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Association of Estimated Long-term Exposure to Air Pollution and Traffic Proximity With a Marker for Coronary Atherosclerosis in a Nationwide Study in China

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Abstract

IMPORTANCE Epidemiologic evidence of the mechanisms of the association between long-term exposure to air pollution and coronary heart disease (CHD) is limited and relies heavily on studies performed in Europe and the United States, where air pollution levels are relatively low. In particular, the association between air pollution and CHD in patients with underlying risks for CHD is understudied.

OBJECTIVE To determine whether air pollution and proximity to traffic are associated with the coronary artery calcium (CAC) score, a key atherosclerotic marker.

DESIGN, SETTING, AND PARTICIPANTS In this prospective, population-based cross-sectional study in a large-scale setting in China, 8867 consecutive patients aged 25 to 92 years with suspected CHD were recruited between November 17, 2015, and September 13, 2017. Participants were excluded if they had previous myocardial infarction, stenting, or coronary artery bypass grafting or incomplete risk factors and exposure data. Each participant underwent assessment of CAC and CHD risk factors at baseline. Data were analyzed from December 2017 to November 2018.

EXPOSURES Annual means of fine particulate matter with aerodynamic diameter less than 2.5 μm (PM$_{2.5}$), nitrogen dioxide (NO$_2$), and ozone (O$_3$) were estimated at the participants’ residences using a validated geostatistical prediction model. Exposure to a nearby roadway was also estimated.

MAIN OUTCOMES AND MEASURES Computed tomography measurement of CAC score.

RESULTS The mean (SD) age of the 8867 participants was 56.9 (10.4) years; 4378 (53.6%) were men. Annual mean (SD) PM$_{2.5}$, NO$_2$, and O$_3$ measurements were 70.1 (20.0), 41.4 (14.7), and 93.9 (10.5) μg/m$^3$, respectively. The mean (SD) CAC score was 91.4 (322.2) Agatston units. Exposure to PM$_{2.5}$ and NO$_2$, adjusting for CHD risk factors and multiple pollutants, were independently associated with increases in CAC scores of 27.2% (95% CI, 10.8% to 46.1%) per 30 μg/m$^3$ PM$_{2.5}$ and 24.5% (95% CI, 3.6% to 49.7%) per 20 μg/m$^3$ NO$_2$. For PM$_{2.5}$, odds of both detectable CAC (Agatston score $>$0; odds ratio, 1.28; 95% CI, 1.13 to 1.45) and severe CAC (Agatston score $>$400; odds ratio, 1.59; 95% CI, 1.20 to 2.12) were increased. Associations of CAC with PM$_{2.5}$ and NO$_2$ were greater among male participants (PM$_{2.5}$: 42.2%; 95% CI, 24.3% to 62.7%; NO$_2$: 45.7%; 95% CI, 25.3% to 69.5%) and elderly participants (PM$_{2.5}$: 50.1%; 95% CI, 28.8% to 75.0%; NO$_2$: 55.5%; 95% CI, 31.8% to 83.6%) and those with diabetes (PM$_{2.5}$: 62.2%; 95% CI, 30.9% to 101.0%; NO$_2$: 31.2%; 95% CI, 13.9% to 51.0%). Independent association with CAC score was 9.0% (95% CI, −1.4% to 20.4%) for O$_3$ per 15 μg/m$^3$ and 2.4% (95% CI, −0.6% to 5.4%) for distance near roadway per 50% decrease.

Key Points

Question Are long-term exposure to ambient air pollution and proximity to traffic associated with subclinical atherosclerosis?

Findings In this population-based cross-sectional study of 8867 Chinese participants, long-term exposure to ambient nitrogen dioxide and fine particulate matter with aerodynamic diameter less than 2.5 μm was independently associated with a higher coronary artery calcium score, a key atherosclerotic marker. Associations with ozone and proximity to traffic were less consistent.

Meaning Long-term exposure to ambient air pollution may be an important risk factor for coronary atherosclerosis.

Supplemental content

Author affiliations and article information are listed at the end of this article.

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CONCLUSIONS AND RELEVANCE  In this large Chinese study, long-term exposures to PM$_{2.5}$ and NO$_2$ were independently associated with severity of CAC. This finding may provide support for the pathophysiological role of coronary atherosclerosis through which air pollution exposure may be associated with CHD.

Introduction

Ambient air pollution is a major contributor to the global burden of disease, accounting for an estimated 4.2 million deaths in 2015. 1 Although the effect of air pollution on public health is potentially very large, the current air pollution guidelines used worldwide have relied principally on findings from studies conducted in Europe and North America where air pollution levels are relatively low compared with other areas experiencing more recent rapid industrialization. 2 Information on associations between exposure and disease from regions with exceptionally high ambient air pollution, such as China, would be useful to produce more accurate estimates of air pollution-attributable disease burden, which in turn can be used to prioritize public health responses, especially in developing countries.

Cardiovascular disease is the leading cause of death worldwide, with available evidence showing that it is associated with long-term exposure to air pollution and proximity to traffic. 3 A likely pathway underlying these associations involves the initiation or acceleration of atherosclerosis. 3 Atherosclerosis is a lifelong process; therefore, the effects of air pollution exposure on atherosclerosis are likely to be long term. If an association between air pollution and subclinical atherosclerosis were established, it could provide an opportunity to intervene before disease is manifested clinically by way of community-level efforts to control pollution exposures. Common approaches to detecting subclinical atherosclerosis include noninvasive measurement of coronary artery calcium (CAC) or carotid intima-media thickness. 4 Compared with carotid intima-media thickness, CAC reflects a more advanced stage of atherosclerosis involving calcification of plaque and is a better predictor of cardiovascular disease. 3 Associations between air pollution and carotid intima-media thickness have been reported in some previous studies. 5-8 Evidence regarding the association between air pollution and coronary atherosclerosis, determined using CAC scoring, is still very limited. 8,9

China is the most populated country in the world and has a large estimated disease burden associated with air pollution. 1 Cohort studies of air pollution and cardiovascular health are still scarce and have been largely limited to the investigation of effects on mortality. 10 Given the increasing public concern regarding air pollution, the Chinese government has recently implemented an unprecedented effort to characterize major air pollutant exposures with more than 1400 widespread monitoring stations established nationwide. 11 Here we use this valuable resource by applying advanced exposure estimation methods for health effect investigation. We hypothesize that long-term exposure to air pollution or proximity to traffic is associated with increased risk of CAC in a well-characterized cohort of Chinese adults.

Methods

Study Population

This cross-sectional study used baseline data collected from a prospective cohort (the CREATION cohort) of 8867 consecutive patients aged 25 to 92 years at Fuwai Hospital in Beijing, China (eMethods in the Supplement). The participants were suspected of having coronary heart disease (CHD) and for this reason underwent cardiac computed tomography to evaluate the presence and amount of CAC between November 17, 2015, and September 13, 2017. Cardiac imaging was ordered...
by primary cardiologists. The participants had relatively low pretest probability of CHD (mean [SD], 27.3% [15.1%]); 2587 patients had no symptoms, 5721 had nonanginal chest pain, 449 had atypical angina, and 110 had typical angina. Participants were excluded if they had previous myocardial infarction, stenting, or coronary artery bypass grafting or incomplete risk factors and exposure data. More details regarding the study cohort are available in Figure 1 and eMethods in the Supplement. All participants provided written informed consent, and the study was approved by the institutional review board of the Chinese Academy of Medical Sciences Fuwai Hospital. This report follows the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) reporting guideline.

CAC Measurement
Details of the measurement of CAC have been published elsewhere. In brief, all patients underwent imaging with a second-generation dual-source computed tomography system (Somatom Definition Flash; Siemens Healthcare) with a standardized scanning protocol, using 128 × 3-mm section collimation, 75-millisecond rotation time, and 120-kV tube voltage. Coronary calcium lesions were defined as having an attenuation threshold greater than or equal to 130 Hounsfield units (HU) and an area greater than or equal to 1 mm². Images were analyzed with the system’s integrated software (Syngo.Via; Siemens Healthcare). The total calcium burden in the coronary arteries was quantified by the scoring algorithm proposed by Agatston et al. The products of the area of each calcified plaque and peak attenuation, defined as 1 (130-199 HU), 2 (200-299 HU), 3 (300-399 HU), and 4 (>400 HU), were summed for the left main coronary artery, left anterior descending coronary artery, left circumflex coronary artery, and right coronary artery to determine the total CAC score.

Exposure Prediction
The method for estimating long-term outdoor air pollution concentrations for each study participant has been described in detail elsewhere. In brief, we developed a hierarchical land-use regression modeling approach for fine particulate matter less than 2.5 μm in aerodynamic diameter (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>) on the basis of annual mean daily monitoring data (2014-2015) from 1419 regulatory monitors nationwide. The models incorporated a wide diversity of geographic features (eg, traffic network, point of interest, population density, and land use), satellite-derived PM<sub>2.5</sub> or NO<sub>2</sub> observations at ground level calibrated by chemical transport model estimates, and meteorological data, with regression residuals smoothed by universal kriging. The performance of

Figure 1. Study Design and Population

Flowchart shows study cohort enrollment criteria. CT indicates computed tomography.
these models ranged from good to excellent, as assessed by the cross-validation (out-of-sample validation; $R^2 = 0.73-0.89$). The model resolution was finer than $1 \times 1$ km$^2$ for the entire country of China. Residence-specific annual mean PM$_{2.5}$ and NO$_2$ concentrations were estimated for 2015 and were used as the primary exposure measures. For O$_3$, we estimated annual mean concentrations by interpolation of O$_3$ observations from the proximal O$_3$ monitoring sites using ordinary kriging. As an alternative assessment of traffic-related associations, we assessed residential proximity to traffic, estimated by the distance to any of the nearest roadways on a natural logarithm scale,$^{15}$ as a secondary exposure variable. This variable is a proxy of exposure to traffic mixtures, including tailpipe and nontailpipe emissions and roadway noise.

In the sensitivity analysis, we tested different exposure time windows to our outcome. For PM$_{2.5}$, we estimated cumulative exposures back for 3-, 4-, 5-, 6-, and 10-year periods before 2015. This analysis took advantage of our single-year fine-scale land-use regression predictions together with the 10-year temporal trend (2004-2014) generated from national satellite-derived data$^{16}$ (eMethods in the Supplement). For NO$_2$ and O$_3$, a monitoring period averaged between 2014 and 2015 was examined.

**Statistical Analysis**

We fit multiple linear regression models in analyzing associations of long-term air pollution exposure (PM$_{2.5}$, NO$_2$, and O$_3$) and proximity to traffic with the CAC score in separate, single-pollutant models. The CAC score was modeled on a continuous scale using a logarithm (CAC score + 1) transformation to account for skewness of the distribution.

We developed statistical models in stages, by incrementally increasing control for a large set of covariates according to a priori knowledge from previous studies.$^{8,17}$ We also used a change-in-estimate criterion by including variables if they changed the effect estimates for the unadjusted association between air pollutants and CAC by more than 10%. The primary model included the following individual-level variables: age, sex, body mass index, smoking (status, duration, and intensity), alcohol consumption, education, and physical activity. To address potential behavioral confounders that differed by geography, we also added area-level variables, including urbanization ($\geq2500$ population per $1 \times 1$-km$^2$ grid), study region (ie, north, southeast, and southwest), Beijing residence (yes or no), and categories of residence distances to Fuwai Hospital (<30, 30-150, 151-300, 301-500, and >500 km) in the primary model. Potential effect modification was evaluated for specific participant characteristics (age and sex), disease risk factors (body mass index, smoking, diabetes, and statin use), and geography (region and urbanization), which have been suggested to act as effect modifiers in prior studies.$^{7,10,18}$ Because cardiovascular disease mortality risk increases in women after menopause,$^{3}$ in an additional analysis, we restricted our analysis to postmenopausal women. To assess exposure-response associations, we refit the models using natural regression splines. To explore nonlinearity associations, we categorized exposure estimation by quartiles.

In sensitivity analyses, we used the same model but restricted analysis to participants with predicted annual exposure concentrations lower than the national standards in China (<35 and <40 $\mu$g/m$^3$ for PM$_{2.5}$ and NO$_2$, respectively) or median exposure for O$_3$ (<100 $\mu$g/m$^3$). Second, we added to the model high-sensitivity C-reactive protein, total cholesterol, high-density lipoprotein cholesterol, low-density lipoprotein cholesterol, and triglyceride levels in the subset of 6538 participants from whom blood samples were collected. Third, the use of lipid-lowering and antihypertensive medications was added to the models. Fourth, individual-level data that included detailed information on the built environment, socioeconomic status (eg, income and employment status), and residential moving history were available for a subset of 1850 participants who completed an additional environmental survey. The sensitivity of our primary model estimates to the addition of each or a group of those covariates was also assessed. Fifth, to account for possible spatial clustering of the outcome, we used a mixed-effects model with a random effect for the 6-digit postal code (n = 917). Sixth, to assess the robustness of effect estimates in the primary model, we estimated the associations in a subset after excluding the most influential participants (Cook
distance > 4/total sample size). Seventh, as noted already, we examined the effects of replacing our
primary exposure variables with exposures of longer durations (2-year means for NO₂ and O₃, and 2-, 3-, 4-, 5-, 6-, and 10-year means for PM₂.₅). Eighth, multipollutant models were used to assess the
independent effects of all the air pollutants and distance variable.

To compare with similar studies, we also assessed the association using 2 binary
categorizations of CAC—detectable (CAC score >0 Agatston units) or severe (CAC score >400
Agatston units)—in logistic regression models. The presence and severity of CAC represent
underlying coronary atherosclerosis in incipient and advanced stages. R statistical software version
3.4.2 (R Foundation) was used for developing the pollution exposure models, and SAS statistical
software version 9.4 (SAS Institute) was used for the health model analyses. The statistical
significance threshold was .05, and all tests were 2-sided.

Results

Study Participants and Environmental Exposures

Of the 8867 participants with a CAC score, 8168 had estimated outdoor residential pollutant
concentrations for the year of the baseline examination. The participants’ mean (SD) age was 56.9
(10.4) years; 4378 (53.6%) were men (Table). More than 30% were smokers, 14.5% had at least a
college education, 49.0% had diagnosed hypertension, and 5.0% had moved residence in the last
5 years.

Annual mean PM₂.₅, NO₂, and O₃ measurements were 70.1 (20.0), 41.4 (14.7), and 93.9 (10.5)
μg/m³, respectively. Annual mean air pollutant concentrations varied substantially among the study
participants (eTable 1 in the Supplement), with all pollutants being highest for participants in the
Hebei province and lowest for those in the south (Figure 2; eFigure 1 in the Supplement). The
participants were exposed to higher concentrations of PM₂.₅ and NO₂ in urban areas than in rural
areas. Predictions for PM₂.₅ were positively correlated with NO₂ and O₃ (Spearman correlation
coefficients, 0.72 and 0.53, respectively) and negatively correlated with proximity to a roadway
(Spearman correlation coefficient, −0.10). Of the participants, 7728 (95.0%) had estimated
exposures greater than 35 μg/m³ for PM₂.₅ and 4551 (56.0%) had estimated exposures greater than
40 μg/m³ for NO₂.

Air Pollution, Proximity to Traffic, and CAC

The mean (SD) CAC score was 91.4 (322.2) Agatston units. Long-term exposure to PM₂.₅, NO₂, and
O₃ as well as distance to the nearest roadway, were all associated with higher CAC score in the crude,
moderately adjusted, and fully adjusted single-pollutant models (fully adjusted single-pollutant
model, PM₂.₅ per 30 μg/m³: 29.6%; 95% CI, 15.7%-45.2%; NO₂ per 20 μg/m³: 33.2%; 95% CI, 16.4%-
52.4%; O₃ per 15 μg/m³: 10.8%; 95% CI, 0.9%-21.8%; distance to roadway per 50% decrease: 3.2%;
95% CI, 0.3%-6.2%) (Figure 3; eTable 2 in the Supplement). Independent association with CAC score
was 9.0% (95% CI, −1.4% to 20.4%) for O₃ per 15 μg/m³ and 2.4% (95% CI, −0.6% to 5.4%) for
distance near roadway per 50% decrease. Associations were not substantially changed by additional
adjustment for medications and blood biomarkers, accounting for neighborhood clustering, use of
alternative exposure durations, and exclusion of data with extreme observations (eTable 2 and
eFigure 2 in the Supplement). The addition of individual home environment variables (eg, environmental tobacco smoking, use of air conditioner, air purifier, cooking stoves, or a ventilation
system) in a subgroup analysis of 1850 participants, the addition of socioeconomic status variables
(eg, income and employment status), or the exclusion from the subgroup analysis of 88 participants
who had moved similarly did not meaningfully change the estimates (eTable 3 in the Supplement).

When we restricted the analyses to the subset of participants with predicted exposure
concentrations below the national standard (or median of exposure distribution), the association
between NO₂ and CAC was statistically significant (NO₂ per 10 μg/m³: 24.0%; 95% CI, 6.4% to
44.5%). The association with O₃ was not significant (O₃ per 10 μg/m³: 7.4%; 95% CI, −0.5% to 16.1%).
Table. Descriptive Statistics of the Study Population at Baseline Classified by High and Low Estimated PM$_{2.5}$ and NO$_2$ Concentrations in 2015 in China

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Entire Cohort (N = 8168)$^a$</th>
<th>PM$_{2.5}$, μg/m$^3$</th>
<th>NO$_2$, μg/m$^3$</th>
<th>P Value$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>≤80 (n = 4250)$^a$</td>
<td>&gt;80 (n = 3918)$^a$</td>
<td>≤42 (n = 4083)$^a$</td>
<td>&gt;42 (n = 4085)$^a$</td>
</tr>
<tr>
<td>Coronary artery calcium present</td>
<td>3676 (45.0)</td>
<td>1828 (43.0)</td>
<td>1848 (47.0)</td>
<td>1748 (42.8)</td>
</tr>
<tr>
<td>Coronary artery calcium score, Agatston units</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>91.4 (322.2)</td>
<td>84.9 (304.5)</td>
<td>98.6 (340.6)</td>
<td>87.9 (325.4)</td>
</tr>
<tr>
<td>Age, mean (SD), y</td>
<td>56.9 (10.4)</td>
<td>56.7 (9.8)</td>
<td>57.1 (11)</td>
<td>56.6 (9.6)</td>
</tr>
<tr>
<td>Male</td>
<td>4378 (53.6)</td>
<td>2163 (50.9)</td>
<td>2215 (56.5)</td>
<td>2086 (51.1)</td>
</tr>
<tr>
<td>Education</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;College</td>
<td>6984 (85.5)</td>
<td>3919 (92.2)</td>
<td>3065 (78.2)</td>
<td>3838 (94)</td>
</tr>
<tr>
<td>College</td>
<td>1045 (12.8)</td>
<td>299 (7.0)</td>
<td>746 (19.1)</td>
<td>233 (5.7)</td>
</tr>
<tr>
<td>&gt;College</td>
<td>139 (1.7)</td>
<td>32 (0.8)</td>
<td>107 (2.7)</td>
<td>12 (0.3)</td>
</tr>
<tr>
<td>Risk factors</td>
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<td></td>
<td></td>
<td></td>
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<tr>
<td>BMI, mean (SD)</td>
<td>25.4 (3.2)</td>
<td>25.3 (3.2)</td>
<td>25.6 (3.3)</td>
<td>25.3 (3.3)</td>
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<tr>
<td>Smoking status</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never and former</td>
<td>5652 (69.2)</td>
<td>2994 (70.5)</td>
<td>2658 (67.9)</td>
<td>2864 (70.2)</td>
</tr>
<tr>
<td>Current</td>
<td>2516 (30.8)</td>
<td>1256 (29.5)</td>
<td>1260 (32.1)</td>
<td>1219 (29.8)</td>
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<tr>
<td>Smoking duration, mean (SD), y</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>≤10</td>
<td>1242 (15.2)</td>
<td>606 (14.2)</td>
<td>636 (16.2)</td>
<td>558 (13.6)</td>
</tr>
<tr>
<td>11-20</td>
<td>4990 (61.1)</td>
<td>2614 (61.6)</td>
<td>2376 (60.7)</td>
<td>2520 (61.8)</td>
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<tr>
<td>&gt;20</td>
<td>1936 (23.7)</td>
<td>1030 (24.2)</td>
<td>906 (23.1)</td>
<td>1005 (24.6)</td>
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<tr>
<td>Alcohol consumption</td>
<td>1764 (21.6)</td>
<td>852 (20.1)</td>
<td>912 (23.3)</td>
<td>820 (20.1)</td>
</tr>
<tr>
<td>Physical activity, times per/wk</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Never</td>
<td>1413 (17.3)</td>
<td>737 (17.3)</td>
<td>676 (17.2)</td>
<td>723 (17.7)</td>
</tr>
<tr>
<td>≤3</td>
<td>3994 (48.9)</td>
<td>2074 (48.8)</td>
<td>1920 (49.0)</td>
<td>2007 (49.1)</td>
</tr>
<tr>
<td>&gt;3</td>
<td>2761 (33.8)</td>
<td>1439 (33.9)</td>
<td>1322 (33.8)</td>
<td>1355 (33.2)</td>
</tr>
<tr>
<td>Hypertension</td>
<td>4002 (49.0)</td>
<td>2052 (49.0)</td>
<td>1950 (49.8)</td>
<td>2009 (49.2)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>1299 (15.9)</td>
<td>614 (14.5)</td>
<td>685 (17.5)</td>
<td>572 (14.0)</td>
</tr>
<tr>
<td>Coronary artery disease$^c$</td>
<td>1715 (21.0)</td>
<td>927 (21.8)</td>
<td>788 (20.1)</td>
<td>898 (22.0)</td>
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<tr>
<td>Medication</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Antihypertensive</td>
<td>3725 (45.6)</td>
<td>1917 (45.1)</td>
<td>1808 (46.1)</td>
<td>1892 (46.3)</td>
</tr>
<tr>
<td>Statin use</td>
<td>2156 (26.4)</td>
<td>1041 (24.5)</td>
<td>1115 (28.4)</td>
<td>1004 (24.6)</td>
</tr>
<tr>
<td>Menopausal women</td>
<td>1732 (45.7)</td>
<td>823 (41.3)</td>
<td>909 (50.6)</td>
<td>791 (40.0)</td>
</tr>
<tr>
<td>Geography</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urban</td>
<td>5628 (68.9)</td>
<td>2309 (54.4)</td>
<td>3319 (84.8)</td>
<td>1870 (45.8)</td>
</tr>
<tr>
<td>Region</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>North</td>
<td>6690 (81.9)</td>
<td>2979 (70.1)</td>
<td>3711 (94.7)</td>
<td>2774 (67.9)</td>
</tr>
<tr>
<td>Southeast</td>
<td>939 (11.5)</td>
<td>732 (17.3)</td>
<td>207 (5.3)</td>
<td>815 (20.0)</td>
</tr>
<tr>
<td>Southwest</td>
<td>539 (6.6)</td>
<td>539 (12.6)</td>
<td>0</td>
<td>494 (12.1)</td>
</tr>
<tr>
<td>Participants with blood samples obtained, No.</td>
<td>6528</td>
<td>3645</td>
<td>2883</td>
<td>3516</td>
</tr>
<tr>
<td>Blood markers, mean (SD)</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Cholesterol, mg/dL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>181.5 (50.2)</td>
<td>181.5 (46.3)</td>
<td>181.5 (50.2)</td>
<td>181.5 (46.3)</td>
</tr>
<tr>
<td>High-density lipoprotein</td>
<td>36.4 (33.6)</td>
<td>33.6 (14.0)</td>
<td>36.4 (50.4)</td>
<td>33.6 (14.0)</td>
</tr>
<tr>
<td>Low-density lipoprotein</td>
<td>84.0 (81.2)</td>
<td>81.2 (28.0)</td>
<td>84.0 (120.4)</td>
<td>81.2 (28.0)</td>
</tr>
<tr>
<td>Triglycerides, mg/dL</td>
<td>150.4 (115.0)</td>
<td>159.3 (123.9)</td>
<td>150.4 (106.2)</td>
<td>159.3 (123.9)</td>
</tr>
<tr>
<td>High-sensitivity C-reactive protein, mg/L</td>
<td>1.9 (4.4)</td>
<td>2 (5.5)</td>
<td>1.8 (2.5)</td>
<td>2 (5.6)</td>
</tr>
<tr>
<td>Moved residence in the last 5 y$^d$</td>
<td>93 (5.0)</td>
<td>47 (5.1)</td>
<td>46 (4.9)</td>
<td>42 (4.8)</td>
</tr>
</tbody>
</table>

Abbreviations: BMI, body mass index (calculated as weight in kilograms divided by height in meters squared); NO$_2$, nitrogen dioxide; PM$_{2.5}$, particulate matter with aerodynamic diameter less than 2.5 μm.

SI conversion factors: To convert total, high-density lipoprotein, and low-density lipoprotein cholesterol to millimoles per liter, multiply by 0.0259; triglycerides to millimoles per liter, multiply by 0.0113; and high-sensitivity C-reactive protein to nanomoles per liter, multiply by 9.524.

$^a$ Except where noted otherwise, data are number (percentage) of participants.

$^b$ P value from univariate regression between coronary artery calcium and individual variables.

$^c$ Defined as 50% or more stenosis in 1 or more major epicardial vessels.

$^d$ Data are based on subpopulation of 1850 participants who had completed detailed information.
The association between PM$_{2.5}$ and CAC in postmenopausal women was significant (PM$_{2.5}$ per 10 μg/m$^3$: 34.5%; 95% CI, 5.8% to 70.9%).

In the multipollutant models with all exposure variables included, the effect estimates for both PM$_{2.5}$ and NO$_2$ were largely unchanged, with a 27.2% (95% CI, 10.8%–46.1%) greater CAC score for a 30-μg/m$^3$ increase in PM$_{2.5}$ and a 24.5% (95% CI, 3.6%–49.7%) greater CAC score per 20-μg/m$^3$ increase in NO$_2$. The concentration-response curve suggested little evidence of a nonlinear association with either PM$_{2.5}$ or NO$_2$. (eFigure 3 in the Supplement). Estimating associations by quartiles of the pollutants confirmed the reasonably linear nature of the concentration-response curves (eFigure 4 in the Supplement).

**Figure 2. Spatial Distribution of Estimated Annual Pollution Concentrations in 2015 in China**

A. Concentration of particulate matter with aerodynamic diameter less than 2.5 μm (PM$_{2.5}$) per 1 × 1 km$^2$. B. Concentration of nitrogen dioxide (NO$_2$) per 1 × 1 km$^2$.

**Figure 3. Coronary Artery Calcium (CAC) Score Associated With an Increase in Long-term Air Pollutant Exposure**

Percentage increase in CAC scores with 95% CIs (error bars). The interquartile range increases in pollutants are 30 μg/m$^3$ for particulate matter with aerodynamic diameter less than 2.5 μm (PM$_{2.5}$), 20 μg/m$^3$ for nitrogen dioxide (NO$_2$), and 15 μg/m$^3$ for ozone (O$_3$), there was a 50% decrease for distance from a roadway. The primary model included age, sex, body mass index, smoking status, smoking years, cigarettes per day, alcohol consumption, education, exercise, urbanization, regions, distance to hospital, and Beijing residence (yes or no). The primary model plus biomarkers also included levels of total cholesterol, high-density lipoprotein cholesterol, low-density lipoprotein cholesterol, triglycerides, and high-sensitivity C-reactive protein. The primary model plus multiple exposures also included PM$_{2.5}$, NO$_2$, O$_3$, and distance to road variables.
In the logistic regression analyses, there were higher odds of the presence of CAC with increasing exposures to all air pollutants and distance to roadway variables (PM$_{2.5}$ per 30 μg/m$^3$: odds ratio [OR], 1.28; 95% CI, 1.13-1.45; NO$_2$ per 20 μg/m$^3$: OR, 1.27; 95% CI, 1.23-1.32; O$_3$ per 15 μg/m$^3$: OR, 1.12; 95% CI, 1.01-1.24; distance to roadway per 50% decrease: OR, 1.04; 95% CI, 1.00-1.07) (Figure 4). Also, the associations with severe CAC scores were greater in participants with increased exposure to PM$_{2.5}$ (OR, 1.59; 95% CI, 1.20-2.12) or NO$_2$ (OR, 1.60; 95% CI, 1.18-2.17).

**Modification of Associations**

We observed stronger associations between air pollutants (PM$_{2.5}$ and NO$_2$) and CAC score in men (PM$_{2.5}$: 42.2%; 95% CI, 24.3% to 62.7%; NO$_2$: 45.7%; 95% CI, 25.3% to 69.5%) and older (age >60 years) participants (PM$_{2.5}$: 50.1%; 95% CI, 28.8% to 75.0%; NO$_2$: 55.5%; 95% CI, 31.8% to 83.6%) (eTable 4 in the Supplement). Participants with diabetes had larger NO$_2$ association estimates than did participants without diabetes (PM$_{2.5}$: 62.2%; 95% CI, 30.9% to 101.0%; NO$_2$: 31.2%; 95% CI, 13.9% to 51.0%), and participants in the north had larger association estimates than did participants in the southeast regions (NO$_2$ per 20 μg/m$^3$, north region: 42.0%; 95% CI, 21.7% to 65.6%; southeast region: −9.2%; 95% CI, −32.1% to 21.4%). There were no clear differences in the pollutant association estimates between local Beijing and non-Beijing residents (local Beijing residents, per 1-SD increase in PM$_{2.5}$: 10.3%; 95% CI, 1.8% to 19.4%; NO$_2$: 9.2%; 95% CI, 0.9% to 18.3%; and O$_3$: −3.2%; 95% CI, −11.5% to 6.0%; and non-Beijing residents, per 1-SD increase in PM$_{2.5}$: 16.1%; 95% CI, 9.4% to 23.1%; NO$_2$: 14%; 95% CI, 7.1% to 21.4%; O$_3$: 9.9%; 95% CI, 3.5% to 16.7%) (eTable 5 in the Supplement).

**Discussion**

In this large Chinese study, long-term exposures to PM$_{2.5}$, NO$_2$, and O$_3$ and proximity to traffic were associated with subclinical coronary atherosclerosis as assessed by the CAC score in residents with potential risk of CHD. The findings for PM$_{2.5}$ and NO$_2$ were robust to the use of various exposure durations and to controlling for major known CHD risk factors and copollutants. The associations were stronger among men, elderly participants, and those with diabetes. Given that CAC is strongly associated with total CHD events,4 coupled with the recently reported finding that higher long-term PM$_{2.5}$ exposure was associated with a higher risk of CHD mortality in China,10 this study may provide evidence suggesting that coronary atherosclerosis is a pathological pathway through which air pollution exposure potentially increases mortality associated with CHD.

Major strengths of our study include the large population sample size, the use of advanced methods for estimating individual-level long-term outdoor exposure concentrations, and the high quality of individual information on both the outcome measures and potential confounding factors.

![Figure 4. Odds Ratio of Detectable and High-Level Coronary Artery Calcium (CAC) Associated With Long-term Exposure Variables](image-url)
Compared with numerous epidemiological studies in China that estimated air pollution exposures from either proximal monitors \(^{22}\) or regional modeling estimates \((\geq 10 \times 10\text{-km spatial resolution})\),\(^{10}\) our exposure models produced individually resolved exposure estimates that allowed us to use accurate fine-scale contrast in investigating the association between air pollution exposure and atherosclerosis.

The associations identified in our study were likely more apparent than those in prior studies owing to the very broad range of ambient air pollution concentrations we observed. In addition, this is a population at risk for coronary atherosclerosis. Prior studies\(^{6,9,19,20}\) showing an association of air pollution or traffic exposure with the degree of CAC were very limited, with mostly null findings. A higher degree of CAC was associated with exposure to nearby roadways in the Ruhr area of Germany,\(^9\) and an association between PM\(_{2.5}\) and thoracic aortic calcifications was recently reported elsewhere.\(^{17}\) In the Multi-ethnic Study of Atherosclerosis,\(^8\) higher exposures to PM\(_{2.5}\) and NO\(_2\) were associated with faster progression of CAC, but cross-sectional associations, such as those seen in this study, have not been observed.\(^6\) Compared with previous studies of calcification in less-polluted regions, the observed magnitude of our estimates seemed smaller for PM\(_{2.5}\) (eFigure 5 in the Supplement). This is in line with a recent Global Burden of Diseases analysis\(^1\) indicating a steeper change in relative risk for countries at much lower levels of PM\(_{2.5}\) compared with those at higher values.

In 2015, more than 95% of the Chinese population was exposed to concentrations of PM\(_{2.5}\) and NO\(_2\) greater than the minimum level of our study.\(^{14}\) Because more than 40% of all deaths in China are attributable to cardiovascular disease,\(^{23}\) the potential contribution of air pollutants to the burden of cardiovascular disease in China is very large. Improving air quality to, for example, the Chinese national standards of 35 \(\mug/m^3\) for PM\(_{2.5}\) and 40 \(\mug/m^3\) for NO\(_2\) may lead to a longer life span.\(^{24}\) The association with NO\(_2\) exposure on CAC persisted even when the analysis was restricted to concentrations less than 40 \(\mug/m^3\), suggesting that associations may occur even below this level.

In our study, we also found that long-term exposure to PM\(_{2.5}\) and NO\(_2\) was associated with increased likelihood of both the presence of detectable CAC at any amount, and the most severe levels of CAC; the risk of severe CAC was most pronounced. This is consistent with 2 US studies\(^{25,26}\) that observed higher risks of mild and severe CHD associated with exposure to PM\(_{2.5}\). Together, these findings and the results of our study indicate that individuals at potential risk for CHD are a population that is especially susceptible to air pollution exposures.

Mechanisms by which PM\(_{2.5}\) exposure affects atherogenesis are not well elucidated. It has been proposed that the generation of oxidative reaction products is initiated by the reaction of pollutants with lipids or cellular membranes in the airways and lung alveoli, with circulating products triggering atherogenic mechanisms such as lipid peroxidation and high-density lipoprotein dysfunction.\(^3\) Persistent activation of this pathway is associated with the development of atherosclerosis.\(^{27}\) We cannot conclude from our study that NO\(_2\) per se is the specific pollutant responsible for these estimated effects. Nitrogen dioxide may simply be an indicator of more harmful traffic mixtures, including ultrafine particles and diesel exhaust black carbon. Diesel exhaust and ultrafine particles have been shown to elicit harmful biological responses in experimental models.\(^{28}\) Indeed, we observed that exposure to NO\(_2\) was associated with CAC and that adjusting for roadway proximity attenuated its association with NO\(_2\). Living close to a roadway is associated with a substantial increase in an individual’s exposure to traffic-related air pollutants, including tailpipe (e.g., NO\(_2\), ultrafine particles, and black carbon) and nontailpipe (e.g., particles from tire wear and friction materials) emissions and noise.\(^{29}\) This finding suggests that the synergistic effect of traffic exposure mixtures might, at least in part, operate through this mechanism.

Prior evidence\(^8\) has supported the role of PM\(_{2.5}\) and traffic-related air pollutants such as NO\(_2\) in the development of atherosclerosis, but less research has focused on the cardiovascular effects of O\(_3\): the long-term effect of exposure to O\(_3\) on subclinical CHD is still largely unknown. Ambient O\(_3\) is a powerful oxidizing agent and common air pollutant worldwide. We observed a significant association...
between O3 and degree of CAC, although this association could not be disentangled from the effects of PM2.5 and NO2 because of the spatial correlations between these exposure variables.

There was evidence that elderly participants and those with diabetes were more susceptible to these exposure effects than other participants. Patients with diabetes have a greater overall coronary plaque burden and a higher rate of multivessel disease. It is possible that air pollution could worsen the underlying diabetes disease course by exacerbating insulin resistance or by instigating adverse biological responses (eg, endothelial dysfunction) that promote future diabetic atherosclerosis. Among women, we observed a stronger association of exposure to PM2.5 with risk of atherosclerosis in postmenopausal women, a group for whom an association between PM2.5 and cardiovascular risk has previously been demonstrated in the United States.

Limitations
This study has several limitations. First, because this was an outpatient population, it may not fully reflect the general population. However, this population had relatively low pretest probability of CHD (27.3%); 29.0% had no symptoms, and 64.0% had nonanginal chest pain. Moreover, this population resembled population-based samples in terms of risk factors and CAC levels. Therefore, this cohort can represent a low-risk natural population. Second, we cannot rule out biases due to unmeasured confounders, such as occupation history. In addition, the main analysis of this study did not account for individuals who moved in recent years, although we did sensitivity analysis in a subset of the samples that showed consistent association. Third, this is a cross-sectional study involving the baseline examination of an ongoing prospective cohort study. Future longitudinal analysis of this prospective cohort may provide additional information.

Conclusions
This study found that long-term exposure to outdoor air pollution and traffic pollutants in China, specifically PM2.5 and NO2, was associated with the development of CAC. This finding should contribute to an understanding of air pollutant effects worldwide, providing both much needed locally generated data and supportive evidence to inform the air pollution standard-setting process on a global scale.

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REFERENCES


**SUPPLEMENT.**

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