Year prior to early childhood visit | -0.0 (-0.1, 0.1) | -0.2 (-0.6, 0.2) | 0.3 (-0.2, 0.8) | 8.8 (-1.4, 20.0) | -0.9 (-5.6, 4.0) |
| Near-residence traffic density | | | | | |
| Birth address | 0.0 (-0.0, 0.1) | 0.0 (-0.1, 0.2) | 0.0 (-0.2, 0.2) | 5.4 (1.3, 9.7) | 0.0 (-1.9, 2.0) |
| Early childhood address | 0.0 (-0.0, 0.1) | -0.0 (-0.2, 0.1) | 0.0 (-0.2, 0.2) | 2.0 (-1.5, 5.6) | 0.8 (-0.9, 2.5) |
| Proximity to major roadway, birth address | | | | | |
| <50m | 0.3 (0.0, 0.7) | 1.7 (0.6, 2.8) | 1.9 (0.6, 3.2) | 40.7 (5.2, 88.1) | 1.1 (-12.3, 16.5) |
| 50-<100m | -0.0 (-0.4,0.3) | 0.0 (-1.2, 1.3) | 0.1 (-1.3, 1.5) | 21.0 (-8.6, 60.2) | -2.0 (-14.6, 12.4) |
| 100-<200 m | 0.4 (0.1, 0.6) | 1.0 (0.1, 1.8) | 0.7 (-0.3, 1.7) | 17.4 (-7.7, 49.2) | -13.1 (-22.7,-2.3) |
| ≥200 m | Reference | Reference | Reference | Reference | Reference |
| Proximity to major roadway, early childhood address | | | | | |
| <50m | 0.1 (-0.2, 0.5) | 0.8 (-0.5, 2.1) | 1.1 (-0.4, 2.7) | 41.7 (3.0, 94.9) | 9.5 (-6.3, 28.0) |
| 50-<100m | -0.0 (-0.4, 0.3) | -0.1 (-1.4, 1.2) | -0.8 (-2.3, 0.6) | -8.0 (-33.2, 26.6) | -9.5 (-22.5, 5.8) |
| 100- < 200 m | 0.1 (-0.2, 0.3) | -0.1 (-1.0, 0.7) | 0.1 (-0.9, 1.1) | 0.4 (-19.0, 24.5) | 2.0 (-8.2, 13.3) |
| ≥200 m | Reference | Reference | Reference | Reference | Reference |
| For black carbon (BC), fine particulate matter (PM _{2.8}) and traffic densit range = 0.33 μg/m ³ for third trimester BC, 0.22 μg/m ³ for BC during th 1454 km * vehicles/day for traffic density at birth and 1247 km * vehicles imate category of roadway proximity versus ≥200 m. Estimates with 95 *Model adjusted for characteristics of child (age, sex and race/ethnicity). | ty exposures, estimates are re year prior to the early chil suday for traffic density at e 5% confidence intervals that , mother (age, education, sm | mean difference (95%, sonfic dhood visit, 2.20 µg/m ³ for th any childhood address). For r t do not cross the null are bc loking during pregnancy) and | lance intervals) in outcome 1 nird trimester PM _{2.5} , 1.33 µg oadway proximity, estimates Ided. neighborhood (census tract | or gach interquartile range inci /m ³ for PM _{2.5} during the year i are mean difference (95% cor median income), as well as sea | rement in exposure (interquartile prior to the early childhood visit, ifidence intervals) for each prox- son and date of health outcome. |

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Table 3 Covariate-adjusted* associations of traffic-related air pollution in pregnancy and in mid-childhood with adiposity and cardiometabolic health in mid-childhood (median: 7.7 years of age)

| | | | ŀ | - | - | |
|---|--|--|--|--|---|---|
| | BIMI z-score (z-units) | lotal tat mass(kg) | Iruncal tat mass(kg) | Leptin (% change) | Adiponectin (% change) | HOMA-IK (% change) |
| Black carbon (BC) exposure (µg/m ³) | | | | | | |
| I nirg trimester | -0.0 (-0.1, 0.1) | -0.Z (-0.9, 0.Z) | -0.1 (-0.2, 0.1) | 4.0 (-0.0, 15.7) | 2.4 (-4.9, 10.3) | 1.1 (-8.8, 12.1) |
| Year prior to mid-childhood visit | -0.1 (-0.2, 0.0) | 0.0 (-0.3, 0.4) | 0.0 (-0.1, 0.2) | 3.5 (-6.8, 15.0) | 1.3 (-5.9, 9.0) | 3.9 (–5.9, 14.6) |
| Fine particulate ($PM_{2.5}$) exposure (μ g/m ³) | | | | | | |
| Third trimester | -0.0 (-0.1, 0.0) | -0.2 (-0.6, 0.1) | -0.1 (-0.2, 0.1) | -5.1 (-14.7, 5.6) | -5.0 (-11.7, 2.3) | 2.8 (-6.6, 13.0) |
| Year prior to mid-childhood visit | -0.2 (-0.4,-0.1) |) -0.6 (-1.2,-0.1 |) -0.3 (-0.5,-0.0 | -12.1 (-24.9, 2.8) | -2.8 (-13.0, 8.6) | -17.8 (-29.2,-4.7) |
| Near-residence traffic density | | | | | | |
| Birth address | 0.0 (-0.0, 0.1) | 0.0 (-0.2, 0.2) | -0.0 (-0.1, 0.1) | 4.0 (-1.3, 9.6) | 0.6 (-3.0, 4.3) | -5.7 (-10.1,-1.1) |
| Mid-childhood address | -0.0 (-0.1, 0.0) | 0.0 (-0.1, 0.2) | 0.0 (-0.1, 0.1) | 1.9 (-2.6, 6.6) | -0.9 (-4.1, 2.3) | 0.1 (-4.2, 4.5) |
| Proximity to major roadway, birth address | | | | | | |
| <50m | 0.1 (-0.2, 0.5) | 2.1 (0.8, 3.5) | 0.9 (0.4, 1.5) | 78.3 (18.5, 168.3) | -13.2 (-34.7, 15.4) | -0.2 (-33.6, 49.8) |
| 50-<100m | -0.0 (-0.4, 0.4) | -0.5 (-2.0, 1.0) | -0.3 (-0.9, 0.4) | -4.9 (-36.0, 41.3) | -1.2 (-25.0, 30.2) | -32.4 (-53.6,-1.4) |
| 100-<200 m | 0.3 (0.0, 0.5) | 1.1 (0.1, 2.0) | 0.4 (-0.0, 0.8) | 1.2 (-21.5, 30.5) | 1.0 (-15.4, 20.5) | -6.3 (-27.2, 20.7) |
| ≥200 m | Reference | Reference | Reference | Reference | Reference | Reference |
| Proximity to major roadway, mid-childhood | σ | | | | | |
| address | | | | | | |
| <50m | -0.0 (-0.4, 0.4) | 0.1 (-1.4, 1.6) | 0.0 (-0.6, 0.7) | 46.6 (-5.3, 127.1) | -13.5 (-36.1, 17.2) | -1.8 (-36.6, 52.3) |
| 50-<100m | -0.1 (-0.5, 0.3) | -1.0 (-2.5, 0.5) | -0.4 (-1.1, 0.2) | -19.1 (-45.5, 20.0) | 0.4 (-23.6, 32.0) | -13.1 (-41.3, 28.6) |
| 100-<200 m | 0.1 (-0.2, 0.3) | 0.4 (-0.6, 1.5) | 0.2 (-0.3, 0.6) | 8.6 (-18.0, 43.9) | 5.6 (-13.1, 28.3) | -5.8 (-29.4, 25.8) |
| ≥200 m | Reference | Reference | Reference | Reference | Reference | Reference |
| For black carbon (BC), fine particulate matter (PM _{2.6}), range = 0.33 μg/m ³ for third trimester BC, 0.20 μg/m ³ 1454 km* vehicles/day for traffic density at birth and 11 of roadway proximity versus ≥200 m. Estimates with 95 "Worde adjusted for characteristics of child (age, sex anc BMI, body mass index; HOMA-IR, homeostasis model | and traffic density exposuru for BC during the year priv 86 km * vehicles/day for m 5% confidence intervals th 1race/ethnicity, mother (ag assessment for insulin res | es, estimates are mean c or to the mid-childhood v id-childhood traffic densit at do not cross the null a ge, education and smokin istance. | lifference (95%, confidence isit, 2.20 µg/m ³ for third t y). For roadway proximity, e bolded. g during pregnancy), neigh | i intervals) in outcome for ea imester PM _{2.5} , 1.66 μg/m ³ f estimates are mean differenc iborhood (census tract media | ch interquartile range incren or PM _{2.5} during the year pri e (95% confidence intervals in income), as well as seasor | tent in exposure (interquartile or to the mid-childhood visit,) for each proximate category and date of health outcome. |

similar associations with mid-childhood outcomes (Table S4). In the penalized spline model, roadway proximity at delivery and mid-childhood truncal fat mass showed a stronger association with closer roadway proximity (Figure S2) with similar results for total fat mass (data not shown).

Discussion

In our analysis of a large prospective cohort, infants whose mothers lived close to a major roadway at the time of delivery had greater adiposity in early and mid-childhood. However, prenatal and early life exposure to air pollutants and traffic density were not consistently associated with adiposity or insulin resistance.

Our findings suggest that features of roadway proximity distinct from air pollution (or from the pollutants we measured) may contribute to later cardiometabolic risk. For example, sleep disruption from roadway noise (24) and light (25), as well as reduced neighborhood walkability (26) are roadway characteristics independently associated with adiposity and dysmetabolism. Alternatively, ultrafine particles (UFPs), which were not measured in our cohort, could have driven the association between residential roadway proximity and cardiometabolic health. UFPs, which have a diameter $< 0.1 \,\mu\text{m}$ and are primarily emitted from vehicle exhaust, have been increasingly implicated in health effects, particularly in urban areas. UFPs increase with vehicle speed and decrease with idling, features common to traffic on major roadways, and they aggregate quickly to form larger particles, so concentrations fall rapidly with distance from roadway (27). Our findings may be impacted by unmeasured confounding by socioeconomic status (SES), although roadway proximity was not as tightly correlated as air pollution with the SES factors measured in our cohort (data not shown). The findings may also reflect random chance, particularly given the small sample sizes in the roadway categories. However, an inverse association between roadway proximity and childhood adiposity in spline models suggests against this possibility.

Our results are consistent with one prior study in which residential roadway proximity (<50 m) but not PM_{2.5} predicted incident type 2 diabetes mellitus in adult women (28). Also, in the Project Viva cohort, impaired neurocognition in childhood was similarly associated with roadway proximity (<50 m) at the time of delivery but not at the time of cognitive testing (29), raising the possibility of an *in utero* programming

effect. Nevertheless, our findings require replication in other populations of pregnant women and children.

We did not observe consistent associations of BC, PM_{2.5} or traffic density exposures with childhood cardiometabolic parameters, although there were a few sporadic associations that did not follow a clear pattern. For example, neighborhood traffic density at the time of delivery and contemporaneous roadway proximity were associated with higher leptin in early childhood. Also, contrary to our a priori hypothesis, PM_{2.5} exposure during the year prior was associated with lower rather than higher BMI z-score, total and truncal fat mass, and HOMA-IR. Although the PM_{2.5} model estimated 10 km × 10 km exposures which could limit local contrast and bias results toward the null, it is unlikely to have led to negative associations. The negative associations are somewhat consistent with one rodent study in which overweight but not normal weight mice exposed to PM_{2.5} in early childhood had non-significantly lower HOMA-IR and body weight (5), and this is in line with an above average BMI z-score of children in our cohort. However, this finding has not been replicated in other animal or human studies, and the biological basis is not clear.

The bulk of the existing rodent and human literature supports an association between air pollutants and cardiometabolic health. In rodents, air pollution exposure led to visceral adiposity and insulin resistance with effects mediated through induction of oxidative stress and systemic inflammation (4,5), as well as neuroinflammation with consequent brain remodeling and altered satiety signals (30). In cohort studies of prenatal exposure, polycyclic aromatic hydrocarbon (a combustion byproduct of fossil fuel and biomass burning) (8) has been associated with early childhood obesity, and Project Viva infants born to mothers living in neighborhoods with higher traffic density had more rapid weight gain and greater risk of weightfor-length >95th percentile by 6 months of age (14). In elementary (9) and teenage (6) cohorts in Southern California, residential traffic pollution at enrollment was associated with BMI over 4-8 years of followup, and elementary school children in China were more likely to be obese if school/residential air pollution (PM_{10} , SO_2 and O_3) was higher during the 2 years preceding the weight measurement (7). Additionally, population-based studies in Iran (10,12) and Germany (11) have demonstrated an association between air pollution exposure and insulin resistance in childhood.

Limitations of Project Viva that may have prevented us from observing a persistent association between early life air pollution exposure and cardiometabolic outcomes include generally low air pollution exposures in the Boston area and a cohort of primarily white children of moderately high SES at relatively low risk for adverse cardiometabolic health. Strengths included use of a large, prospective cohort with multiple potential confounding variables, several measures of air pollution exposure with daily spatiotemporal resolution and evaluation of both anthropometric and serum markers of dysmetabolism at two time points in childhood.

In conclusion, infants whose mothers lived close to a major roadway at the time of delivery were at risk for adverse cardiometabolic parameters in early and mid-childhood. However, we found no evidence of a persistent effect of prenatal or early life BC or $PM_{2.5}$ exposures on childhood cardiometabolic profile in a population with relatively high SES exposed to modest levels of air pollution.

Conflicts of Interest Statement

The authors declare no conflict of interest.

Author contributions

A.F.F. conceived this analysis and drafted the manuscript. A.F.F., H.L.-G., W.P. and S.L.R.-S. performed the analysis. All authors critically reviewed the manuscript.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Table S1. Characteristics of participants in the fullcohort and for the subset included in analyses.

Table S2. Relationship between third trimester blackcarbon (BC) exposure and other measures of air pol-lution.

Table S3. Relationship between third trimester blackcarbon (BC) exposure and distance to roadway.

Table S4. Covariate-adjusted associations of roadway proximity at delivery and adiposity and cardiometabolic health in early childhood (median: 3.3 years of age) and mid-childhood (median: 7.7 years of age) without versus with inverse probability weighting (IPW) to account for loss to follow-up.

Figure S1. Project Viva cohort sample size and participation. 1418 participants had data available for at least one exposure and one outcome studied.

Figure S2. Covariate-adjusted mean difference (95% confidence intervals) in truncal fat mass in mid-childhood (median: 7.7 years of age) associated with residential major roadway proximity at time of delivery using a penalized spline model. (A) shows the full range of roadway distances and (B) is cropped to show only distances < 1000 m.