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# Associations of early pregnancy air pollution with adverse birth outcomes and infant neurocognitive development in the Complex Lipids in Mothers and Babies (CLIMB) cohort

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- 2 and infant neurocognitive development in the Complex Lipids in Mothers
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- 25 Abstract
- Objectives: to investigate the associations of traffic-related air pollution exposures in early
- 27 pregnancy with birth outcomes and infant neurocognitive development
- **Design:** cohort study
- **Setting:** in the maternity clinics of two centres (the First Affiliated Hospital of Chongqing
- 30 Medical University (FCQMU) and Chongqing Health Centre for Women and Children
- 31 (CHC)).
- Participants: women who were between 20 and 40 years of age and were at 11–14 weeks
- 33 gestation with a singleton pregnancy were eligible for participation. Women were excluded if
- 34 they had a history of premature delivery before 32 weeks of gestation, maternal milk allergy
- or aversion, or severe lactose intolerance. 1,273 pregnant women enrolled in 2015-2016 and
- 36 1,174 live births were included in this analysis.
- **Interventions:** Air pollution concentrations at their home addresses, including particulate
- matter (PM) with diameter  $\leq 2.5 \mu m$  (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>), during pre-
- 39 conception and each trimester period (TI, T2, T3) were estimated using land-use regression
- 40 (LUR) models.
- Primary and secondary outcome measures: birth outcomes (i.e., birthweight, birth length,
- 42 preterm birth (PTB), low birth weight (LBW), large for gestational age (LGA) and small for
- 43 gestational age (SGA) status) and neurodevelopment outcomes measured by the Chinese
- version of Bayley Scales of Infant Development (CBSID)).
- **Results:** An association between SGA and per Interquartile range (IQR) increases in NO<sub>2</sub> was
- 46 found during the whole pregnancy (Odd ratio (OR): 1.60, 99% confidence interval (CI): 1.03,
- 47 2.48) after co-adjusted for PM<sub>2.5</sub>. Both PM<sub>2.5</sub> and NO<sub>2</sub> exposure in the 90 days prior to
- 48 conception (but not during) were associated with lower PDI score (β: -6.15, 99% CI: -9.69, -
- 49 2.61; β: -2.83, 99% CI: -4.27, -0.93, respectively).
- 50 Conclusions: NO<sub>2</sub> levels during pregnancy were associated with increased risk of SGA, while
- both PM<sub>2.5</sub> and NO<sub>2</sub> pre-conception were associated with adverse neurodevelopment outcomes
- at 12 months of age.
- **Keywords:** Air pollution; birth outcomes; child cognition

#### Article summary

# Strength and limitation

- We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification if using monitoring stations.
- This is one of the few studies to investigate both pre-conception and prenatal PM<sub>2.5</sub> and NO<sub>2</sub> exposure with neurodevelopment outcomes among young infants, in the context of a relatively high air pollution urban environment.
- Our sample size was relatively small, limiting the statistical power to assess several outcomes.
- We defined exposure windows for clinically-defined trimesters.
- The performance of the NO<sub>2</sub> spatiotemporal model was moderate (COR-R<sup>2</sup>: 0.39), which may introduce exposure misclassification and therefore bias in the coefficients.

#### Introduction

Air pollution is a major environmental factor that has been linked to a range of adverse health outcomes in children. Maternal exposure to air pollutants during pregnancy, especially PM<sub>2.5</sub> and NO<sub>2</sub>, has been found to be associated with adverse birth outcomes, including PTB(1), term low birth weight (TLBW)(2), and small for gestational age (SGA) status (3). According to the developmental origins of health and disease (DOHaD) hypothesis, prenatal environmental exposures to air pollution may lead to adverse birth outcomes and subsequently increase the susceptibility to the development of certain diseases later in life (4). A number of epidemiological studies have linked prenatal air pollutant exposure to neurodevelopmental disorders such as autism spectrum disorder (ASD), attention deficit hyperactivity disorder (ADHD), and cognitive impairment (5). Although the underlying biological mechanisms are still unclear, some studies indicated that prenatal air pollution exposure may induce systemic oxidative stress that triggers intrauterine inflammation, leading to damage to several fetal organs, including the brain (6, 7).

It is also unclear that whether the adverse effects of air pollution start earlier before conception.

81 Three months before conception was considered as a critical developmental window for

gametogenesis. Air pollution exposure during three months preconception or early stages of

pregnancy may have adverse effects on gametogenesis of sperm (8, 9) and ova cells (10).

84 Exposures to PM<sub>2.5</sub> in preconception period have been associated with various

neurodevelopmental outcomes, such as neural tube defects (11), lower psychomotor

development scores (12), higher risk of ASD (13, 14), and higher risk of intellectual disability

87 (15). While there is growing evidence for the effects of preconception  $PM_{2.5}$  exposure on the

risk of adverse neurodevelopmental outcomes, no study examined the effects of preconception

NO<sub>2</sub> exposure. There are also inconsistencies across studies and heterogeneities in health

90 outcomes and air pollutant levels (12).

91 Many studies have reported the effects of prenatal exposure to air pollution on

92 neurodevelopmental function in children. However, the reported associations vary, due to the

93 heterogeneous assessments of air pollutants exposure and neurodevelopmental outcomes (5,

94 16). Moreover, most of these studies were conducted in relatively developed countries where

95 pollution is lower; little evidence has come from populations in developing countries such as

China (17-20), that are more likely to experience more severe air pollution exposure.

The current study leveraged the Complex Lipids in Mothers and Babies (CLIMB) cohort, a prospective birth cohort recruited in Chongqing, China(21), with trimester-specific maternal PM<sub>2.5</sub> and NO<sub>2</sub> air pollution exposure derived from a spatio-temporal LUR model (22). The aim of this analysis was to examine the association between PM<sub>2.5</sub> and NO<sub>2</sub> air pollution preand during pregnancy, with birth and infant neurocognitive development outcomes at 12 months of age.

A key aspect in all studies like this one is the accuracy of documenting exposure; a recent Chinese study determined air pollution exposure based on the nearest monitoring station data (18) may not reflect the temporal and spatial variability of pollutant exposures among participants. The current study employed common air pollutant exposure models based on advanced geographic information systems (GIS), to address some of the limitations of previous studies (23).

In addition, the timing of in exposure is also critical in determining the effects of exposure on developmental outcomes. Indeed, the evidence from previous studies on the sensitive time window for exposure pre- and during pregnancy remains inconclusive. Some studies have indicated that the early-to-mid pregnancy phase may be a critical period in terms of the impact of air pollution on neurodevelopment(17, 24). Early pregnancy is particularly important for neurogenesis and neuromigration, making it a susceptible period (25). However, actual findings have been varied, with some studies showing stronger associations for middle or late pregnancy (18, 19, 26). More studies identifying critical periods are needed to enhance our understanding of how pre-conception and prenatal air pollution exposure affect neurodevelopment. With this cohort, we are able to examine the effects of exposure preconception, at each trimester, as well as across the entire pregnancy.

#### Methods

#### **Study population**

A description of participant recruitment in the CLIMB cohort has been described previously (27). In brief, women who were between 20 and 40 years of age and were at 11–14 weeks gestation with a singleton pregnancy were eligible for participation. Women were excluded if they had a history of premature delivery before 32 weeks of gestation, maternal milk allergy or aversion, or severe lactose intolerance.

From September 2015 to November 2016, a total of 1,500 women were recruited into the cohort.

Participants attended six visits at the First Affiliated Hospital of Chongqing Medical University
and Chongqing Health Centre for Women and Children: 11–14 weeks' gestation (visit 1), 22–
28 week's gestation (visit 2), 32–34 week's gestation (visit 3), at birth (visit 4), 6 weeks
postnatal (visit 5), and 12 months postnatal (visit 6).

Women who withdrew from the study (n = 146), terminated their pregnancy (n = 29), miscarried (n = 12) or were lost to follow-up (n = 40) were excluded from the analysis, leaving a sample size of 1,273 women. Analyses were restricted to mothers whose detailed residential addresses during pregnancy were known (**Figure 1**). A total of 1,174 live births were thus included in the pregnancy and neonatal outcomes analysis. Subsequently, at 1 year follow-up, 946 children were included in the analysis of neurodevelopment outcomes.

# **Exposure assessment**

The address of participants was collected at the first visit. Exposure assessment based on spatiotemporal land use regression (LUR) models for PM<sub>2.5</sub> and NO<sub>2</sub> were developed for the study region. The study area focused on the urban center of the Chinese municipality of Chongqing (Figure 2). A description of the methodology of exposure modelling has been reported previously (22). Briefly, the models included both spatial and temporal components of exposure. PM<sub>2.5</sub> and NO<sub>2</sub> concentration data were collected from 17 routine monitoring sites operated by the Chongqing Environmental Monitoring Center in 2015-2016. For the spatial component of models, we calculated annual average concentrations of each pollutant in 2015, and fit linear regression models using five groups of geographic data (road network, land use, topography, vegetation, and population density) as spatial predictor variables. For the temporal component of models, we calculated the residuals from the spatial component at each monitoring site on a daily basis by subtracting the predicted annual average concentration from the observed daily average concentrations measured in 2015 and 2016, and then fitted generalised additive models (GAM) using seven groups of meteorological data (temperature, amount of rainfall, rainfall events, relative humidity, horizontal visibility, wind direction and wind speed) as temporal predictor variables. The meteorological variables were used to account for the influence of weather on the change in air pollution concentration over time. To account for the remaining spatial autocorrelation, the smoothed terms of longitude and latitude were fit to spatiotemporal residuals which were calculated by subtracting the sum of the spatial

temporal predictions from the measured daily average concentrations in 2015 and 2016. The performance of the PM<sub>2.5</sub> spatiotemporal models was good (COR-R<sup>2</sup>: 0.72) and the NO<sub>2</sub> spatiotemporal model was moderate (COR-R<sup>2</sup>: 0.39) when providing concentration estimates in absolute terms.

Combining the family address coordinates of each pregnant woman and the gestation period of the pregnancy (calculated from the date of last menstrual period to the date of delivery), we used this spatiotemporal model to estimate the average exposure of each pregnant woman in 90 days prior to pregnancy (90D), first trimester (T1), second trimester (T2), third trimester (T3) and whole pregnancy period (WP), respectively.

#### **Outcomes**

### **Birth Outcomes**

Birth outcomes were determined by experienced obstetricians and abstracted from the medical records. Birth outcomes included: birthweight (in grams), birth length (in centimetres), PTB, low birth weight (LBW), large for gestational age (LGA) and SGA status (28). PTB was defined as delivery before 37 weeks. LBW was defined as weighing less than 2500 g at birth. LGA and SGA were indicated by birth weight greater than and less than the 90th and 10th percentile within this study for the gestational age by sex respectively (29). Term low birth weight was not considered due to a small sample size of only 8 cases.

#### **Neurodevelopment outcomes**

The Chinese version of Bayley Scales of Infant Development (CBSID) was used to assess mental and psychomotor development for infants in this study. The CBSID is appropriate for evaluation of infants from 2–30 months old (30) and takes into consideration each infant's age in days. Infants were assessed at around 12 months (range from 11 months and 15 days to 12 months and 15 days) by a trained examiner, with ages corrected for preterm birth. These scales have been formally adapted to the Chinese language and locally standardized to become culturally appropriate, with two main indexes: the Mental Development Index (MDI) and the Psychomotor Development Index (PDI). The MDI component comprised 163 items and assessed age-appropriate items related to cognitive functioning, personal and social

development, and language development (see eTable 1 in the Supplement). The PDI component comprised 81 items and assessed age-appropriate fine and gross motor skills (see eTable 2 in the Supplement). The test provided raw scores for mental and psychomotor development that were converted to standardized (in terms of age in days) MDI and PDI scores, based on norms for the Chinese population. As with other forms of the Bayley test these index scores have a mean of 100, and a standard deviation of 15, with a lower score reflecting poorer performance (31). If an infant refused to cooperate with the examiners to finish the task, a second assessment was arranged within two weeks. If the infant could not cooperate at the second BSID assessment, their data were classified as missing.

#### **Covariates**

Socio-demographic data were collected through interviews by trained nurses. The following potential confounders were identified: maternal age at enrolment (in years), infant sex (male/female), maternal BMI at 11–14 weeks' gestation (kg/m²), parity (Yes/No), monthly household income level (categorized as: <2,000 yuan, 2,000–7,000 yuan, 7,000–10,000 yuan, or >10,000 yuan), season of birth (categorized as: Spring (Mar-May), Summer (Jun-Aug), Autumn (Sep-Nov) or Winter (Dec-Feb)). Marital status (single/married) and smoking or drinking during pregnancy (Yes/no) were not taken into account in this analysis because of the homogeneity of the study population (i.e., 98.6% women were married and 99.6% women reported not smoking or drinking alcohol during pregnancy).

## Statistical analyses

Data were described in terms of mean  $\pm$  SD or median (IQR) for continuous variables, or as percentages for categorical variables. Modelled PM<sub>2.5</sub> and NO<sub>2</sub> exposure levels in 90D, T1, T2, T3 and WP were considered separately. We examined the correlation between each of the exposures in the different pregnancy periods. For birth outcomes, multivariable linear regression was used for continuous outcomes (e.g., birth weight and birth length) to estimate  $\beta$  coefficient and their 95% confidence intervals (CIs) and multivariable logistic regression for binary outcomes (e.g., PTB, LBW, LGA and SGA status) to estimate odds ratio (OR) and 99% CIs. For mental and psychomotor development (e.g., MDI and PDI scores), multivariate linear), multivariable linear regression models were fit to estimate  $\beta$  coefficient and their 99%CIs.

Models were adjusted for maternal age at enrolment, infant sex, maternal BMI at 11–14 weeks gestation, primiparity, monthly household income level, and season of birth. We also ran coexposure models to estimate associations of one air pollutant with outcomes adjusted for the other air pollutant (i.e., PM<sub>2.5</sub> in T1 adjusted for NO<sub>2</sub> in T1). Effect estimates are reported for each IQR increase of PM<sub>2.5</sub> and NO<sub>2</sub>. We also tested for indirect effects of NO<sub>2</sub> on PDI mediated by SGA with "medsem" commands. All analyses were performed using STATA version 17. A p-value of <0.01 was considered statistically significant to address multiple comparisons in the analyses.



229	Results

# **Study Participants**

Participant characteristics are presented in **Table 1**. Of those participating women, the mean age was 28 years and mean BMI was 21.5 kg/m<sup>2</sup>. 98.0% of women were of Han ethnicity, 77.9% were primiparous, and 67.6% had completed tertiary education. 33 (2.8%), 30 (2.6%), 108

(9.2%), 84 (7.2%) of the 1,174 births considered in this analysis were classified as PTB, LBW,

LGA and SGA, respectively. For those 946 children who completed the BSID test, the mean

MDI score was 94.7 (SD: 17.7) and the mean MDI score was 87.4 (SD: 14.9).

# **Exposure assessment**

Median PM<sub>2.5</sub> exposure levels were 57.31  $\mu$ g/m3 (IQR: 5.76) and median NO<sup>2</sup> exposure levels were 50.46  $\mu$ g/m3 (IQR: 5.51) during the whole pregnancy period (eTable 3 in the Supplement). For PM<sub>2.5</sub>, the concentration in the pre-conception and T1 were considerably lower than other periods, close to 10  $\mu$ g/m3. The between-trimester and 90D values for NO<sub>2</sub> were generally moderately correlated (Pearson's r > 0.5). The correlation coefficients of PM<sub>2.5</sub> were more variable between time periods reflecting the high variability of PM<sub>2.5</sub> concentrations, with values ranging from -0.78 to +0.68. Correlations between PM<sub>2.5</sub> and NO<sub>2</sub> in the same pregnancy period were moderately correlated (Pearson's r ~0.6, eTable 4 in the Supplement).

#### **Association with birth outcomes**

In the unadjusted models (**eTable 5 in the Supplement**), higher exposure concentrations of PM<sub>2.5</sub> in T3 were significantly associated with lower birth length (β: -0.32, 99% CI: -0.57, -0.07; per IQR increase). A risk between SGA and increases in NO<sub>2</sub> (per IQR) was found in T2 (OR: 1.46, 99% CI: 1.01, 2.11), T3 (OR: 1.58, 99% CI: 1.03, 2.42) and in the whole pregnancy period (OR: 1.44, 99% CI: 1.04, 2.00). We observed no evidence of associations of NO<sub>2</sub> with overall birth weight, birth length and other adverse birth outcomes (e.g., PTB, LBW, and LGA). In the adjusted models (**Table 2**), we found slightly reduced effect size for NO<sub>2</sub> and SGA in the whole pregnancy period (OR: 1.33, 99% CI:0.92, 1.75) compared with the unadjusted model. We observed no evidence of associations with birth length in the adjusted models. After

co-adjustment for PM<sub>2.5</sub> (see eTable 6 in the Supplement), the association of NO<sub>2</sub> with SGA
 became significant in the whole pregnancy period (OR: 1.60, 99% CI: 1.03, 2.48)

#### Association with infant neurodevelopment outcomes

- In unadjusted models, PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower MDI and PDI scores in offspring ( $\beta$ : -3.54, 99% CI: -5.94, -1.13;  $\beta$ : -3.42, 95% CI: -5.44, -1.40) (eTable 7 in the Supplement). We also observed an unexpected positive association between PM<sub>2.5</sub> exposures in second trimester with MDI ( $\beta$ : 4.21, 99% CI: 1.87, 6.56) and PDI ( $\beta$ :2.63, 99%CI: 0.65, 4.61). Exposure to NO<sub>2</sub> was associated with lower PDIs in the 90 days prior to conception (-2.86, 99% CI: -4.46, -1.26), and T3 (-1.97, 99% CI: -3.70, -0.23). We did not observe any association between NO<sub>2</sub> and MDI in any pregnancy periods.
- In the adjusted models (Table 5), we found PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower PDI scores (β: -6.15, 99% CI: -9.69, -2.61). Similarly, there was also a significant association of increased NO<sub>2</sub> exposure and lower PDI score (β: -2.83, 99% CI: -4.27, -0.93). However, the positive association between PM<sub>2.5</sub> exposures in second trimester with PDI (β: 3.76, 99% CI: 0.49, 7.02) remained. We did not observe any association with MDI in any pregnancy periods.
  - In the co-exposure models (**see eTable 8 in the Supplement**), PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower PDI scores (β: -4.74, 99% CI: -8.67, -0.81). We also observed a positive association between PM<sub>2.5</sub> exposures in second trimester with PDI (β: 5.51, 99% CI: 1.86, 9.16). Exposure to NO<sub>2</sub> was significantly associated with lower PDI in the second trimester (β: -2.11, 99% CI: -4.11, -0.12) and whole pregnancy period (β: -1.68, 99% CI: -3.28, -0.08). In the mediation analyses, we did not observe any indirect effect for PM<sub>2.5</sub> and NO<sub>2</sub> on BSID scores mediated by SGA in all pregnancy periods (**see eTable 9 and 10 in the Supplement**).

#### Discussion

We analyzed associations between modelled PM<sub>2.5</sub> and NO<sub>2</sub> pre- and during pregnancy with birth and neurodevelopment outcomes in singleton children born in a south-western metropolis of China in 2015-16. We found the likelihood of SGA increased by 60% per IQR higher

exposure to  $NO_2$  in the whole pregnancy periods after adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11-14 weeks' gestation, primiparity, monthly household income level, and season of births and  $PM_{2.5}$ . For childhood cognitive development, increased exposure to  $PM_{2.5}$  and  $NO_2$  in the 90 days prior to conception were both associated with lower PDI scores, with the effect size per IQR being higher for  $PM_{2.5}$  than for  $NO_2$ . We also found a positive association between  $PM_{2.5}$  exposures in second trimester with PDI. While SGA was associated with  $NO_2$  exposures, SGA was not found to mediate the effects of  $PM_{2.5}$  and  $NO_2$  on BSID scores in this study.

Many studies from other geographic areas, including Europe (32-34), the United States (24, 26), and Asia (17, 35-37) have found that prenatal air pollution exposure has a negative impact on a variety of neurodevelopmental outcomes. Our finding of a negative association between prenatal NO<sub>2</sub> air pollution exposure and infant neurocognitive development is consistent with these reports. A recent Chinese birth cohort study of 15,778 child-mother pairs in Foshan reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with increased risk of suspected developmental delay (OR: 1.06, 95% CI: 0.94, 1.19) measured by a five-domain scale and developmental quotient (DQ) (17). A birth cohort study of 520 mother-child pairs in South Korea reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with impairment of psychomotor development ( $\beta = -1.30$ , p = 0.05) but – as in the present study not with cognitive function ( $\beta = -0.84$ , p = 0.20) (35). However, results from previous research varied by air pollutants. For example, a Chinese study of 1193 mother-newborn pairs in Changsha found significant associations between PM<sub>2.5</sub> exposure in trimester 2 and lower neurobehavioral developmental scores, while other air pollutants such as PM<sub>10</sub>, carbon monoxide (CO), and Sulfur dioxide (SO<sub>2</sub>) had null or even reverse associations. In this study, we observed that the negative effect of NO<sub>2</sub> exposure during pregnancy on PDI is significant at 5% level, whereas no such effect was found for PM<sub>2.5</sub>; in the co-exposure model, this negative effect of NO<sub>2</sub> was stronger and became significant at 1% level after adjustment for PM<sub>2.5</sub>. This heterogeneity may relate to the time of exposure assessment, the type of instruments or evaluators, and the levels of air pollution. In addition, air pollution mixtures may have differed among the study regions, thus there are several potential explanations for the heterogeneity of the findings.

To date, most studies on prenatal air pollution exposure and child neurodevelopment have been conducted in developed countries with relatively low levels of air pollution. In this study, the level of air pollution was higher (median PM<sub>2.5</sub>: 57.31 µg/m<sup>3</sup>, IQR: 5.76; median NO<sub>2</sub>:

50.46  $\mu$ g/m³, IQR: 5.51) compared to studies in developed countries such as Europe and the United States. In a multi-centre European cohort, the mean PM<sub>2.5</sub> and NO<sub>2</sub> exposure levels during pregnancy were 13.4  $\mu$ g/m³ and 11.5  $\mu$ g/m³ (32). Researchers found that the psychomotor development score significantly decreased by 0.68 points (95% CI: -1.25, -0.11) for every 10  $\mu$ g/m³ increase in NO<sub>2</sub>, and there was also a non-significant decrease of 1.64 points (95% CI: -3.47, 0.18) for every 5  $\mu$ g/m³ increase in PM<sub>2.5</sub> during pregnancy (32). Noticeably, we did not find associations of either NO<sub>2</sub> or PM<sub>2.5</sub> during pregnancy with neurodevelopmental delay. Factors such as the types of pollutants and concentrations may differ between China and other regions with a lower air pollution level, leading to variations in the observed effects.

Contrary to expectations, we found significant positive associations between prenatal exposure to PM<sub>2.5</sub> air pollution in the second trimester and PDI. Given the prior literature and the high variability observed here, we believe that this is likely spurious/sample specific. Several epidemiological studies have reported associations between prenatal exposure to high levels of PM<sub>2.5</sub> and lower neurodevelopment in children ranging in age from 6 months to 6 years (12, 33, 38-40). In agreement with our findings, a multi-centre cohort study from six European countries investigated the effects of prenatal exposure to multiple air pollutants including PM<sub>2.5</sub>, PM<sub>10</sub>, coarse particles, NO<sub>2</sub> and nitrogen oxides (NOx) among 9482 children between 1 and 6 years; the authors found nonsignificant positive associations between prenatal PM<sub>2.5</sub> exposure and normal neurodevelopment (β: 1.64, 95% CI: -3.47, 0.18; per 5 µg/m<sup>3</sup> increase in PM<sub>2.5</sub>) (32). Similarly, another study examining the effects of multiple pollutant exposures on early childhood cognition at 40 days of age in a highly exposed area of Spain also found PM<sub>10</sub>, PM<sub>coarse</sub>, PM<sub>2.5absorbance</sub>, NO<sub>2</sub>, NO<sub>x</sub>, and Ozone (O<sub>3</sub>) were linked to lower motor function in children, except for PM<sub>2.5</sub> (41). The inconsistent findings could be because of heterogeneity between studies in terms of exposure (e.g., exposure assessment methods used, PM<sub>2.5</sub> exposure levels, or composition of  $PM_{2.5}$ ).

Aside from the conflicting findings regarding prenatal PM<sub>2.5</sub> exposure and neurodevelopmental outcomes, results regarding the most potential sensitive time windows before and during pregnancy are also inconclusive. Some studies suggested that early-to-mid pregnancy might be a potential sensitive period (17, 24), while other studies found stronger associations for midate pregnancy, thus results are equivocal (18, 19, 26). The potential biological mechanisms by which air pollution could affect neurodevelopment are not yet clearly understood. There is evidence suggesting that exposure to prenatal PM<sub>2.5</sub> could potentially induce maternal immune activation during pregnancy (42). Higher levels of cytokines or reactive oxygen species may

potentially interfere fetal neurodevelopment through three mechanisms: crossing the placental barrier into the fetal body, inducing fetal immune dysregulation, and contributing to inadequate placental perfusion that affects nutritional processes and oxygenation of maternal blood(43). More research is needed to investigate trimester effects of air pollution on neurodevelopment and provide better understanding on the underlying biological mechanisms. Our study is the first to consider an exposure window 90 days prior to conception for NO<sub>2</sub>. A novel observation is that effects of NO<sub>2</sub> or PM<sub>2.5</sub> air pollution on child cognition can be seen at least 90 days prior to conception, representing a potentially vulnerable periods in relation to air pollution on neurodevelopment. Similar results were found in previous study recruited 1329 mother-child pairs in Wuhan, China (12). This study reported a higher level of PM<sub>2.5</sub> during preconception (Median:  $76.1 \,\mu\text{g/m}^3$ ) and in the first trimester (Median:  $82.3 \,\mu\text{g/m}^3$ ). This study found for each doubling of PM<sub>2.5</sub> exposure during preconception, children's PDI scores was reduced by 8.23 (95% CI: -10.01, -6.44) points. A potential explanation is that preconception air pollution exposures induce genetic and epigenetic alterations in sperm, that increase the risk of adverse health outcomes in offspring(44, 45). To date, all studies examined the effect of maternal preconception exposure while omitting paternal exposures (16). Future studies should consider the effect of preconception paternal exposure in relation to childhood health outcomes.

This study has several strengths. We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification if using monitoring stations. This is one of the few studies to investigate both pre-conception and prenatal  $PM_{2.5}$  and  $NO_2$  exposure with neurodevelopment outcomes among young infants, in the context of a relatively high air pollution urban environment.

A major limitation of this study was that our sample size was relatively small, limiting the statistical power to assess several outcomes, although the higher exposures in Chongqing than in some other studies may increase probability of detecting effects. In terms of limitations, due to a lack of information on participant time-activity patterns, exposure estimates in this study refer only to ambient concentrations at home addresses, and no other activity spaces (e.g., indoor, workplace, commuting) were considered. We may have thus underestimated total air pollution exposure. Second, we defined exposure windows for clinically-defined trimesters; sensitive periods may be shorter or longer than 3 months, or they may exist in the overlap of multiple trimesters. However, we were unable to investigate the sensitive time windows using established methods such as distributed lag non-linear models due to the lack of highly time-resolved air pollution estimates. Third, the performance of the NO<sub>2</sub> spatiotemporal model was

moderate (COR-R<sup>2</sup>: 0.39), which may introduce exposure misclassification and therefore bias in the coefficients. Finally, we were unable to include some other air pollutants such as polycyclic aromatic hydrocarbons (PAH), black carbon (BC) and Ozone, which have bene found particular harmful to neurodevelopment in children (46).

#### Conclusion

This study provides evidence for an association between  $NO_2$  exposure pre- and during pregnancy with birth and neurodevelopmental outcomes in a birth cohort in Chongqing, China. Exposure to  $NO_2$  and  $PM_{2.5}$  exposure before pregnancy was associated with a lower psychomotor development score and further studies are warranted.



399	List of abbreviations
	ADHD

ADHD Attention deficit hyperactivity disorder

ASD Autism spectrum disorder

BMI Body mass index

BSID Bayley Scales of Infant Development

CBSID Chinese version of Bayley Scales of Infant Development

CI Confidence interval

CLIMB Complex Lipids in Mothers and Babies

CO Carbon monoxide

DOHaD Developmental origins of health and disease

DQ Developmental quotient

GIS Geographic information systems

IQR Interquartile range

LBW Low birth weight

LGA Large for gestational age

LUR Land-use Regression

MDI Mental Development Index

NO<sub>x</sub> Nitrogen oxides

NO<sub>2</sub> Nitrogen dioxide

OR Odd ratio

 $O_3$  Ozone

PDI Psychomotor Development Index

PM Particulate matter

PM<sub>2.5</sub> Particulate matter with diameter  $\leq$ 2.5µm

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PTB	Preterm birth
SGA	Small for gestational age
$SO_2$	Sulfur dioxide
TLBW	Term low birth weight
T1	First trimester
T2	Second trimester
T3	Third trimester
WP	Whole pregnancy period
90D	90 days prior to pregnancy

#### **Declarations**

#### Ethics approval and consent to participate

- 403 Ethical approval for this study was granted by the Ethics Committee of Chongqing Medical
- 404 University (#2014034). The participants provided their written informed consent to
- participate in this study. Written informed consent was obtained from the individual(s) for the
- 406 publication of any potentially identifiable images or data included in this article.

#### Availability of data and materials

- The data that support the findings of this study are available from Chongqing Medical
- 409 University but restrictions apply to the availability of these data, which were used under
- license for the current study, and so are not publicly available. Data are however available
- from the authors upon reasonable request and with permission of Chongqing Medical
- 412 University.

#### 413 Conflicts of interests

The authors declare that they have no conflicts of interests.

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#### 422 Author statement

- 423 Y. X., T.L.H., H.Z. and P.B. conceived and designed research; T.Z., Y.X. and H.Z. recruited
- 424 the patients and collected the samples; T.K., A.H., and J.G. constructed the air pollution
- model; Y.C. analyzed, interpreted the data and prepared the figures; Y.C and T.K were major
- 426 contributors in writing the manuscript text; YSC, JC, TLH, YX, ALH, and PNB substantively
- revised the manuscript; All authors read and approved the final manuscript.

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#### Tables and figures

Figure 1 Flow chart of the study population in CLIMB

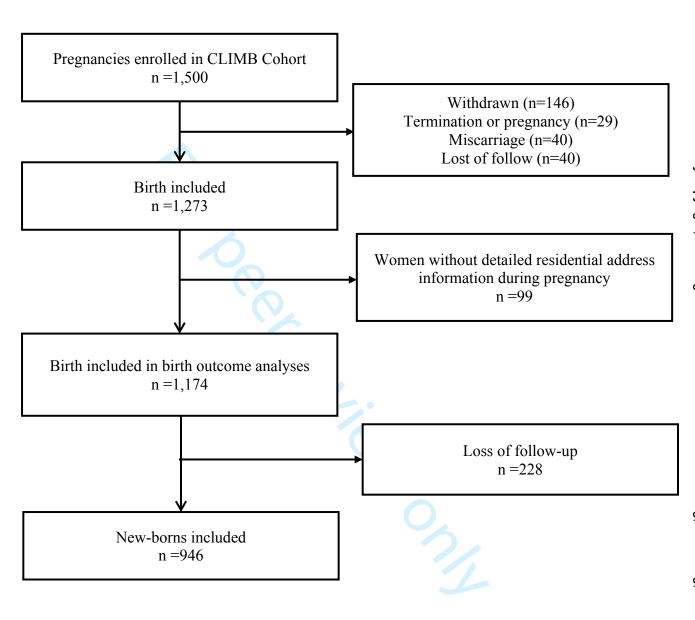


Figure 2 Study area and location of monitoring sites (OpenStreetMap contributors, 2015; https://data.nextgis.com/en/region/CN-50/).

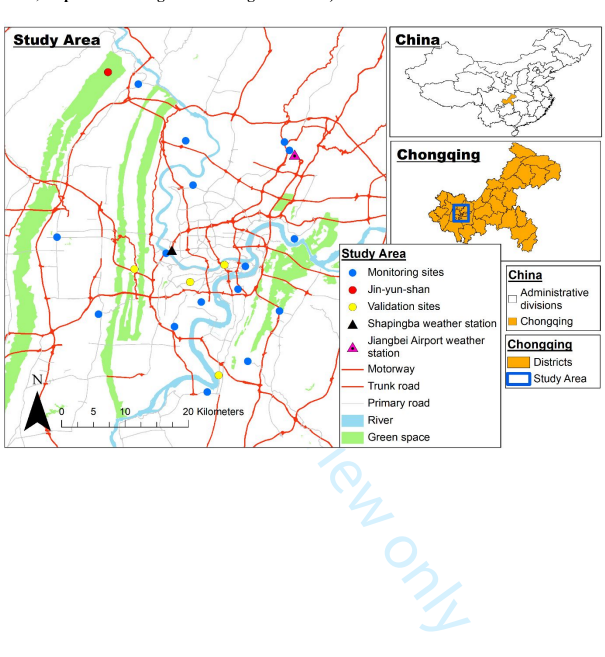


Table 1 Characteristics of study sample in the CLIMB cohort (N = 1,174)

Characteristic of mothers	N	$n (\%)$ or mean $\pm SD$	Characteristic of child	N	$n (\%)$ or mean $\pm SD$
Maternal age (Years)	1,174	$28.7 \pm 3.5$	(week)	1,174	$39.4 \pm 1.5$
BMI (kg/m²)	1,174	$21.5 \pm 2.9$	Birth weight (g)	1,165	$3314.4 \pm 428.8$
Han ethnicity (%)	1,174		Birth length (cm)	1,149	$49.7 \pm 1.9$
Yes	,	1,151 (98.0%)	Apgar score at 1 min	1,035	$9.4 \pm 1.3$
No		23 (2.0%)	Apgar score at 5 min	1,035	
Marital status (%)	1,174		New born sex	1,172	
Single		16 (1.4%)	Female		561 (47.9%)
Married		1,158 (98.6%)	Male		9.9 ± 3.0  561 (47.9%) 611 (52.1%)  33 (2.8%) 1,141 (97.2%)
Primiparity (%)	1,174		Birth outcomes		
Yes		914 (77.9%)	Preterm birth (PTB)	1,174	
No		260 (22.1%)	Yes		33 (2.8%)
History of miscarriage or abortion (%)	1,174		No		1,141 (97.2%)
Yes		553 (47.1%)	Low birth weight (LBW)	1,174	
No		621 (52.9%)	Yes		30 (2.6%)
Smoking/drinking during pregnancy (%)	1,174		No		30 (2.6%) 1,141 (97.2%)
Yes		5 (0.4%)	Large for gestational age (LGA)	1,174	
No		1,169 (99.6%)	Yes		108 (9.2%)
Education level	946		No		1,066 (90.8%)
Low: High school or below		306 (32.3%)	Small for gestational age (SGA)	1,174	84 (7.2%)
High: College/uni or above		640 (67.6%)	Yes		84 (7.2%)
Job	946		No		1,090 (92.8%)
Full-time		762 (80.5%)	BSID test	946	
Housewife		82 (8.7%)	MDI (mean ± SD)		94.7 ± 17.7
Others		102 (10.8%)	PDI (mean ± SD)		$1,090 (92.8\%)$ $94.7 \pm 17.7$ $87.4 \pm 14.9$ $411 (35.01\%)$ $263 (22.40\%)$
Household income (Monthly)	946		Birth season	1,174	•
<2000 RMB		186 (19.7%)	Spring (Mar-May)		411 (35.01%)
2000-4000 RMB		329 (34.8%)	Summer (Jun-Aug)		263 (22.40%)
4000-7000 RMB		292 (30.9%)	Autumn (Sep-Nov)		198 (16.87%)
7000-10000 RMB		139 (14.7%)	Winter (Dec-Feb)		302 (25.72%)

			ВМЈ Ор	en	/bmjopen-2023 d by copyright,		Page
435 <b>T</b>	Table 2 Associations between	en PM <sub>2.5</sub> and NO <sub>2</sub> expos	ure in different pre	gnancy periods and	adverse Firth out	comes (adjusted m	odels)
		Mean diffe	erence		ng for Odd	l ratios	
		Birth weight, grams	Birth length, cm	PTB (case: 33)	LBW (cass 50)	LGA (case: 108)	SGA (case: 84)
		99% CI	99% CI	99% CI	2024. D relatec 99% CI	99% CI	99% CI
I	Per IQR increase in	(N=941)	(N=927)	(N=945)	N=945) to to the	(N=945)	(N=945)
Estimated	90 days prior to conception	59.73 (-40.56, 160.01)	0.15 (-0.28, 0.58)	0.24 (0.06, 1.01)	0.49 (0.13) (0.49) (0.13) (0.49)	1.4 (0.59, 3.33)	1.66 (0.58, 4.73)
exposure	First trimester	6.21 (-99.01, 111.42)	0.04 (-0.42, 0.50)	0.88 (0.19, 4.02)	0.76 (0.124)	0.86 (0.36, 2.05)	1.33 (0.45, 3.95)
to PM <sub>2.5</sub>	Second trimester	-37.64 (-129.82, 54.54)	0.02 (-0.38, 0.42)	1.62 (0.37, 7.05)	1.34 (0.24, 0.24)	1 (0.46, 2.20)	0.94 (0.41, 2.15)
	Third trimester	4.2 (-97.55, 105.95)	-0.17 (-0.62, 0.27)	0.92 (0.20, 4.17)	ق · ق 0.92 (0.2 <b>≥</b> , 4 <b>3</b> )6)	1.29 (0.53, 3.13)	0.83 (0.33, 2.06)
	Total pregnancy	8.01 (-56.57, 72.59)	0.02 (-0.27, 0.30)	0.77 (0.31, 1.91)	0.62 (0.2 <b>3</b> , 1 <b>5</b> 6)	1.15 (0.65, 2.02)	0.84 (0.45, 1.57)
Estimated	90 days prior to conception	-1.03 (-54.75, 52.68)	-0.04 (-0.27, 0.20)	0.84 (0.37, 1.91)	بق ع. 1.04 (0.4 <mark>%</mark> , 2 <mark>%</mark> 2)	1.31 (0.82, 2.10)	1.45 (0.88, 2.38)
exposure	First trimester	-9.78 (-63.78, 44.22)	0.04 (-0.19, 0.28)	0.9 (0.41, 2.00)	1.03 (0.4 <u>3</u> , 2.32)	1.21 (0.76, 1.92)	1.57 (0.94, 2.62)
to NO <sub>2</sub>	Second trimester	-20.82 (-71.18, 29.54)	-0.06 (-0.28, 0.16)	1.31 (0.61, 2.81)	1.34 (0.6 <b>2</b> , 2.38)	1.21 (0.78, 1.89)	1.36 (0.85, 2.19)
	Third trimester	-9.5 (-70.66, 51.65)	-0.01 (-0.28, 0.26)	0.79 (0.32, 1.97)	0.95 (0.3 <del>0</del> <b>0</b> , 2 <b>0</b> 3)	1.42 (0.83, 2.42)	1.51 (0.84, 2.71)
	Total pregnancy	-8.45 (-46.96, 30.06)	0 (-0.17, 0.17)	0.97 (0.54, 1.74)	المجادة 1.04 (0.5%, 1%)	1.2 (0.86, 1.69)	1.33 (0.92, 1.91)

Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

# and BSID scores (adjusted models)

		Mean dit	ference
		MDI 99% CI	PDI 99% CI
F	Per IQR increase in	(N=946)	(N=946)
Estimated	90 days prior to conception	-1.98 (-6.21, 2.25)	-6.15 (-9.69, -2.61)
exposure	First trimester	-1.66 (-6.08, 2.76)	-2.11 (-5.84, 1.62)
to PM <sub>2.5</sub>	Second trimester	3.79 (-0.08, 7.66)	3.76 (0.49, 7.02)
	Third trimester	-2.73 (-7.01, 1.55)	-1.37 (-4.99, 2.26)
	Total pregnancy	-0.27 (-2.99, 2.45)	0.23 (-2.07, 2.53)
Estimated	90 days prior to conception	-0.72 (-2.98, 1.55)	-2.83 (-4.72, -0.93)
exposure	First trimester	0.59 (-1.68, 2.86)	-1.91 (-3.83, 0.00)
to NO <sub>2</sub>	Second trimester	0.56 (-1.56, 2.68)	-0.75 (-2.54, 1.04)
	Third trimester	0.51 (-2.06, 3.09)	-1.92 (-4.09, 0.26)
	Total pregnancy	0.41 (-1.22, 2.03)	-1.15 (-2.52, 0.21)

Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11-14 weeks' gestation, primiparity, monthly household income level, and season of births.

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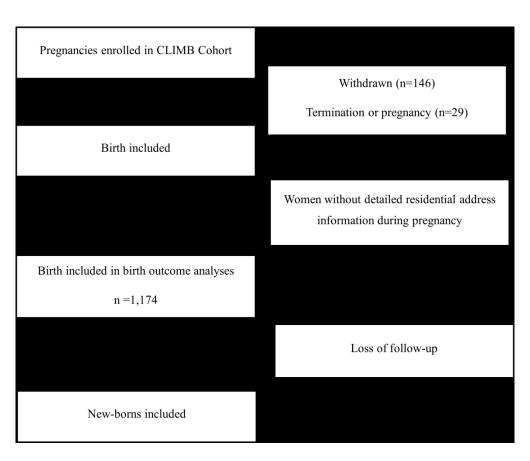


Figure 1 Flow chart of the study population in CLIMB  $441\times367$ mm (130 x 130 DPI)

Figure 2 Study area and location of monitoring sites (OpenStreetMap contributors, 2015; https://data.nextgis.com/en/region/CN-50/).

402x284mm (87 x 87 DPI)

Associations of early pregnancy air pollution with adverse birth outcomes and infant neurocognitive development in the Complex Lipids in Mothers and Babies (CLIMB) cohort

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# **Supplement**

eTable 1 Mental Development Index (Chinese version)

智力量表

(※可偶尔观察到)

序号	月龄	条目	计分
1	0.1	对铃声反应	
2	0.1	抱起时安静	
3	0.1	对摇鼓声反应	
4	0.1	对尖声反应:(电灯开关)	
5	0.1	短暂地注视红环	<b>/</b> ,*
6	0.2	短暂地注视人	
7	0.4	稍长时间地注视红环	
8	0.5	眼的水平协调活动 ( 红环 )	
9	0.7	眼的水平向天活动 (光)	
10	0.7	眼睛追随移动的人	
11	0.7	对说话声反应	
12	0.8	眼的垂直协调活动 (光)	
13	0.9	发声一至两次	
14	1	眼的垂直协调活动 ( 红环 )	
15	1.2	眼的旋转协调活动(光)	
16	1.2	眼的旋转细条活动 ( 光环 )	

17	1.3	※自由环视周围		
18	1.5	社交笑:测试者谈话与微笑时		
19	1.6	眼转向红环		
20	1.6	眼转向光		
21	1.6	※发声至少四次		
22	1.7	期待性兴奋		
23	1.7	对面部的纸有反应		
24	1.9	能用视觉辨认母亲		
25	1.9	社交笑:测试者微笑与安静时		
26	2	※对测试者的微笑和说话有发声反应		
27	2.1	※用眼睛寻找声源(详细说明)		
28	2.2	※发出两种不同的声音	1 . •	
29	2.2	对手的遮蔽眨眼		
30	2.2	对面孔的消失有反应		4.
31	2.4	注视方木		
32	2.6	从一物转看另一物		Up.
33	2.6	眼睛追随铅笔		
34	2.7	对抱起有预感性的调节反应		
35	2.9	目光追随横过桌面的球		
36	2.9	头追随悬摆的环		
37	3.1	头追随逐渐消失的勺子		
38	3.2	操作红环		
39	3.3	简单地玩摇鼓		

40	3.4	※轻轻地抚摸桌沿	
41	3.4	※意识到陌生环境	
42	3.5	头转向铃声	
43	3.6	头转向摇鼓声	
44	3.6	※手碰手的玩耍	
45	3.6	将红环送入口中	
46	3.7	伸手够悬环	
47	3.8	看自己的手	
48	4.2	接近悬环 (优势手)	
49	4.4	※发声时的姿态(描述)	
50	4.4	※主动抚摸桌沿	
51	4.4	接近镜像	. •
52	4.4	注意小糖丸	
53	4.6	伸手取方木	
54	4.7	喜欢嬉戏	
55	4.9	伸手时眼手协调	
56	4.9	拾起方木 ( 优势手 )	
57	5	保持两块方木	
58	5	持久地看红环	
59	5	头部跟着掉下的勺转动	
60	5	探索性地玩纸	
61	5	对镜像微笑	
62	5	坚持够东西	

63	5.1	在小床内重新找到摇鼓	
64	5.1	※辨别生人	
65	5.4	举起倒扣的茶杯	
66	5.5	※敲打玩耍	
67	5.5	探索性地玩细绳	
68	5.5	伸手取第二块方木	
69	5.6	※由一手向另一手传递物体	
70	5.8	※对产生响声感兴趣	
71	5.9	灵巧而直接地拾起方木	
72	6	※对镜像开玩笑	
73	6	用把柄举起茶杯	
74	6	寻找掉落的勺子	. •
75	6.1	牵拉细绳获取红环	10.
76	6.1	保留三块方木中的两块	- W
77	6.6	※发出四个不同的音节	
78	6.8	能配合玩游戏	
79	7	恰当地牵拉细绳获取红环	
80	7.1	玩摇铃,对细节感兴趣	
81	7.4	企图获得三块方木	
82	7.4	有目的地摇铃	
83	7.5	※选择性地倾听熟悉的词语	
84	8	※对 da-da 或类同词	
85	8.1	暴露玩具	

86	8.2	注意测试者的乱写	
87	8.3	将手指插入桩板洞中	
88	8.6	观看书中图画	
89	8.9	对他人的言语要求有反应	
90	9.1	拿起茶杯获得方木	
91	9.8	寻找盒子里面的东西	
92	10.3	遵照命令将方木放入茶杯 ( 放入数 )	
93	10.7	企图模仿乱写	
94	10.8	模仿用勺子搅拌	
95	10.9	遵照命令停止	
96	10.9	推动小汽车	
97	11	模仿地拍打哨娃	
98	11.1	※重复引入发笑的把戏	
99	11.2	解开裹着的方木	
100	11.2	将三块方木放入杯中	
101	11.4	※快速而不清的表达	
102	11.4	揭开兰盒子的盖	
103	11.5	翻开书页	
104	11.5	摇晃悬环的	
105	11.8	将骰子放入盒中(6个)	
106	12	恰当地握持画笔	
107	12.2	模仿说单词 (记录用过的词 )	
108	12.4	重复地插一根桩钉	

109	12.5	用手势表达想要的东西	
110	12.9	自动乱写	
111	12.9	能说两个词	
112	13	搭两层塔	
113	13.1	出示鞋子或其他衣服或自己的玩具	
114	13.2	从瓶中移出小糖丸	
115	13.3	掺九块方木放入杯中	
116	14.3	※盖上圆盒	
117	14.4	兰色模板:放置一个圆形模块	
118	14.8	用棍子够取玩具	
119	15.4	搭三层塔	
120	15.7	在 70 秒钟内插完桩钉	1
121	16.1	指出娃娃身体的各部分:三个部位以上	1/6
122	16.3	粉红模板:放置圆形模块	
123	16.6	兰色模板:放置两个圆形模块	
124	17.2	用笔模仿画一划	
125	17.5	在 42 秒钟内插完桩钉	
126	17.6	说出一物名	
127	17.7	对娃娃执行指令(在通过部位打钩:椅、杯、鼻)	
128	18.1	用语言表达要求	
129	18.6	不用于一划的乱写?	
130	18.8	兰色模板:放置两个圆快和方块	
131	18.8	指出三幅画	

		ВМЈ Ор	en	
132	19.1	能说两个单词的句子		
133	19.2	说出一副画名		
134	19.2	说出两幅画名		
135	19.3	找出两物		
136	19.8	在 30 秒钟内插完桩钉		
137	20.4	粉红模板:完成		
138	20.4	搭六层塔		
139	20.5	兰色模板,放置六个模块		
140	21	指出五副画		
141	21.1	说出三物名		
142	21.2	勉强合格地安装破娃娃		
143	21.2	区别两物:杯、盘、盒	1.	
144	22.8	辨认钟表:第四张图 1,2,3,4,5		
145	22.9	说出三幅画名		11.
146	23.8	粉红模板(反转)		
147	24.3	近似地安装破娃娃		
148	24.6	区别三物:杯、盘、盒		
149	24.7	兰色模板,在150秒钟内完成		
150	25	搭八层塔		
151	25.1	指出七副画		
152	25.1	用方木搭火车		
153	25.7	说出五副画名		
154	26.3	模仿笔划:垂直线和水平线		

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155	27.1	辨认钟表:第2张图
156	27.6	理解两个方位词
157	28	在 22 秒钟内插完桩钉
158	28.5	兰色模板:90 秒钟内完成
159	29.5	折纸
160	29.6	兰色模板:60秒钟内完成
161	30+	正确安装破娃娃
162	30+	"—"的概念
163	30+	理解三个方位词

eTable 2 Psychomotor Development Index (Chinese version)

# 运动量表

## (※可偶尔观察到,△可在施测智力量表时观察到)

序号	月龄	条目	计分
1	0.1	抱起靠肩时抬头	
2	0.1	抱起靠肩时调整姿势	
3	0.1	侧头	
4	0.1	爬起	
5	0.8	△保留红环	
6	0.8	※伸臂玩耍	
7	0.8	※踢腿玩耍	11
8	0.8	头起竖起:垂直位	16
9	1.6	头部稳定地竖起	
10	1.7	抬头(背悬位)	
11	1.8	由侧卧转向仰卧	
12	2.2	在俯卧位时用双臂撑起自己	
13	2.2	支撑下坐起	
14	2.5	保持头部稳定	
15	2.6	※双手张开占优势	
16	3.3	头平衡	
17	3.4	※尺侧一手掌抓握方木	

18	3.5	轻度支撑坐位	
19	4.3	※由仰卧转向侧卧	
20	4.7	努力想坐起	
21	5.0	部分的拇指相对(桡侧一手掌)拾起方木	
22	5.1	独坐片刻	
23	5.1	※单手抽取	
24	5.2	※转腕	
25	5.2	牵拉坐起	
26	5.6	△试图获取小糖丸	
27	5.7	独立 30 秒钟或以上	
28	5.8	由仰卧转向俯卧	
29	6.2	稳定地独坐	•
30	6.5	独坐时协调好	
31	6.6	※舀起小糖丸	
32	6.6	△完全的拇指相对拾起方木	
33	7	早期跨步运动	
34	7.5	牵拉站起	
35	7.6	※不完全的拇指相对抓糖丸	
36	7.6	走路之前的行进方式(俯卧、手膝、手足、其他)	
37	8.3	使两个勺子或方木在中线相碰	
38	8.5	跨步运动	
39	8.6	自己坐起	
40	8.6	借助家具站起	

	ĺ		1
41	8.9	精细地抓糖丸(灵巧地钳夹)	
42	9.6	拍手(中线技巧)	
43	9.8	坐下	
44	10	扶助下行走	
45	11.1	独站	
46	12	投球	
47	12.1	独走	
48	12.4	起立!	
49	13.2	扶助下右足独站	
50	13.7	扶助下左足独站	
51	14.1	侧身走	
52	14.5	扶助上楼梯	
53	14.7	倒退走	1/6
54	15.1	扶助下楼梯	
55	17.6	试图站在行木上	
56	18.7	左足独站	
57	19.3	单足踏在行木上走	
58	19.9	起立Ⅱ	
59	20.1	右足独站	
60	21.1	走直线:大致方向	
61	23.1	行木:双足站立	
62	24	踮脚走几步	
63	24.3	独自上楼梯:双足	

		BMJ Open	
64	24.4	双足跳离地面	
65	25.3	独自下楼梯	
66	25.6	行木:企图跨步	
67	25.6	倒行两米半	
68	25.7	自第一级台阶下跳下	
69	29.2	自第二级台阶下跳下	
70	29.8	踮脚走两米半	
71	29.9	跳远: 10 至 35cm (记录距离)	
72	30+	起立: Ш	
73	30+	上楼梯:双足交替向前	
74	30+	行木:交替步伐走部分路程	
75	30+	保持双足走在直线上(两米半)	
76	30+	跳远:35cm 至 60cm	16
77	30+	跳过:5cm 高的绳子	
78	30+	跳远:60cm 至 85cm	
79	30+	独脚跳两次以上	<b>U</b>
80	30+	下楼梯:双足交替向前	
81	30+	跳过 20cm 高的绳子	

BMJ Open Sop							
`PM2 5 and N	O2 exposure le	evel in 90 days prior	to concention, each t	₹ Å	d T3) and combined	across	
		over mose and prior	to conception, each	475 on 2 Jul	a 15) ana comomea		
			Estimated ex	sposure (μg/mg) 20			
N	Minimum	25th percentile	$Mean \pm SD$		75th percentile	Maximum	
	<u> </u>			wnloa of text			
1,174	38.17	44.00	$52.91 \pm 10.99$	48.43 <b>a a a a a a a a a a</b>	62.06	80.53	
1,174	37.26	43.77	$52.07 \pm 10.98$	47.26 1 7 31	61.08	82.41	
1,174	38.46	47.57	$58.64 \pm 12.21$	57.97	67.19	90.02	
1,174	37.03	47.25	$61.83 \pm 16.04$	58.82 ± 28.7	75.95	96.48	
1,174	46.69	54.85	$57.48 \pm 3.97$	57.3 1 5 76	60.61	66.98	
			10.	ning,			
1,174	25.86	45.49	$49.59 \pm 6.34$	49.94 <b>8</b> 7	53.76	70.48	
1,174	20.81	44.60	$48.8 \pm 6.43$	48.92 <b>5</b> ± 8 <b>5</b> 51	53.10	69.31	
1,174	28.93	47.18	$50.98 \pm 6.23$	51.20 ± 7 = 7 = 7 = 7 = 7 = 7 = 7 = 7 = 7 = 7	54.90	70.42	
1,174	20.57	47.20	$51.79 \pm 6.78$	52.45 ± 9.47	56.67	75.12	
1,174	27.50	47.89	$50.52 \pm 5.08$	50.4 🚉 5 🕻 1	53.40	67.53	
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	N 1,174 1,174 1,174 1,174 1,174 1,174 1,174 1,174 1,174 1,174	N Minimum  1,174 38.17 1,174 37.26 1,174 38.46 1,174 37.03 1,174 46.69  1,174 25.86 1,174 20.81 1,174 28.93 1,174 20.57 1,174 27.50	PM2.5 and NO2 exposure level in 90 days prior (WP) (n = 1,174)  N Minimum 25th percentile  1,174 38.17 44.00 1,174 37.26 43.77 1,174 38.46 47.57 1,174 37.03 47.25 1,174 46.69 54.85  1,174 25.86 45.49 1,174 20.81 44.60 1,174 28.93 47.18 1,174 20.57 47.20 1,174 27.50 47.89	PM2.5 and NO2 exposure level in 90 days prior to conception, each and the example of the exampl	PM2.5 and NO2 exposure level in 90 days prior to conception, each trimester ( $\frac{1}{100}$ ) $\frac{1}{100}$	PM2.5 and NO2 exposure level in 90 days prior to conception, each trimester (1814), \$2.5, and T3) and combined (WP) (n = 1,174)    Estimated exposure (1923)   \$2.5, and T3) and combined (WP) (n = 1,174)   \$2.5 \text{the percentile}    Mean \(\pm \) SD   Media (1.5)   \$2.5 \\\ \pm \)   \$2.5 \\\ \pm \)   \$4.00   \$5.2.91 \(\pm \)   \$1.0.99   \$48.43 \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	

e 45 of 58				В	MJ Open		/bmjope				
	eTable 4 Pearson's correlati	ons of PM <sub>2.5</sub> and	$\mathrm{NO}_2$ betwee	en each of th	e five differe	nt pregnancy	n-2023-082475 on 2 July 2024. D Enseigner yright, including for uses related perioding for uses related	N = 1,174)			
Estima	ted exposure to			PM <sub>2.5</sub>			seign s relat		NO <sub>2</sub>		
		90 days prior to conception	First trimester	Second trimester	Third trimester	Total pregnancy	90 days potext and data mining, Al training, and similar 0.554 0.334	First trimester	Second trimester	Third trimester	Total pregnancy
PM <sub>2.5</sub>	90 days prior to conception	1	1				aded uperie tt and	I			
	First trimester	-0.065	1				from eur (#   data				
	Second trimester	-0.779	-0.2012	1			MES BES min				
	Third trimester	0.288	-0.7613	-0.1688	1		) · ing, /				
	Total pregnancy	-0.534	-0.2709	0.6838	0.3858	1	Jopen VI tra				
NO <sub>2</sub>	90 days prior to conception	0.6383	0.0684	-0.4588	0.3714	0.0376	1 in 1.bm				
	First trimester	0.1537	0.6352	-0.0159	-0.4927	-0.0633	0.554	1			
	Second trimester	-0.431	0.0714	0.6269	-0.0133	0.7251	0.3345	0.5399	1		
	Third trimester	0.3027	-0.5213	0.0528	0.6817	0.4432	0.714 <b>%</b>	0.2159	0.5145	1	
	Total pregnancy	0.0057	0.0781	0.2862	0.0737	0.4779	0.678 <b>gologi</b>	0.7435	0.8755	0.7331	1
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el	Table 5 Associations between				rse birthæutkomes		ls)
		Mean d	lifference		for 2 Odd	l ratios	
		Birth weight, grams	Birth length, cm	PTB (case: 33)	Odd Enseignemen LBW (Grelated to 99% CIG	LGA (case: 108)	SGA (case: 84)
		99% CI	99% CI	99% CI	99% CIde to 1999 1999 1999 1999 1999 1999 1999 1	99% CI	99% CI
]	Per IQR increase in	(N=1,165)	(N=1,149)	(N=1,174)	to winload (N=1,174xt) xt;	(N=1,174)	(N=1,174)
Estimated	90 days prior to conception	9.28 (-44.04, 62.60)	-0.09 (-0.32, 0.14)	0.98 (0.46, 2.09)	and erie 2582)	1.2 (0.79, 1.81)	0.98 (0.60, 1.58)
exposure	First trimester	21.95 (-29.14, 73.04)	0.14 (-0.09, 0.36)	0.98 (0.48, 2.02)		0.97 (0.64, 1.47)	0.78 (0.48, 1.27)
to PM <sub>2.5</sub>	Second trimester	-18.21 (-70.25, 33.84)	0.04 (-0.19, 0.27)	0.85 (0.40, 1.79)	0.61 (0.26, <u>B</u> 40)	0.92 (0.60, 1.40)	1.33 (0.84, 2.10)
	Third trimester	-37.38 (-95.31, 20.56)	-0.32 (-0.57, -0.07)	1.35 (0.61, 2.99)	1.51 (0.56, 547)	1.08 (0.68, 1.72)	1.12 (0.67, 1.88)
Estimated	Total pregnancy  90 days prior to conception	-20.02 (-66.93, 26.89)	-0.1 (-0.30, 0.10)	0.81 (0.42, 1.55)	0.69 (0.35, 137)	1.00 (0.69, 1.46)	1.2 (0.78, 1.84)
exposure	First trimester	-13.23 (-55.66, 29.19)	-0.12 (-0.31, 0.06)	1.2 (0.65, 2.19)	1.62 (0.55, 210)	1.21 (0.85, 1.70)	1.24 (0.84, 1.83)
to NO <sub>2</sub>	Second trimester	0.3 (-42.65, 43.25)	0.08 (-0.11, 0.27)	1.01 (0.55, 1.85)	1.15 (0.61, 217)	1.17 (0.83, 1.66)	1.27 (0.86, 1.87)
<b>30</b> 1 ( <b>3</b> 2	Third trimester	-22.85 (-63.02, 17.32)	-0.04 (-0.22, 0.13)	1.11 (0.63, 1.95)	1.08 (0. <b>5</b> 0, <b>1</b> 95)	1.06 (0.77, 1.47)	1.46 (1.01, 2.11)
	Total pregnancy	-32.72 (-78.00, 12.57)	-0.16 (-0.36, 0.03)	1.13 (0.60, 2.16)	1.35 (0.88, 269)	1.24 (0.86, 1.80)	1.58 (1.03, 2.42)
	r -0	-16.58 (-51.79, 18.63)	-0.03 (-0.18, 0.12)	1.03 (0.63, 1.69)	1.13 (0.67, 🖁 91)	1.16 (0.87, 1.55)	1.44 (1.04, 2.00)

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e'	Table 6 Associations between	en PM <sub>2.5</sub> and NO <sub>2</sub> exposur	re in different pregna	ancy periods and adv	/erse birthædutkomes	(co-exposure mode	els)
		Mean diffe	erence		g for Odd	ratios	
		Birth weight, grams	Birth length, cm	PTB (case: 33)	LBW (cass 5)	LGA (case: 108)	SGA (case: 84)
		99% CI	99% CI	99% CI	2024. E seignen s relate	99% CI	99% CI
F	Per IQR increase in	(N=941)	(N=927)	(N=945)	N=945) N=945) N=945)	(N=945)	(N=945)
Estimated	90 days prior to conception	75.00 (-36.61, 186.61)	0.23 (-0.25, 0.71)	0.21 (0.04, 1.03)	0.41 (0.10 and 0.41)	1.14 (0.43, 3.03)	1.18 (0.36, 3.87)
exposure	First trimester	19.59 (-99.85, 139.04)	0 (-0.52, 0.52)	0.97 (0.17, 5.55)	0.66 (0.0 (0.0 (0.0 (0.0 (0.0 (0.0 (0.0	0.67 (0.25, 1.79)	0.73 (0.21, 2.61)
to PM <sub>2.5</sub>	Second trimester	-25.62 (-129.13, 77.90)	0.08 (-0.36, 0.53)	1.34 (0.25, 7.29)	0.94 (0.13)	0.83 (0.34, 2.01)	0.69 (0.27, 1.75)
	Third trimester	13.77 (-99.47, 127.00)	-0.2 (-0.69, 0.29)	1.12 (0.20, 6.12)	0.94 (0.18 4.89)	1 (0.38, 2.68)	0.57 (0.21, 1.56)
	Total pregnancy	21.13 (-54.55, 96.81)	0.02 (-0.31, 0.35)	0.73 (0.26, 2.07)	0.52 (0.18 in 1.48)	0.98 (0.51, 1.88)	0.55 (0.27, 1.15)
Estimated	90 days prior to conception	-18.63 (-78.33, 41.07)	-0.09 (-0.35, 0.17)	1.24 (0.49, 3.11)	1.3 (0.52, 3).29	1.27 (0.74, 2.16)	1.39 (0.79, 2.46)
exposure	First trimester	-14.53 (-75.84, 46.78)	0.05 (-0.22, 0.31)	0.91 (0.36, 2.28)	1.14 (0.44 <u>8</u> .2.98)	1.33 (0.78, 2.27)	1.70 (0.93, 3.11)
to NO <sub>2</sub>	Second trimester	-14.46 (-71.01, 42.09)	-0.08 (-0.32, 0.17)	1.22 (0.51, 2.91)	1.36 (0.55 3.36)	1.27 (0.77, 2.10)	1.50 (0.88, 2.55)
	Third trimester	-13.13 (-81.19, 54.93)	0.04 (-0.25, 0.34)	0.77 (0.28, 2.13)	0.97 (0.3 6 2.8)	1.41 (0.78, 2.57)	1.77 (0.92, 3.40)
	Total pregnancy	-15.02 (-60.15, 30.11)	0 (-0.20, 0.19)	1.08 (0.55, 2.12)	1.28 (0.64, 2.5)	1.21 (0.81, 1.80)	1.60 (1.03, 2.48)

Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

Mean	difference
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<sup>5</sup> 58			BMJ Open  BMJ open	/bmjopen-2023
eTable ´	7 Associations between PM <sub>2.5</sub> and	nd NO <sub>2</sub> exposure in differen	pregnancy periods and BSID scores	an Sijusted models)
		Mean d	fference of	5 on 2 July 2024. Downloaded from http://bmjopen.bmj.com/ on June 11, 2025 at A
		MDI 99% CI	PDI 99% CI	July 20
	Per IQR increase in	(N=946)	(N=946)	024. D
Estima	ted 90 days prior to conception	-3.54 (-5.94, -1.13)	-3.42 (-5.44, -1.40)	oownic
exposu	re First trimester	-1.07 (-3.51, 1.37)	0.04 (-2.01, 2.10)	oaded uperic
to PM	Second trimester	4.21 (1.87, 6.56)	2.63 (0.65, 4.61)	from (Al
	Third trimester	-1.43 (-4.03, 1.17)	-1.76 (-3.94, 0.42)	http://
	Total pregnancy	1.64 (-0.43, 3.71)	0.5 (-1.24, 2.25)	• • • • • • • • • • • • • • • • • • •
Estima	ted 90 days prior to conception	-1.90 (-3.82, 0.02)	-2.86 (-4.46, -1.26)	en.bm
exposu	re First trimester	-0.08 (-2.05, 1.90)	-1.17 (-2.83, 0.48)	j.com
to NC	Second trimester	1.81 (-0.03, 3.66)	0 (-1.56, 1.55)	on Ju
	Third trimester	0.04 (-2.04, 2.11)	-1.97 (-3.70, -0.23)	une 11
	Total pregnancy	0.67 (-0.95, 2.28)	-1.08 (-2.44, 0.28)	, 2025
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Mean	difference	
viean	annerence	

			BMJ Open	/bmjopen	Page 50 o
eTable 8 Asso	ociations between PM <sub>2.5</sub> and	d NO <sub>2</sub> exposure in differ	BMJ Open ent pregnancy periods and BSI an difference	ight, 2023 1D scores to 22 xposure mod	lels)
		Mea	n difference	5 on 2	
		MDI 99% CI	PDI 99% CI	July 2 Ens uses	
Pe	er IQR increase in	(N=946)	(N=946)	2024. I eigner relate	
Estimated	90 days prior to conception	-1.73 (-6.43, 2.98)	-4.74 (-8.67, -0.81)	Down! nent S d to te	
exposure to	First trimester	-2.84 (-7.85, 2.18)	-0.45 (-4.68, 3.77)	oaded Superio ext and	
PM <sub>2.5</sub>	Second trimester	4.19 (-0.15, 8.53)	5.51 (1.86, 9.16)	from from data	
	Third trimester	-3.84 (-8.60, 0.93)	0.04 (-3.99, 4.06)	http:// BES) minin	
	Total pregnancy	-0.85 (-4.04, 2.33)	1.69 (-0.99, 4.37)	bmjop g, Al t	
Estimated	90 days prior to conception	-0.31 (-2.83, 2.20)	-1.72 (-3.82, 0.38)	n 2 July 2024. Downloaded from http://bmjopen.bmj.com/ on June 11, 2025 at A Enseignement Superieur (ABES) . for uses related to text and data mining, Al training, and similar technologies.	
exposure to	First trimester	1.28 (-1.30, 3.86)	-1.80 (-3.98, 0.37)	ij.com g, and	
$NO_2$	Second trimester	-0.48 (-2.85, 1.90)	-2.11 (-4.11, -0.12)	on Ju	
	Third trimester	1.52 (-1.34, 4.38)	-1.92 (-4.34, 0.49)	ine 11	
	Total pregnancy	0.67 (-1.23, 2.57)	-1.68 (-3.28, -0.08)	, 2025 nolog	
Models adjus	ted for maternal age at enrolm	nent, infant's sex, maternal	BMI at 11–14 weeks' gestation,		
primiparity, n	nonthly household income lev	el, and season of births.		ence	
				Biblio	
				graph	
				gence Bibliographique de l	
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		Total Effect	P value	Direct Effect	Poo	Indirect Effect	P value
Estimated	90 days prior to conception	-3.54	0.000	-3.543	<b>30</b>	0.007	0.745
exposure to	First trimester	-1.07	0.259	-1.083		0.013	0.715
PM2.5	Second trimester	4.21	0.000	4.239	0 20 20 20 20 20 20 20 20 20 20 20 20 20	0.024	0.599
	Third trimester	-1.43	0.157	-1.422	0 <u>3</u> 13.86	-0.006	0.779
	Total pregnancy	1.64	0.042	1.65	o <sup>হ্ন</sup> তুর্বুব্রে	-0.012	0.695
Estimated	90 days prior to conception	-1.90	0.011	1.896		0.007	0.799
exposure to	First trimester	-0.08	0.921	0.067	0 <u>3</u> 33	0.009	0.744
NO2	Second trimester	1.81	0.011	1.844	0 <b>1</b> 010	-0.03	0.596
	Third trimester	0.04	0.962	0.059	0 4 2 4	-0.021	0.728
	Total pregnancy	0.67	0.288	0.686	0 <b>2</b> 27 <b>4</b>	-0.021	0.672
					10		

eTable 10 Mediation effect of SGA on the associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposure in different pregnancy periods and PDI scores

		Total Effect	P value	Direct Effect	Povaline	Indirect Effect	P value
Estimated	90 days prior to conception	-3.422	0.000	-3.421	0 <u>2</u> 0₽€	0.001	0.931
exposure to	First trimester	0.045	0.955	0.049	2024 000 000 000 000 000	-0.005	0.872
PM2.5	Second trimester	2.632	0.001	2.631	0 <u>5</u> 05 B	0.000	0.997
	Third trimester	-1.758	0.038	-1.762		0.003	0.849
	Total pregnancy	0.504	0.456	0.501	0 <u>545</u> 8	0.003	0.894
Estimated	90 days prior to conception	-2.862	0.000	-2.869		0.007	0.762
exposure to	First trimester	-1.174	0.067	-1.179	0306	0.005	0.826
NO2	Second trimester	-0.003	0.996	-0.010	0 8 8	0.007	0.872
	Third trimester	-1.966	0.004	-1.985	0 200 200	0.018	0.711
	Total pregnancy	-1.079	0.041	-1.092	0 <del>2</del> 03	0.013	0.751

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Complete this checklist by entering the page numbers from your manuscript where readers will find each of the items listed below.

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Page

Reporting Item Number

## Title and abstract

Title #1a Indicate the study's design with a commonly used term in the title or the abstract

Abstract	<u>#1b</u>	Provide in the abstract an informative and balanced	2, 3	
		summary of what was done and what was found		
Introduction				
Background /	<u>#2</u>	Explain the scientific background and rationale for the	4, 5	770
rationale		investigation being reported		ected
Objectives	<u>#3</u>	State specific objectives, including any prespecified	5	Protected by copyright
		hypotheses		yrign
				_
Methods				ncluding for uses
Study design	<u>#4</u>	Present key elements of study design early in the paper	5,6	or use
				s related
Setting	<u>#5</u>	Describe the setting, locations, and relevant dates,	5,6	ed to
		including periods of recruitment, exposure, follow-up, and		eria
		data collection		ind da
Eligibility criteria	<u>#6a</u>	Give the eligibility criteria, and the sources and methods of	5,6	data minin
		selection of participants. Describe methods of follow-up.		9, A
				craining,
Eligibility criteria	<u>#6b</u>	For matched studies, give matching criteria and number of	6	
		exposed and unexposed		a simi
Variables	<u>#7</u>	Clearly define all outcomes, exposures, predictors,	6, 7, 8	and similar technologies
		potential confounders, and effect modifiers. Give diagnostic		nolog
		criteria, if applicable		les.
Data sources /	<u>#8</u>	For each variable of interest give sources of data and	6, 7, 8	
measurement		details of methods of assessment (measurement).		
		Describe comparability of assessment methods if there is		
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		BMJ Open	Pag	e 56 of 58
		more than one group. Give information separately for for		BMJ
		exposed and unexposed groups if applicable.		Open: f
Bias	<u>#9</u>	Describe any efforts to address potential sources of bias	6, 7, 8	BMJ Open: first published as 10.1136/bmjopen-2023-082475 on 2 July 2024. Downloaded from ht Enseignement Superieur (ABI Protected by copyright, including for uses related to text and data m
Study size	<u>#10</u>	Explain how the study size was arrived at	6	shed as 10 Pro
Quantitative	<u>#11</u>	Explain how quantitative variables were handled in the	8, 9	).1136/ tected
variables		analyses. If applicable, describe which groupings were		bmjop by cop
		chosen, and why		s 10.1136/bmjopen-2023-082475 on 2 July 2024. Do Enseignem Protected by copyright, including for uses related
Statistical	<u>#12a</u>	Describe all statistical methods, including those used to	8, 9	-08247! includi
methods		control for confounding		5 on 2 . ng for
				July 20: Enseiç uses re
				24. Dov gneme slated t
Statistical	<u>#12b</u>	Describe any methods used to examine subgroups and	8, 9	wnloac nt Sup o text
methods		interactions		led from erieur and da
Statistical	<u>#12c</u>	Explain how missing data were addressed	8, 9	=. m <b></b>
methods				//bmjo ) ing, Al
Statistical	<u>#12d</u>	If applicable, explain how loss to follow-up was addressed	8, 9	pen.brr trainin
methods				nj.com/ o g, and s
Statistical	<u>#12e</u>	Describe any sensitivity analyses	8, 9	p://bmjopen.bmj.com/ on June 11, 2025 at Agence Bibliographique de l S) . ning, Al training, and similar technologies.
methods				11, 20: chnolo
				25 at Aç ogies.
Results				yence B
<i>เ ง</i> ธอนเเอ				ibliogr
Participants	<u>#13a</u>	Report numbers of individuals at each stage of study—eg	10	aphiqu
	Ear man	numbers potentially eligible, examined for eligibility,		ie de l
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		confirmed eligible, included in the study, completing follow-		
		up, and analysed. Give information separately for for		
		exposed and unexposed groups if applicable.		
Participants	<u>#13b</u>	Give reasons for non-participation at each stage	10, 20	
Participants	<u>#13c</u>	Consider use of a flow diagram	10, 20	Protected
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Descriptive data	<u>#14a</u>	Give characteristics of study participants (eg demographic,	10,22	right, i
		clinical, social) and information on exposures and potential		ncludi
		confounders. Give information separately for exposed and		ng for
		unexposed groups if applicable.		uses r
Descriptive data	#14b	Indicate number of participants with missing data for each	N/A	elated
Decempary data	<u># 1 10</u>	variable of interest	14// (	to text
		variable of interest		and d
				ata mi
Descriptive data	<u>#14c</u>	Summarise follow-up time (eg, average and total amount)	N/A	ning, A
				training, and similar technologies
Outcome data	<u>#15</u>	Report numbers of outcome events or summary measures	10,22	ıd simi
		over time. Give information separately for exposed and		lar tec
		unexposed groups if applicable.		hnolog
				gies.
	<b>!!</b> 40		10 11	
Main results	<u>#16a</u>	Give unadjusted estimates and, if applicable, confounder-	10, 11	
		adjusted estimates and their precision (eg, 95% confidence		

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		interval). Make clear which confounders were adjusted for	
		and why they were included	
Main results	<u>#16b</u>	Report category boundaries when continuous variables were categorized	10, 11
Main results	<u>#16c</u>	If relevant, consider translating estimates of relative risk	N/A
		into absolute risk for a meaningful time period	
			;
Other analyses	<u>#17</u>	Report other analyses done—eg analyses of subgroups	11
		and interactions, and sensitivity analyses	
Discussion			
Key results	<u>#18</u>	Summarise key results with reference to study objectives	11, 12
Limitations	<u>#19</u>	Discuss limitations of the study, taking into account sources	14
		of potential bias or imprecision. Discuss both direction and	
		magnitude of any potential bias.	,
Interpretation	<u>#20</u>	Give a cautious overall interpretation considering	12, 13
		objectives, limitations, multiplicity of analyses, results from	·
		similar studies, and other relevant evidence.	
Generalisability	<u>#21</u>	Discuss the generalisability (external validity) of the study	14
		results	(
Other Information	1		

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Funding #22 Give the source of funding and the role of the funders for the present study and, if applicable, for the original study on which the present article is based

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# **BMJ Open**

Associations of air pollution exposures in preconception and pregnancy with birth outcomes and infant neurocognitive development: analysis of the Complex Lipids in Mothers and Babies (CLIMB) prospective cohort in Chongqing, China

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- 1 Associations of air pollution exposures in preconception and pregnancy with birth
- 2 outcomes and infant neurocognitive development: analysis of the Complex Lipids in
- 3 Mothers and Babies (CLIMB) prospective cohort in Chongqing, China
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#### **Abstract**

**Objectives:** to investigate the associations of traffic-related air pollution exposures in early pregnancy with birth outcomes and infant neurocognitive development

**Design:** cohort study

**Setting:** eligible women attended six visits in the maternity clinics of two centres (the First Affiliated Hospital of Chongqing Medical University and Chongqing Health Centre for Women and Children).

**Participants:** women who were between 20 and 40 years of age and were at 11–14 weeks gestation with a singleton pregnancy were eligible for participation. Women were excluded if they had a history of premature delivery before 32 weeks of gestation, maternal milk allergy or aversion, or severe lactose intolerance. 1,273 pregnant women enrolled in 2015-2016 and 1,174 live births were included in this analysis.

**Exposures:** Air pollution concentrations at their home addresses, including particulate matter (PM) with diameter  $\leq$ 2.5µm (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>), during pre-conception and each trimester period were estimated using land-use regression models.

**Outcome measures:** birth outcomes (i.e., birthweight, birth length, preterm birth (PTB), low birth weight (LBW), large for gestational age (LGA) and small for gestational age (SGA) status) and neurodevelopment outcomes measured by the Chinese version of Bayley Scales of Infant Development (CBSID).

**Results:** An association between SGA and per Interquartile range (IQR) increases in NO<sub>2</sub> was found in the first trimester (Odd ratio (OR): 1.57, 95% confidence interval (CI): 1.06, 2.32) and during the whole pregnancy (OR: 1.33, 99% CI: 1.01, 1.75). Both PM<sub>2.5</sub> and NO<sub>2</sub> exposure in the 90 days prior to conception were associated with lower Psychomotor Development Index (PDI) scores ( $\beta$ : -6.15, 95% CI: -8.84, -3.46;  $\beta$ : -2.83, 95% CI: -4.27, -1.39, respectively). Increased NO<sub>2</sub> exposure was associated with an increased risk of psychomotor development delay (PDD) during different trimesters of pregnancy.

Conclusions: Increased exposure to NO<sub>2</sub> during pregnancy were associated with increased risk of SGA and psychomotor development delay, while increased exposure to both PM<sub>2.5</sub> and NO<sub>2</sub> pre-conception were associated with adverse psychomotor development outcomes at 12 months of age.

## **Article summary**

## Strengths and limitations of this study

- We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification.
- This study uniquely explored the impacts of both pre-conception and prenatal exposure to PM<sub>2.5</sub> and NO<sub>2</sub> on neurodevelopmental outcomes in young infants, within an urban environment characterized by relatively high air pollution levels.
- Our sample size was relatively small, limiting the statistical power to assess several outcomes.
- We defined exposure windows for clinically-defined trimesters.



#### Introduction

 Air pollution is a major environmental factor that has been linked to a range of adverse health outcomes in children. Maternal exposure to air pollutants during pregnancy, especially particulate matter (PM) with diameter  $\leq 2.5 \mu m$  (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>), has been found to be associated with adverse birth outcomes, including pre-term birth (PTB)(1), term low birth weight (TLBW) (2), and small for gestational age (SGA) status (3). According to the developmental origins of health and disease (DOHaD) hypothesis, prenatal exposures to air pollution may lead to adverse birth outcomes and subsequently increase the susceptibility to the development of certain diseases later in life (4). A number of epidemiological studies have linked prenatal air pollution exposure with neurodevelopmental disorders such as autism spectrum disorder (ASD), attention deficit hyperactivity disorder (ADHD), and cognitive impairment (5). Although the underlying biological mechanisms are still unclear, some studies indicated that prenatal air pollution exposure may induce systemic oxidative stress that triggers intrauterine inflammation, leading to damage to several fetal organs, including the brain (6, 7).

It is also unclear that whether the adverse effects of air pollution may start earlier before conception. Three months before conception was considered as a critical developmental window for gametogenesis. Air pollution exposure during the three-month preconception period may have adverse effects on gametogenesis of sperm (8, 9) and ova cells (10). Exposures to PM<sub>2.5</sub> in preconception period have been associated with various neurodevelopmental outcomes, such as neural tube defects (11), lower psychomotor development scores (12), higher risk of ASD (13, 14), and higher risk of intellectual disability (15). Further research is required due to inconsistencies across studies in terms of studied health outcomes and exposure levels of air pollution (12). Additionally, while there is growing evidence for the effects of preconception PM<sub>2.5</sub> exposure on the risk of adverse neurodevelopmental outcomes, no study to date has examined the effects of preconception NO<sub>2</sub> exposure. Exposure to NO<sub>2</sub> during pregnancy may be linked to compromised neural development in children, particularly affecting fine psychomotor skills(16). Studying PM<sub>2.5</sub> along with NO<sub>2</sub> may allow us to explore how multiple pollutants affect birth outcomes and infant neurocognitive development independently and jointly. Moreover, both PM<sub>2.5</sub> and NO<sub>2</sub> are regulated traffic-related air pollutants in many countries. Understanding their impacts on birth and infant neurocognitive development can provide valuable insights for policymakers and public health authorities to develop effective air quality regulations and interventions.

Many studies have reported the effects of prenatal exposure to air pollution on neurodevelopmental function in children. However, the reported associations vary, due to the heterogeneous assessments of air pollution and neurodevelopmental outcomes (5, 17).

The current study leveraged the Complex Lipids in Mothers and Babies (CLIMB) cohort, a prospective birth cohort recruited in Chongqing, China (18), with trimester-specific maternal PM<sub>2.5</sub> and NO<sub>2</sub> air pollution exposure derived from a spatio-temporal land use regression (LUR) model (19). The aim of this analysis was to examine the associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposures during pre- and during pregnancy, with birth and infant neurocognitive development outcomes at 12 months of age.

A key aspect in all studies like this one is the accuracy of documenting exposure; a recent Chinese study determined air pollution exposure based on data from the nearest monitoring station (20) may not reflect the fine temporal and spatial variability of pollutant exposures among participants. Our study employed common air pollution exposure models based on advanced geographic information systems (GIS), to address some of the limitations of previous studies (21).

In addition, the timing of exposure is also critical in determining the effects of exposure on developmental outcomes. Indeed, the evidence from previous studies on the sensitive time windows for exposure pre- and during pregnancy remains inconclusive. Some studies have indicated that the early-to-mid pregnancy phase may be a critical period in terms of the impact of air pollution on neurodevelopment(22, 23). Early pregnancy is particularly important for neurogenesis and neuromigration, making it a susceptible period (24). However, some studies reported stronger associations for middle or late pregnancy (20, 25, 26). More studies identifying critical periods are needed to enhance our understanding of how pre-conception and prenatal air pollution exposure affect neurodevelopment. With this cohort, we are able to examine the effects of exposure pre-conception, at each trimester, and the entire pregnancy.

#### Methods

#### Study population

Participant recruitment in the CLIMB cohort has been described previously (27). In brief, women who were between 20 and 40 years of age and were at 11–14 weeks gestation with a singleton pregnancy were eligible for participation. Women were excluded if they had a history of premature delivery before 32 weeks of gestation, maternal milk allergy or aversion, or severe lactose intolerance.

From September 2015 to November 2016, a total of 1,500 women were recruited into the cohort. Participants attended six visits at the First Affiliated Hospital of Chongqing Medical University and Chongqing Health Centre for Women and Children: 11–14 weeks' gestation (visit 1), 22–28 week's gestation (visit 2), 32–34 week's gestation (visit 3), at birth (visit 4), 6 weeks postnatal (visit 5), and 12 months postnatal (visit 6).

Women who withdrew from the study (n = 146), terminated their pregnancy (n = 29), miscarried (n = 12) or were lost to follow-up (n = 40) were excluded from the analysis, leaving a sample size of 1,273 women. Analyses were restricted to mothers whose detailed residential addresses during pregnancy were known (**Figure 1**). A total of 1,174 live births were thus included in the pregnancy and neonatal outcomes analysis. Subsequently, at 1 year follow-up, 946 children were included in the analysis of neurodevelopment outcomes.

## **Exposure assessment**

The address of participants was collected at the first visit. Exposure assessment based on spatiotemporal land use regression (LUR) models for PM<sub>2.5</sub> and NO<sub>2</sub> were developed for the study region. The study area focused on the urban center of the Chinese municipality of Chongqing (**Figure 2**). A description of the methodology of exposure modelling has been reported previously (19). Briefly, the models included both spatial and temporal components of exposure. PM<sub>2.5</sub> and NO<sub>2</sub> concentration data were collected from 17 routine monitoring sites operated by the Chongqing Environmental Monitoring Center in 2015-2016. For the spatial component of models, we calculated annual average concentrations of each pollutant in 2015, and fit linear regression models using five groups of geographic data (road network, land use, topography, vegetation, and population density) as spatial predictor variables. For the temporal component of models, we calculated the residuals from the spatial component at each

monitoring site on a daily basis by subtracting the predicted annual average concentration from the observed daily average concentrations measured in 2015 and 2016, and then fitted generalised additive models (GAM) using seven groups of meteorological data (temperature, amount of rainfall, rainfall events, relative humidity, horizontal visibility, wind direction and wind speed) as temporal predictor variables. The meteorological variables were used to account for the influence of weather on the change in air pollution concentration over time. To account for the remaining spatial autocorrelation, the smoothed terms of longitude and latitude were fit to spatiotemporal residuals which were calculated by subtracting the sum of the spatial temporal predictions from the measured daily average concentrations in 2015 and 2016. The performance of the PM<sub>2.5</sub> spatiotemporal models was good (Correlation (COR)-R<sup>2</sup>: 0.72) and the NO<sub>2</sub> spatiotemporal model was moderate (COR-R<sup>2</sup>: 0.39) when providing concentration estimates in absolute terms.

Combining the family address coordinates of each pregnant woman and the gestation period of the pregnancy (calculated from the date of last menstrual period to the date of delivery), we used this spatiotemporal model to estimate the average exposure of each pregnant woman in 90 days prior to pregnancy (90D), first trimester (T1), second trimester (T2), third trimester (T3) and whole pregnancy period (WP), respectively.

#### **Outcomes**

#### **Birth Outcomes**

Birth outcomes were determined by experienced obstetricians and abstracted from the medical records. Birth outcomes included: birthweight (in grams), birth length (in centimetres), PTB, low birth weight (LBW), large for gestational age (LGA) and SGA status (28). PTB was defined as delivery before 37 weeks. LBW was defined as weighing less than 2500 g at birth. LGA and SGA were indicated by birth weight greater than and less than the 90th and 10th percentile within this study for the gestational age by sex respectively (29). Term low birth weight was not considered due to a small sample size of only 8 cases.

#### **Neurodevelopment outcomes**

The Chinese version of Bayley Scales of Infant Development (CBSID) was used to assess mental and psychomotor development for infants in this study. The CBSID is appropriate for evaluation of infants from 2–30 months old (30) and takes into consideration each infant's age in days. Infants were assessed at around 12 months (range from 11 months and 15 days to 12 months and 15 days) by a trained examiner, with ages corrected for preterm birth. These scales have been formally adapted to the Chinese language and locally standardized to become culturally appropriate, with two main indexes: the Mental Development Index (MDI) and the Psychomotor Development Index (PDI). The MDI component comprised 163 items and assessed age-appropriate items related to cognitive functioning, personal and social development, and language development (see eTable 1 in the Supplement). The PDI component comprised 81 items and assessed age-appropriate fine and gross motor skills (see eTable 2 in the Supplement). The test provided raw scores for mental and psychomotor development that were converted to standardized (in terms of age in days) MDI and PDI scores, based on norms for the Chinese population. As with other forms of the Bayley test these index scores have a mean of 100, and a standard deviation of 15, with a lower score reflecting poorer performance (31). If an infant refused to cooperate with the examiners to finish the task, a second assessment was arranged within two weeks. If the infant could not cooperate at the second BSID assessment, their data were classified as missing. In addition to the continuous scores, we define mental developmental delay (MDD) and psychomotor developmental delay (PDD) if the score is less than 85(32).

#### **Covariates**

 Socio-demographic data were collected through interviews by trained nurses. The following potential confounders were identified: maternal age at enrolment (in years), infant sex (male/female), maternal BMI at 11–14 weeks' gestation (kg/m²), parity (Yes/No), monthly household income level (categorized as: <2,000 yuan, 2,000–7,000 yuan, 7,000–10,000 yuan, or >10,000 yuan), season of birth (categorized as: Spring (Mar-May), Summer (Jun-Aug), Autumn (Sep-Nov) or Winter (Dec-Feb)). Marital status (single/married) and smoking or drinking during pregnancy (Yes/no) were not taken into account in this analysis because of the homogeneity of the study population (i.e., 98.6% women were married and 99.6% women reported not smoking or drinking alcohol during pregnancy). We did not adjust dietary

supplements during pregnancy because all pregnant women routinely take folic acid in this cohort.

#### Statistical analyses

Data were described in terms of mean ± SD or median (IQR) for continuous variables, or as percentages for categorical variables. Modelled PM<sub>2.5</sub> and NO<sub>2</sub> exposure levels in 90D, T1, T2, T3 and WP were considered separately. We examined the Spearman correlation between each of the exposures in the different pregnancy periods. For birth outcomes, multivariable linear regression was used for continuous outcomes (e.g., birth weight and birth length) to estimate β coefficient and their 95% confidence intervals (CIs) and multivariable logistic regression for binary outcomes (e.g., PTB, LBW, LGA and SGA status) to estimate odds ratio (OR) and 95% CIs. For mental and psychomotor development (e.g., MDI and PDI scores), multivariable linear regression models were fit to estimate β coefficient and their 95% CIs. We also conducted multivariable logistic regression analysis for binary neurodevelopment outcomes (i.e., MDD and PDD). Models were adjusted for maternal age at enrolment, infant sex, maternal BMI at 11–14 weeks gestation, primiparity, monthly household income level, and season of birth. We also ran co-exposure models to estimate associations of one air pollutant whilst additionally adjusting for the other air pollutant (i.e., PM<sub>2.5</sub> effects in T1 adjusted for NO<sub>2</sub> in T1). Effect estimates are reported for each IQR increase of PM<sub>2.5</sub> and NO<sub>2</sub>. All analyses were performed using STATA version 17. A p-value of <0.05 was considered statistically significant to address multiple comparisons in the analyses.

### Patient and public involvement

Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

#### **Results**

## **Study Participants**

Participant characteristics are presented in **Table 1**. Of those participating women, the mean age was 28.7 years and mean BMI was 21.5 kg/m². 98.0% of women were of Han ethnicity, 77.9% were primiparous, and 67.6% had completed tertiary education. 33 (2.8%), 30 (2.6%), 108 (9.2%), 84 (7.2%) of the 1,174 births considered in this analysis were classified as PTB, LBW, LGA and SGA, respectively. For those 946 children who completed the BSID test, the mean MDI score was 94.7 (SD: 17.7) and the mean PDI score was 87.4 (SD: 14.9). The proportions of participants with MDD (MDI <85) and PDD (PDI < 85) were 27.1% and 42.4%, respectively.

## **Exposure assessment**

Median PM<sub>2.5</sub> exposure concentrations were 57.31  $\mu$ g/m³ (IQR: 5.76) and median NO<sub>2</sub> exposure levels were 50.46  $\mu$ g/m³ (IQR: 5.51) during the whole pregnancy period (eTable 3 in the Supplement). For PM<sub>2.5</sub>, the concentration in the pre-conception and T1 were considerably lower than other periods, close to 10  $\mu$ g/m³. The between-trimester and 90D values for NO<sub>2</sub> were generally moderately correlated (Pearson's r > 0.5). The correlation coefficients of PM<sub>2.5</sub> were more variable between time periods reflecting the high variability of PM<sub>2.5</sub> concentrations, with values ranging from -0.78 to +0.68. Correlations between PM<sub>2.5</sub> and NO<sub>2</sub> in the same pregnancy period were moderately correlated (Pearson's r ~0.6, eTable 4 in the Supplement).

Table 1 Characteristics of study sample in the CLIMB cohort (N = 1,174)

6Characteristic of mothers	N	n (%) / mean ± SD	Characteristic of child	N	n (%) /mean ± SD
7Maternal age (Years)	1,174	$28.7 \pm 3.5$	Gestational week (week)	1,174	$39.4 \pm 1.5$
8BMI (kg/m²)	1,174	$21.5 \pm 2.9$	Birth weight (g)	1,165	$3314.4 \pm 428.8$
9Han ethnicity (%)	1,174		Birth length (cm)	1,149	$49.7 \pm 1.9$
10 Yes 1 No		1,151 (98.0%)	New born sex	1,172	
1No		23 (2.0%)	Female		561 (47.9%)
Marital status (%)	1,174		Male		611 (52.1%)
1\$ingle		16 (1.4%)	Birth outcomes		_
1Married		1,158 (98.6%)	Preterm birth (PTB)	1,174	Pro
1 <b>8</b> rimiparity (%)	1,174		Yes		33 (2.8%) <b>§</b>
1¥es		914 (77.9%)	No		1,141 (97.2%)
18No		260 (22.1%)	Low birth weight (LBW)	1,174	33 (2.8%) 1,141 (97.2%)  30 (2.6%) 30 (2.6%) 1,141 (97.2%) 108 (9.2%)
<sup>1</sup> History of miscarriage or	1,174	· /	Yes		30 (2.6%)
<sup>20</sup> abortion (%)					· Ý
<sup>2</sup> Yes		553 (47.1%)	No		1,141 (97.2%) 🕏
2Yes 22No 23		621 (52.9%)	Large for gestational age	1,174	<u>, , , , , , , , , , , , , , , , , , , </u>
			(LGA)	,	n <u>cl</u>
24 2\$moking/drinking during	1,174		Yes		108 (9.2%) _ 트
pregnancy (%)	,				) j
2¥es		5 (0.4%)	No		1,066 (90.8%)
2 <b>%</b> Io		1,169 (99.6%)	Small for gestational age	1,174	S
29		, ,	(SGA)	,	es
3€ducation level	946		Yes		$\begin{array}{c ccccccccccccccccccccccccccccccccccc$
<sup>3</sup> Low: High school or below		306 (32.3%)	No		1,090 (92.8%)
High: College/uni or above		640 (67.6%)	BSID test	946	<u> </u>
34 35 36 35 36 36 36 36 36 36 36 36 36 36 36 36 36	946		MDI score		94.7 ± 17.7
Full-time		762 (80.5%)	PDI score		$87.4 \pm 14.9$
Housewife		82 (8.7%)	Mental development	946	D D
of thers		102 (10.8%)	Delay (MDI < 85)		276 (27.1%) <b>a</b>
3 Household income (Monthly)	946		Normal (MDI $\geq$ 85)		741 (72.9%)
392000 RMB		186 (19.7%)	Psychomotor Development	946	<u> </u>
4 <b>0</b> 000-4000 RMB		329 (34.8%)	Delay (PDI < 85)	,.0	431 (42.4%) <b>5</b>
4 <b>4</b> 000-7000 RMB		292 (30.9%)	Normal (PDI $\geq$ 85)		586 (57.6%) ≥
4₹000-10000 RMB		139 (14.7%)	Season of birth	1,174	
43		15 (1, 10)	Spring (Mar-May)	-,-, .	411 (35.01%)
44			Summer (Jun-Aug)		و (22.40%)
45			Autumn (Sep-Nov)		198 (16.87%)
46			Winter (Dec-Feb)		302 (25.72%) <u>w</u>
47					<u> </u>
48					ar
49 50					tec
51					μν
52					olo
53					411 (35.01%) aining, and 263 (22.40%) 198 (16.87%) 302 (25.72%) milar technologies
54					· ·

### **Association with birth outcomes**

- In the unadjusted models (eTable 5 in the Supplement), higher exposure concentrations of
- $PM_{2.5}$  in T3 were significantly associated with lower birth length ( $\beta$ : -0.32, 95% CI: -0.51, -
- 28 0.13; per IQR increase). We also observed increased NO<sub>2</sub> in T3 were significantly associated
- 29 with lower birth length (β: -0.16, 95% CI: -0.32, -0.01; per IQR). A risk between SGA and
- 30 increases in NO<sub>2</sub> (per IQR) was found in T2 (OR: 1.46, 95% CI: 1.10, 1.93), T3 (OR: 1.58,
- 31 95% CI: 1.14, 2.18) and in the whole pregnancy period (OR: 1.44, 95% CI: 1.13, 1.85). We
- observed no evidence of associations of NO<sub>2</sub> with overall birth weight, birth length and other
- adverse birth outcomes (e.g., PTB, LBW, and LGA).
- In the adjusted models (**Table 2**), we found increased effect size for NO<sub>2</sub> and SGA in T2
- 35 (OR: 1.57, 95% CI: 1.06, 2.32), and slightly reduced effects size for NO<sub>2</sub> and SGA in the
- whole pregnancy period (OR: 1.33, 95% CI: 1.01, 1.75) compared with the unadjusted
- 37 model. We observed no evidence of associations with birth length in the adjusted models.
- After co-adjustment for PM<sub>2.5</sub> (see eTable 6 in the Supplement), the association of NO<sub>2</sub>
- 39 with SGA was also found in T1 (OR: 1.70, 95% CI: 1.07, 2.69), T3 (OR: 1.77, 95% CI: 1.08,
- 40 2.91) and in the whole pregnancy period (OR: 1.60, 95% CI: 1.15, 2.23).

Page 15 of 48  1 2 3 41		ween PM <sub>2.5</sub> and NO <sub>2</sub> ex	BMJ Open		bmjopen-2023-08gg by copyright, incipated dverse	mes (adjusted mod	els)
4 5		Mean	difference		ding fo Odd	ratios	
7 8	Per IQR increase in	Birth weight, grams (95% CI)	Birth length, cm (95% CI)	PTB (case: 33) (95% CI)	LBW (45 (25 (25 (25 (25 (25 (25 (25 (25 (25 (2	LGA (case: 108) (95% CI)	SGA (case: 84) (95% CI)
9 10		(N=941)	(N=927)	(N=945)		(N=945)	(N=945)
1Estimated	90 days prior to conception	59.73 (-16.52, 135.98)	0.15 (-0.176, 0.48)	0.24 (0.06, 1.00)	0.49 (8.38)	1.40 (0.72, 2.71)	1.66 (0.75, 3.68)
12 exposure	First trimester	6.21 (-73.79, 86.20)	0.04 (-0.308, 0.388)	0.88 (0.28, 2.80)	0.76 (9.31 2.81)	0.86 (0.45, 1.67)	1.33 (0.58, 3.04)
	Second trimester	-37.64 (-107.73, 32.44)	0.02 (-0.283, 0.326)	1.62 (0.53, 4.96)	1.34 (3.58 4.68)	1.00 (0.55, 1.83)	0.94 (0.50, 1.76)
14to PM <sub>2.5</sub>	Third trimester	4.20 (-73.17, 81.57)	-0.17 (-0.509, 0.162)	0.92 (0.29, 2.90)	0.92 (2.85)	1.29 (0.65, 2.53)	0.83 (0.42, 1.66)
16	Total pregnancy	8.01 (-41.10, 57.11)	0.02 (-0.198, 0.230)	0.77 (0.38, 1.54)	0.62 (1.25)	1.15 (0.75, 1.77)	0.84 (0.52, 1.35)
17Estimated	90 days prior to conception	-1.03 (-41.88, 39.81)	-0.04 (-0.215, 0.139)	0.84 (0.45, 1.57)	1.04 (9.54 1.98)	1.31 (0.91, 1.88)	1.45 (0.99, 2.12)
18 19exposure	First trimester	-9.78 (-50.84, 31.28)	0.04 (-0.133, 0.223)	0.90 (0.49, 1.65)	1.03 (3.96 1.91)	1.21 (0.85, 1.72)	1.57 (1.06, 2.32)
_	Second trimester	-20.82 (-59.11, 17.47)	-0.06 (-0.222, 0.112)	1.31 (0.73, 2.34)	1.34 (2.75 2.40)	1.21 (0.86, 1.70)	1.36 (0.95, 1.95)
$\frac{20}{21}$ to $NO_2$	Third trimester	-9.50 (-56.00, 36.99)	-0.01 (-0.213, 0.191)	0.79 (0.40, 1.59)	0.95 ( 1.94)	1.42 (0.94, 2.13)	1.51 (0.97, 2.36)
22 23	Total pregnancy	-8.45 (-37.73, 20.83)	0.00 (-0.125, 0.130)	0.97 (0.62, 1.51)	1.04 ( <b>9</b> .66-1.64)	1.20 (0.93, 1.56)	1.33 (1.01, 1.75)

2½ Il significant findings in the table are bold.
2½ Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

## Association with infant neurodevelopment outcomes

- In unadjusted models, PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with
- 45 lower MDI and PDI scores in offspring (β: -3.54, 95% CI: -5.37, -1.71; β: -3.42, 95% CI: -
- 46 4.96, -1.89) (**Table 3**). We also observed an unexpected positive association between PM<sub>2.5</sub>
- 47 exposures in second trimester with MDI (β: 4.21, 95% CI: 2.43, 6.00) and PDI (β: 2.63, 95%
- 48 CI: 1.12, 4.14). Exposure to NO<sub>2</sub> was associated with lower MDI (-1.90, 95% CI: -3.36, -0.44)
- and PDI in the 90 days prior to conception (-2.86, 95% CI: -4.08, -1.65). NO<sub>2</sub> exposure was
- also associated with lower PDI scores in T3 (-1.97, 95% CI: -3.29, -0.65) and in the whole
- 51 pregnancy periods (-1.08, 95% CI: -2.11, -0.05). We did not observe any association between
- 52 NO<sub>2</sub> and MDI in any pregnancy periods.

- In the adjusted models (Table 3), we found PM<sub>2.5</sub> exposure in the 90 days prior to conception
- was associated with lower PDI scores (β: -6.15, 95% CI: -8.84, -3.46). Similarly, there was
- also a significant association of increased NO<sub>2</sub> exposure and lower PDI score in the 90 days
- 56 prior to conception (β: -2.83, 95% CI: -4.27, -1.39), T1 (β: -1.91, 95% CI: -3.37, -0.46), T3 (β:
- 57 -1.92, 95% CI: -3.57, -0.26) and whole pregnancy period (β: -1.15, 95% CI: -2.19, -0.11). The
- positive association between  $PM_{2.5}$  exposures in second trimester with PDI ( $\beta$ : 3.76, 95% CI:
- 59 1.27, 6.24) remained. We did not observe any association with MDI in any pregnancy periods.
- In the co-exposure models (**Table 3**),  $PM_{2.5}$  exposure in the 90 days prior to conception was
- associated with lower PDI scores (β: -4.74, 95% CI: -7.73, -1.75). We also observed a positive
- association between  $PM_{2.5}$  exposures in second trimester with PDI ( $\beta$ : 5.51, 95% CI:2.73, 8.28).
- Exposure to NO<sub>2</sub> was significantly associated with lower PDI in 90D (β: -1.72, 95% CI: -3.31,
- -0.12), T1 (β: -1.80, 95% CI: -3.46, -0.15), T2 (β: -2.11, 95% CI: -3.63, -0.60), T3 (β: -1.92,
- 65 95% CI: -3.76, -0.09) and whole pregnancy period (β: -1.68, 95% CI: -2.89, -0.46).

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6 Crude models		Adjusted models*		Co-exposure models**			
7 8	Per IQR increase in	MDI (95% CI) (N=946)	PDI 95% CI (N=946)	MDI (95% CI) (N=945)	PDI 95% & Huly (N=945% 22	MDI (95% CI) (N=945)	PDI 95% CI (N=945)
16 stimated	90 days prior to conception	-3.54 (-5.37, -1.71)	-3.42 (-4.96, -1.89)	-1.98 (-5.19, 1.23)	-6.15 (-8.84, 33, 46)	-1.73 (-5.30, 1.85)	-4.74 (-7.73, -1.75)
<sup>1</sup> exposure	First trimester	-1.07 (-2.93, 0.79)	0.04 (-1.52, 1.61)	-1.66 (-5.02, 1.70)	-2.11 (-4.95, <b>3</b> , <b>3</b> )	-2.84 (-6.65, 0.97)	-0.45 (-3.67, 2.76)
12 13to PM <sub>2.5</sub>	Second trimester	4.21 (2.43, 6.00)	2.63 (1.12, 4.14)	3.79 (0.85, 6.73)	3.76 (1.27, <b>6</b> 2 4 3	4.19 (0.89, 7.49)	5.51 (2.73, 8.28)
14	Third trimester	-1.43 (-3.41, 0.55)	-1.76 (-3.42, -0.10)	-2.73 (-5.99, 0.53)	-1.37 (-4.12, 🗟 🗟 🗞	-3.84 (-7.46, -0.22)	0.04 (-3.02, 3.09)
15	Total pregnancy	1.64 (0.06, 3.21)	0.5 (-0.82, 1.83)	-0.27 (-2.34, 1.80)	0.23 (-1.52, <b>1.2</b> 8)	-0.85 (-3.28, 1.57)	1.69 (-0.35, 3.73)
Estimated	90 days prior to conception	-1.90 (-3.36, -0.44)	-2.86 (-4.08, -1.65)	-0.72 (-2.43, 1.00)	-2.83 (-4.27, at 3 9)	-0.31 (-2.22, 1.60)	-1.72 (-3.31, -0.12)
1&exposure	First trimester	-0.08 (-1.57, 1.42)	-1.17 (-2.43, 0.08)	0.59 (-1.14, 2.32)	-1.91 (-3.37,	1.28 (-0.68, 3.24)	-1.80 (-3.46, -0.15)
19 20 to NO <sub>2</sub>	Second trimester	1.81 (0.41, 3.22)	0.00 (-1.18, 1.18)	0.56 (-1.05, 2.17)	-0.75 (-2.11 <b>,3</b> ).6	-0.48 (-2.28, 1.33)	-2.11 (-3.63, -0.60)
20 10 11 02	Third trimester	0.04 (-1.54, 1.62)	-1.97 (-3.29, -0.65)	0.51 (-1.45, 2.47)	-1.92 (-3.57, <b>≥</b> 0.2€)	1.52 (-0.66, 3.69)	-1.92 (-3.76, -0.09)
22	Total pregnancy	0.67 (-0.56, 1.89)	-1.08 (-2.11, -0.05)	0.41 (-0.83, 1.64)	-1.15 (-2.19, a).1 <mark>2</mark> ).1	0.67 (-0.77, 2.12)	-1.68 (-2.89, -0.46)

<sup>23</sup>All significant findings in the table are bold.

29 djusted for the other air pollutant. ar technologies une 11, 2025 at Agence Bibliographique de l 

<sup>2</sup>¾ Il significant findings in the table are bold.

2⁴ Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

<sup>25 \*</sup>Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births, and

- In the adjusted model, the risk of PDD was found to increase by 112% and 42% with each per-IQR increase in PM<sub>2.5</sub> (OR: 2.12, 95% CI: 1.45, 3.11) and NO<sub>2</sub> (OR: 1.42, 95% CI: 1.16, 1.75) in the 90 days prior to conception (**Table 4**). There was also a significant association between increased NO<sub>2</sub> exposure and the risk of PDD in T1 (OR: 1.29, 95% CI: 1.05, 1.58), T3 (OR: 1.27, 95% CI: 1.01, 1.60), and the whole pregnancy period (OR: 1.17, 95% CI: 1.02, 1.36). We
- did not observe any association with MDD in any pregnancy periods.

Page 19 of 48  1 2 3 74	Table 4 Associations betw	ween PM <sub>2.5</sub> and NO <sub>2</sub> (		MJ Open at pregnancy periods a	/bmjopen-2023-085875 by copyright, incaddin and mental	chomotor developme	ental delay
6		Crude	models	Adjusted r	nodels*	Co-exposure	e models**
7 8	Per IQR increase in	MDD (95% CI) (N=946)	PDD (95% CI) (N=946)	MDD (95% CI) (N=945)	PDD (95% CHE) (N=94%) 8 2	MDD (95% CI) (N=945)	PDD (95% CI) (N=945)
1 Estimated	90 days prior to conception	1.45 (1.16, 1.83)	1.49 (1.20, 1.83)	0.95 (0.64, 1.42)	2.12 (1.45, 2 1)	0.97 (0.63, 1.51)	1.78 (1.17, 2.71)
<sup>1</sup> lexposure	First trimester	1.05 (0.83, 1.33)	1.04 (0.84, 1.28)	1.14 (0.73, 1.79)	1.42 (0.96, 🛂 🖔	1.35 (0.80, 2.25)	1.16 (0.74, 1.82)
12 13to PM <sub>2.5</sub>	Second trimester	0.63 (0.49, 0.80)	0.77 (0.63, 0.95)	0.81 (0.54, 1.22)	0.72 (0.51, 2.3)	0.83 (0.52, 1.31)	0.57 (0.38, 0.85)
14	Third trimester	1.23 (0.96, 1.58)	1.19 (0.95, 1.49)	1.25 (0.82, 1.90)	1.17 (0.80, 🖁 👼 👸	1.39 (0.87, 2.23)	0.98 (0.64, 1.49)
15	Total pregnancy	0.84 (0.69, 1.03)	0.98 (0.82, 1.18)	1.07 (0.82, 1.39)	1.07 (0.84, <b>ā. శ్రీ</b>	1.17 (0.86, 1.59)	0.90 (0.68, 1.20)
16Estimated	90 days prior to conception	1.20 (0.99, 1.45)	1.41 (1.19, 1.67)	0.97 (0.77, 1.21)	1.42 (1.16, 🖁 🞝 🕏	0.97 (0.76, 1.25)	1.24 (0.99, 1.56)
17 18exposure	First trimester	0.97 (0.80, 1.17)	1.18 (0.99, 1.40)	0.91 (0.72, 1.13)	1.29 (1.05, 3, 58)	0.84 (0.65, 1.09)	1.24 (0.99, 1.57)
$\frac{19}{20}$ to NO <sub>2</sub>	Second trimester	0.79 (0.66, 0.95)	1.04 (0.88, 1.22)	0.94 (0.76, 1.15)	1.14 (0.95,	0.98 (0.77, 1.24)	1.31 (1.06, 1.63)
20	Third trimester	1.04 (0.85, 1.28)	1.25 (1.04, 1.50)	0.94 (0.73, 1.21)	1.27 (1.01, 3.60)	0.86 (0.65, 1.14)	1.28 (0.99, 1.65)
21	Total pregnancy	0.92 (0.79, 1.07)	1.16 (1.01, 1.33)	0.94 (0.80, 1.11)	1.17 (1.02, <b>3</b> .3 <b>6</b> )	0.9 (0.75, 1.08)	1.21 (1.02, 1.43)
All giomific	cont findings in the table are be	1.1			<del>5</del> 🚉		

24ll significant findings in the table are bold.

24 Il significant findings in the table are bold.
24 Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

25\*Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly hause old income level, and season of births, and

<sup>26</sup>adjusted for the other air pollutant.

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### **Discussion**

We analyzed associations between modelled PM<sub>2.5</sub> and NO<sub>2</sub> pre- and during pregnancy with birth and neurodevelopment outcomes in singleton children born in a south-western metropolis of China in 2015-16. We found the likelihood of SGA increased by 33% per IQR higher exposure to NO<sub>2</sub> in the whole pregnancy periods after adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births and PM<sub>2.5</sub>. For childhood cognitive development, increased exposure to PM<sub>2.5</sub> and NO<sub>2</sub> in the 90 days prior to conception were both associated with lower PDI scores, with the effect size per IQR being higher for PM<sub>2.5</sub> than for NO<sub>2</sub>. Increased NO<sub>2</sub> exposure was associated with an increased risk of PDD during different trimesters of pregnancy.

Many studies from other geographic areas, including Europe (33-35), the United States (22, 26), and Asia (23, 36-38) have found that prenatal air pollution exposure has a negative impact on a variety of neurodevelopmental outcomes. Our finding of a negative association between prenatal NO<sub>2</sub> air pollution exposure and infant neurocognitive development is consistent with these reports. A recent Chinese birth cohort study of 15,778 child-mother pairs in Foshan reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with an increased risk of suspected developmental delay (OR: 1.06, 95% CI: 0.94, 1.19) measured by a five-domain scale and developmental quotient (DQ) (23). A birth cohort study of 520 mother-child pairs in South Korea reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with impairment of psychomotor development ( $\beta = -1.30$ , p = 0.05) but – as in the present study not with cognitive function ( $\beta = -0.84$ , p = 0.20) (36). However, results from previous research varied by air pollutants. For example, a Chinese study of 1193 mother-newborn pairs in Changsha found significant associations between PM<sub>2.5</sub> exposure in trimester two and lower neurobehavioral developmental scores, while other air pollutants such as PM<sub>10</sub>, carbon monoxide (CO), and Sulfur dioxide (SO<sub>2</sub>) had null or even reverse associations. In this study, we observed that the negative effect of NO<sub>2</sub> exposure during pregnancy on PDI is significant at 5% level; this negative effect of NO2 still remained after adjustment for PM2.5. This heterogeneity may relate to the temporality of exposure assessment, types of outcome assessment instruments or evaluators, and the levels of air pollution. In addition, air pollution mixtures may have differed among the study regions, thus there are several potential explanations for the heterogeneity of the findings. We also observed negative correlations between certain exposures, indicating the need to consider potential collinearity in our twopollutant models. In Chongqing, a major industrial city in southwest China, air pollution may

come from industrial and traffic emissions, construction activities, and dust, and negative correlations may occur if different sources contribute disproportionately to each pollutant. 

Their correlations may also be affected by seasonal changes and variations in weather patterns.

Future research should also explore the impact of source-specific air pollution on children's

cognitive health.

> To date, most studies on prenatal air pollution exposure and child neurodevelopment have been conducted in developed countries with relatively low levels of air pollution. In this study, the level of air pollution was higher (median PM<sub>2.5</sub>: 57.31 μg/m<sup>3</sup>, IQR: 5.76; median NO<sub>2</sub>: 50.46 µg/m<sup>3</sup>, IQR: 5.51) compared to studies in developed countries such as Europe and the United States. In a multi-centre European cohort, the mean PM<sub>2.5</sub> and NO<sub>2</sub> exposure concentration during pregnancy were 13.4 µg/m<sup>3</sup> and 11.5 µg/m<sup>3</sup> (33). Researchers found that the psychomotor development score significantly decreased by 0.68 points (95% CI: -1.25, -0.11) for every 10 µg/m<sup>3</sup> increase in NO<sub>2</sub>, and there was also a non-significant decrease of 1.64 points (95% CI: -3.47, 0.18) for every 5 µg/m<sup>3</sup> increase in PM<sub>2.5</sub> during pregnancy (33). Factors such as the types of pollutants and concentrations may differ between China and other regions with a lower air pollution level, leading to variations in the observed effects.

> Contrary to expectations, we found significant positive associations between prenatal exposure to PM<sub>2.5</sub> air pollution in the second trimester and PDI. However, no association was observed between PM<sub>2.5</sub> exposures in the second trimester and the risk of PDD. Given the existing literature and the conflicted observation here, we believe that this is likely to be spurious/sample specific. Some plausible explanations include the uneven distribution of PDI scores, the potentially inappropriate selection of the cut-off value of 85 (which may not effectively discriminate between groups), or the possibility that the observed outcome occurred by chance. Several epidemiological studies have reported associations between prenatal exposure to high levels of PM<sub>2.5</sub> and lower neurodevelopment in children ranging in age from 6 months to 6 years (12, 34, 39-41). In agreement with our findings, a multi-centre cohort study from six European countries investigated the effects of prenatal exposure to multiple air pollutants including PM<sub>2.5</sub>, PM<sub>10</sub>, coarse particles, NO<sub>2</sub> and nitrogen oxides (NOx) among 9482 children between 1 and 6 years; the authors found nonsignificant positive associations between prenatal  $PM_{2.5}$  exposure and normal neurodevelopment ( $\beta$ : 1.64, 95% CI: -3.47, 0.18; per 5 μg/m<sup>3</sup> increase in PM<sub>2.5</sub>) (33). Similarly, another study examining the effects of multiple pollutant exposures on early childhood cognition at 40 days of age in a highly exposed area of Spain also found PM<sub>10</sub>, PM<sub>coarse</sub>, PM<sub>2.5absorbance</sub>, NO<sub>2</sub>, NO<sub>x</sub>, and Ozone (O<sub>3</sub>) were linked to lower

motor function in children, except for PM<sub>2.5</sub> (42). The inconsistent findings could be because of heterogeneity between studies in terms of exposure (e.g., exposure assessment methods used,

 $PM_{2.5}$  exposure levels, or composition of  $PM_{2.5}$ ).

 Aside from the conflicting findings regarding prenatal  $PM_{2.5}$  exposure and neurodevelopmental outcomes, results regarding the most potential sensitive time windows before and during pregnancy are also inconclusive. Some studies suggested that early-to-mid pregnancy might be a potential sensitive period (22, 23), while other studies found stronger associations for middle-

to- late pregnancy, thus results are equivocal (20, 25, 26).

The potential biological mechanisms by which air pollution could affect neurodevelopment are not yet clearly understood. There is evidence suggesting that exposure to prenatal PM<sub>2.5</sub> could potentially induce maternal immune activation during pregnancy (43). Higher levels of cytokines or reactive oxygen species may potentially interfere fetal neurodevelopment through three mechanisms: crossing the placental barrier into the fetal body, inducing fetal immune dysregulation, and contributing to inadequate placental perfusion that affects nutritional processes and oxygenation of maternal blood(44). More research is needed to investigate trimester effects of air pollution on neurodevelopment and provide better understanding on the underlying biological mechanisms. Our study is the first to consider an exposure window 90 days prior to conception for NO<sub>2</sub>. A novel observation is that effects of NO<sub>2</sub> or PM<sub>2.5</sub> air pollution on child cognition can be seen at least 90 days prior to conception, representing a potentially vulnerable periods in relation to air pollution on neurodevelopment. Similar results were found in previous study recruited 1329 mother-child pairs in Wuhan, China (12). This study reported a higher level of PM<sub>2.5</sub> during preconception (Median: 76.1 µg/m<sup>3</sup>) and in the first trimester (Median: 82.3 μg/m<sup>3</sup>). This study found for each doubling of PM<sub>2.5</sub> exposure during preconception, children's PDI scores was reduced by 6.15 (95% CI: -8.84, -3.46) points. A potential explanation is that preconception air pollution exposures induce genetic and epigenetic alterations in sperm, that increase the risk of adverse health outcomes in offspring (45, 46). To date, all studies examined the effect of maternal preconception exposure while omitting paternal exposures (17). Future studies should consider the effect of preconception paternal exposure in relation to childhood health outcomes.

This study has several strengths. We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification if using monitoring stations. This is an novel study to investigate both pre-conception and prenatal

  $PM_{2.5}$  and  $NO_2$  exposure with neurodevelopment outcomes among young infants, in the context of a relatively high air pollution urban environment. The exposure levels in our study were similar as those in comparable urban areas in Chinese cities. A study in Shanghai, China reported an average  $NO_2$  exposure during pregnancy from 2014 to 2015, predicted by the LUR model, of 48.23  $\mu$ g/m³ (Mean  $PM_{2.5}$  in our study: 50.52  $\mu$ g/m³) (47). Similarly, a study in Tianjin found the annual average  $PM_{2.5}$  exposure to be 62  $\mu$ g/m³ in 2017 (Mean  $NO_2$  in our study: 57.48  $\mu$ g/m³) (48). Wu et al. developed a LUR model for  $PM_{2.5}$  in the main urban area of Chongqing (49). This model predicted an annual average  $PM_{2.5}$  concentration of 40.6  $\mu$ g/m³ (49), whereas our prediction is higher at 55.9  $\mu$ g/m³ (19). This difference can be attributed to the temporal variations. Wu et al. used monitoring data from 2013, while we utilized data from 2015. It could be considered that our GAM model, with its temporal component, could explain temporal variations and is more suitable for pregnancy-specific exposure estimates.

A major limitation of this study was that our sample size was relatively small, limiting the statistical power to assess several outcomes, although the higher exposures in Chongqing than in some other studies may increase probability of detecting effects. In terms of limitations, due to a lack of information on participant time-activity patterns, exposure estimates in this study refer only to ambient concentrations at home addresses, and no other activity spaces (e.g., indoor, workplace, commuting) were considered. We may have thus underestimated total air pollution exposure. Second, we defined exposure windows for clinically-defined trimesters; sensitive periods may be shorter or longer than 3 months, or they may exist in the overlap of multiple trimesters. However, we were unable to investigate the sensitive time windows using established methods such as distributed lag non-linear models due to the lack of highly timeresolved air pollution estimates. Third, the performance of the NO<sub>2</sub> spatiotemporal model was moderate (COR-R<sup>2</sup>: 0.39), which may introduce exposure misclassification and therefore bias in the coefficients. Finally, we were unable to include some other air pollutants such as polycyclic aromatic hydrocarbons (PAH), black carbon (BC) and Ozone, which have bene found particular harmful to neurodevelopment in children (50). Although we have accounted for most of the important confounders in this study, unfortunately, we did not collect information on the feeding patterns of infants. This may undermine the validity and reliability of our findings.

#### Conclusion

This study provides evidence for an association between NO<sub>2</sub> exposure prior to- and during pregnancy with birth and neurodevelopmental outcomes in a birth cohort in Chongqing, China. Exposure to NO<sub>2</sub> and PM<sub>2.5</sub> exposure before pregnancy was associated with a lower psychomotor development score. Increased NO<sub>2</sub> exposure was linked to a risk of psychomotor development delay during various pregnancy trimesters.



## 212 List of abbreviations

dy mass index  Vley Scales of Infant Development  Inese version of Bayley Scales of Infant Development  Infidence interval  Implex Lipids in Mothers and Babies
vley Scales of Infant Development nese version of Bayley Scales of Infant Development nfidence interval
nese version of Bayley Scales of Infant Development  infidence interval
nfidence interval
mplex Lipids in Mothers and Babies
bon monoxide
rrelation
velopmental origins of health and disease
velopmental quotient
ographic information systems
erquartile range
w birth weight
ge for gestational age
nd-use Regression
ntal Developmental Delay
ntal Development Index
rogen oxides
rogen dioxide
d ratio

PDD	Psychomotor Developmental Delay
PDI	Psychomotor Development Index
PM	Particulate matter
PM <sub>2.5</sub>	Particulate matter with diameter ≤2.5μm
PTB	Preterm birth
SGA	Small for gestational age
$SO_2$	Sulfur dioxide
TLBW	Term low birth weight
T1	First trimester
T2	Second trimester
T3	Third trimester
WP	Whole pregnancy period
90D	90 days prior to pregnancy

#### **Declarations**

## Ethics approval and consent to participate

- 216 Ethical approval for this study was granted by the Ethics Committee of Chongqing Medical
- 217 University (#2014034). The participants provided their written informed consent to
- 218 participate in this study. Written informed consent was obtained from the individual(s) for the
- publication of any potentially identifiable images or data included in this article.

## Data availability statement

- The data that support the findings of this study are available from Chongqing Medical
- 222 University but restrictions apply to the availability of these data, which were used under
- license for the current study, and so are not publicly available. Data are however available
- from the authors upon reasonable request and with permission of Chongqing Medical
- 225 University.

# **Competing interests**

The authors declare that they have no conflicts of interests.

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## **Contributors**

- Y. X., T.L.H., H.Z. and P.B. conceived and designed research; T.Z., Y.X. and H.Z. recruited
- the patients and collected the samples; T.K., A.H., and J.G. constructed the air pollution
- 243 model; Y.C. analyzed, interpreted the data and prepared the figures; Y.C and T.K were major

contributors in writing the manuscript text; YSC, JC, TLH, YX, ALH, and PNB substantively revised the manuscript; All authors read and approved the final manuscript.

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- 408 Figures
- 409 Figure 1 Flow chart of the study population in CLIMB
- 410 Figure 2 Study area and location of monitoring sites (OpenStreetMap contributors,

411 2015; https://data.nextgis.com/en/region/CN-50/).



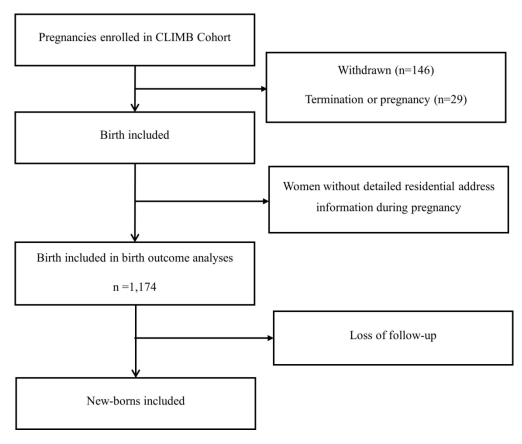


Figure 1 Flow chart of the study population in CLIMB  $173 \times 144 \text{mm} (300 \times 300 \text{ DPI})$ 

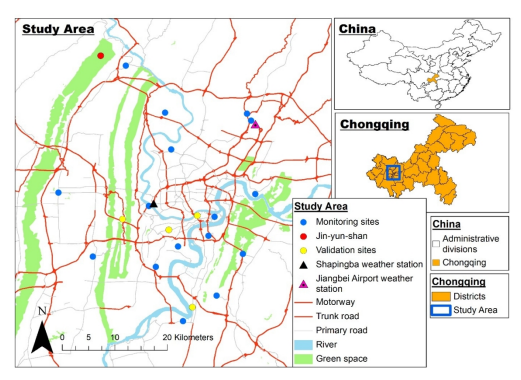


Figure 2 Study area and location of monitoring sites (OpenStreetMap contributors, 2015; https://data.nextgis.com/en/region/CN-50/).

159x112mm (220 x 220 DPI)

Associations of air pollution exposures in preconception and pregnancy with birth outcomes and infant neurocognitive development: analysis of the Complex Lipids in Mothers and Babies (CLIMB) prospective cohort in Chongqing, China

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## **Supplement**

# eTable 1 Mental Development Index (Chinese version)

## 智力量表

(\*可偶尔观察到)

序号	月龄	条目	计分
1	0.1	对铃声反应	
2	0.1	抱起时安静	
3	0.1	对摇鼓声反应	
4	0.1	对尖声反应: (电灯开关)	
5	0.1	短暂地注视红环	
6	0.2	短暂地注视人	
7	0.4	稍长时间地注视红环	
8	0.5	眼的水平协调活动(红环)	
9	0.7	眼的水平向天活动 (光)	
10	0.7	眼睛追随移动的人	
11	0.7	对说话声反应	
12	0.8	眼的垂直协调活动 (光)	
13	0.9	发声一至两次	
14	1	眼的垂直协调活动(红环)	
15	1.2	眼的旋转协调活动 (光)	
16	1.2	眼的旋转细条活动 (光环)	
17	1.3	*自由环视周围	
18	1.5	社交笑: 测试者谈话与微笑时	
19	1.6	眼转向红环	
20	1.6	眼转向光	
21	1.6	*发声至少四次	
22	1.7	期待性兴奋	
23	1.7	对面部的纸有反应	
24	1.9	能用视觉辨认母亲	
25	1.9	社交笑: 测试者微笑与安静时	
26	2	*对测试者的微笑和说话有发声反应	
27	2.1	*用眼睛寻找声源(详细说明)	
28	2.2	*发出两种不同的声音	
29	2.2	对手的遮蔽眨眼	
30	2.2	对面孔的消失有反应	
31	2.4	注视方木	
32	2.6	从一物转看另一物	
33	2.6	眼睛追随铅笔	
34	2.7	对抱起有预感性的调节反应	
35	2.9	目光追随横过桌面的球	
36	2.9	头追随悬摆的环	

37	3.1	
38	3.2	操作红环
39	3.3	简单地玩摇鼓
40	3.4	*轻轻地抚摸桌沿
41	3.4	*意识到陌生环境
42	3.5	头转向铃声
43	3.6	头转向摇鼓声
44	3.6	*手碰手的玩耍
45	3.6	将红环送入口中
46	3.7	伸手够悬环
47	3.8	看自己的手
48	4.2	接近悬环 (优势手)
49	4.4	*发声时的姿态(描述)
50	4.4	*主动抚摸桌沿
51	4.4	接近镜像
52	4.4	注意小糖丸
53	4.6	伸手取方木
54	4.7	喜欢嬉戏
55	4.9	伸手时眼手协调
56	4.9	拾起方木 (优势手)
57	5	保持两块方木
58	5	持久地看红环
59	5	头部跟着掉下的勺转动
60	5	探索性地玩纸
61	5	对镜像微笑
62	5	坚持够东西
63	5.1	在小床内重新找到摇鼓
64	5.1	*辨别生人
65	5.4	举起倒扣的茶杯
66	5.5	*敲打玩耍
67	5.5	探索性地玩细绳
68	5.5	伸手取第二块方木
69	5.6	*由一手向另一手传递物体
70	5.8	*对产生响声感兴趣
71	5.9	灵巧而直接地拾起方木
72	6	*对镜像开玩笑
73	6	用把柄举起茶杯
74	6	寻找掉落的勺子
75	6.1	牵拉细绳获取红环
76	6.1	保留三块方木中的两块
77	6.6	*发出四个不同的音节
78	6.8	能配合玩游戏
79	7	恰当地牵拉细绳获取红环
80	7.1	玩摇铃, 对细节感兴趣
0		

82	7.4	
	7.4	
83	7.5	*选择性地倾听熟悉的词语
84	8	*对 da-da 或类同词
85	8.1	暴露玩具
86	8.2	注意测试者的乱写
87	8.3	将手指插入桩板洞中
88	8.6	观看书中图画
89	8.9	对他人的言语要求有反应
90	9.1	拿起茶杯获得方木
91	9.8	寻找盒子里面的东西
92	10.3	遵照命令将方木放入茶杯(放入数)
93	10.7	企图模仿乱写
94	10.8	模仿用勺子搅拌
95	10.9	遵照命令停止
96	10.9	推动小汽车
97	11	模仿地拍打哨娃
98	11.1	*重复引入发笑的把戏
99	11.2	解开裹着的方木
100	11.2	将三块方木放入杯中
101	11.4	*快速而不清的表达
102	11.4	揭开兰盒子的盖
103	11.5	翻开书页
104	11.5	摇晃悬环的
105	11.8	将骰子放入盒中(6个)
106	12	恰当地握持画笔
107	12.2	模仿说单词(记录用过的词)
108	12.4	重复地插一根桩钉
109	12.5	用手势表达想要的东西
110	12.9	自动乱写
111	12.9	能说两个词
112	13	搭两层塔
113	13.1	出示鞋子或其他衣服或自己的玩具
114	13.2	从瓶中移出小糖丸
115	13.3	掺九块方木放入杯中
116	14.3	*盖上圆盒
117	14.4	兰色模板: 放置一个圆形模块
118	14.8	用棍子够取玩具
119	15.4	搭三层塔
120	15.7	在 70 秒钟内插完桩钉
121	16.1	指出娃娃身体的各部分:三个部位以上
122	16.3	粉红模板:放置圆形模块
123	16.6	兰色模板: 放置两个圆形模块
124	17.2	用笔模仿画一划
125	17.5	在 42 秒钟内插完桩钉
126	17.6	说出一物名
120	1	20 K2 H

127	17.7	   对娃娃执行指令(在通过部位打钩:椅、杯、鼻)
128	18.1	用语言表达要求
129	18.6	不用于一划的乱写?
130	18.8	兰色模板: 放置两个圆快和方块
131	18.8	指出三幅画
132	19.1	能说两个单词的句子
133	19.2	说出一副画名
134	19.2	说出两幅画名
135	19.3	找出两物
136	19.8	在 30 秒钟内插完桩钉
137	20.4	粉红模板:完成
138	20.4	搭六层塔
139	20.5	兰色模板,放置六个模块
140	21	指出五副画
141	21.1	说出三物名
142	21.2	勉强合格地安装破娃娃
143	21.2	区别两物:杯、盘、盒
144	22.8	辨认钟表: 第四张图 1, 2, 3, 4, 5
145	22.9	说出三幅画名
146	23.8	粉红模板(反转)
147	24.3	近似地安装破娃娃
148	24.6	区别三物: 杯、盘、盒
149	24.7	兰色模板,在 150 秒钟内完成
150	25	搭八层塔
151	25.1	指出七副画
152	25.1	用方木搭火车
153	25.7	说出五副画名
154	26.3	模仿笔划:垂直线和水平线
155	27.1	辨认钟表: 第2张图
156	27.6	理解两个方位词
157	28	在 22 秒钟内插完桩钉
158	28.5	兰色模板: 90 秒钟内完成
159	29.5	折纸
160	29.6	兰色模板: 60 秒钟内完成
161	30+	正确安装破娃娃
162	30+	"—"的概念
163	30+	理解三个方位词

### 运动量表

(\*可偶尔观察到,△可在施测智力量表时观察到)

序号	月龄	条目	计分
1	0.1	抱起靠肩时抬头	
2	0.1	抱起靠肩时调整姿势	
3	0.1	侧头	
4	0.1	爬起	
5	0.8	△保留红环	
6	0.8	*伸臂玩耍	
7	0.8	*踢腿玩耍	
8	0.8	头起竖起:垂直位	
9	1.6	头部稳定地竖起	
10	1.7	抬头 (背悬位)	
11	1.8	由侧卧转向仰卧	
12	2.2	在俯卧位时用双臂撑起自己	
13	2.2	支撑下坐起	
14	2.5	保持头部稳定	
15	2.6	*双手张开占优势	
16	3.3	头平衡	
17	3.4	*尺侧一手掌抓握方木	
18	3.5	轻度支撑坐位	
19	4.3	*由仰卧转向侧卧	
20	4.7	努力想坐起	
21	5.0	部分的拇指相对(桡侧一手掌)拾起方木	
22	5.1	独坐片刻	
23	5.1	*单手抽取	
24	5.2	*转腕	
25	5.2	牵拉坐起	
26	5.6	△试图获取小糖丸	
27	5.7	独立 30 秒钟或以上	
28	5.8	由仰卧转向俯卧	
29	6.2	稳定地独坐	
30	6.5	独坐时协调好	
31	6.6	*舀起小糖丸	
32	6.6	△完全的拇指相对拾起方木	
33	7	早期跨步运动	
34	7.5	牵拉站起	
35	7.6	*不完全的拇指相对抓糖丸	
36	7.6	走路之前的行进方式 (俯卧、手膝、手足、其他)	
37	8.3	使两个勺子或方木在中线相碰	
38	8.5	跨步运动	

39	8.6	自己坐起
40	8.6	借助家具站起
41	8.9	精细地抓糖丸 (灵巧地钳夹)
42	9.6	拍手(中线技巧)
43	9.8	坐下
44	10	扶助下行走
45	11.1	独站
46	12	投球
47	12.1	独走
48	12.4	起立丨
49	13.2	扶助下右足独站
50	13.7	扶助下左足独站
51	14.1	侧身走
52	14.5	扶助上楼梯
53	14.7	倒退走
54	15.1	扶助下楼梯
55	17.6	试图站在行木上
56	18.7	左足独站
57	19.3	单足踏在行木上走
58	19.9	起立Ⅱ
59	20.1	右足独站
60	21.1	走直线: 大致方向
61	23.1	行木: 双足站立
62	24	踮脚走几步
63	24.3	独自上楼梯: 双足
64	24.4	双足跳离地面
65	25.3	独自下楼梯
66	25.6	行木: 企图跨步
67	25.6	倒行两米半
68	25.7	自第一级台阶下跳下
69	29.2	自第二级台阶下跳下
70	29.8	踮脚走两米半
71	29.9	跳远: 10 至 35cm (记录距离)
72	30+	起立:
73	30+	上楼梯:双足交替向前
74	30+	行木: 交替步伐走部分路程
75	30+	保持双足走在直线上 (两米半)
76	30+	跳远: 35cm 至 60cm
77	30+	跳过: 5cm 高的绳子
78	30+	跳远: 60cm 至 85cm
79	30+	独脚跳两次以上
80	30+	下楼梯: 双足交替向前
81	30+	跳过 20cm 高的绳子

eTable 3 Distributions of PM2.5 and NO2 exposure level in 90 days prior to conception, each trime stern T1, T2, and T3) and combined across whole pregnancy period (WP) (n = 1,174)

8 9		Estimated exposure (µg no N								
10 11	·	N	Minimum	25 <sup>th</sup> percentile	Mean ± SD	Mediang + JQR	75 <sup>th</sup> percentile	Maximum		
12 13	Estimated exposure to PM <sub>2.5</sub>					wnlo				
14 15 16 17 18 19 20 21 22 23 24 25	90 days prior to conception	1,174	38.17	44.00	$52.91 \pm 10.99$	48.4 3 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	62.06	80.53		
	First trimester	1,174	37.26	43.77	$52.07 \pm 10.98$	47.2 <b>@ (5)</b> .31	61.08	82.41		
	Second trimester	1,174	38.46	47.57	$58.64 \pm 12.21$	57.9 B B B .62	67.19	90.02		
	Third trimester	1,174	37.03	47.25	$61.83 \pm 16.04$	58.8 <b>2</b> ± <b>28</b> .7	75.95	96.48		
	Total pregnancy	1,174	46.69	54.85	$57.48 \pm 3.97$	57.3 ± = .76	60.61	66.98		
	Estimated exposure to NO <sub>2</sub>				1/0	n.bm Jining				
	90 days prior to conception	1,174	25.86	45.49	$49.59 \pm 6.34$	49.9 <b>4</b> ± <b>8</b> .27	53.76	70.48		
26	First trimester	1,174	20.81	44.60	$48.8 \pm 6.43$	48.9 <b>2</b> .± <b>8</b> .51	53.10	69.31		
27 28 29 30 31 32	Second trimester	1,174	28.93	47.18	$50.98 \pm 6.23$	51.2 ± \$ .72	54.90	70.42		
	Third trimester	1,174	20.57	47.20	$51.79 \pm 6.78$	52.4 <b>9</b> ± <b>9</b> .47	56.67	75.12		
	Total pregnancy	1,174	27.50	47.89	$50.52 \pm 5.08$	50.4 <b>8</b> ± <b>8</b> 51	53.40	67.53		

eTable 4 Pearson's correlations of PM<sub>2.5</sub> and NO<sub>2</sub> between each of the five different pregnancy time periods (N = 1,174)

Estimated exposure to		$\mathrm{PM}_{2.5}$				NO <sub>2</sub>					
		90 days prior	First	Second	Third	Total	90 days prigr	First	Second	Third	Total
		to conception	trimester	trimester	trimester	pregnancy	to conception 202	trimester	trimester	trimester	pregnancy
PM <sub>2.5</sub>	90 days prior to conception	1					4. Donemo				
	First trimester	-0.065	1				ont S to te				
	Second trimester	-0.779	-0.2012	1			padec uperi xt and				
	Third trimester	0.288	-0.7613	-0.1688	1		l from eur ( <i>i</i> d data				
	Total pregnancy	-0.534	-0.2709	0.6838	0.3858	1	ABES ABES				
NO <sub>2</sub>	90 days prior to conception	0.6383	0.0684	-0.4588	0.3714	0.0376	1 ing,				
	First trimester	0.1537	0.6352	-0.0159	-0.4927	-0.0633	0.5545	1			
	Second trimester	-0.431	0.0714	0.6269	-0.0133	0.7251	0.3345	0.5399	1		
	Third trimester	0.3027	-0.5213	0.0528	0.6817	0.4432	0.71498	0.2159	0.5145	1	
	Total pregnancy	0.0057	0.0781	0.2862	0.0737	0.4779	0.678 emil	0.7435	0.8755	0.7331	1

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eTable 5 Associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposure in different pregnancy periods and adverse birgh outcomes (unadjusted models)

5					g or			
6 7	Mean difference			Odd ratios				
8 Per IQR increase in		Birth weight, grams (95%	Birth length, cm	PTB (case: 33)	LB <b>W</b> 55 (50)	LGA (case: 108)	SGA (case: 84)	
9	rei iQK increase in	CI)	(95% CI)	(95% CI)	( <b>9</b> 50/2)	(95% CI)	(95% CI)	
10		(N=941)	(N=927)	(N=945)	<b>4</b> 5)	(N=945)	(N=945)	
Estimated	90 days prior to conception	9.28 (-31.26, 49.83)	-0.09 (-0.27, 0.09)	0.98 (0.56, 1.74)	1.35 <b>4</b> ( <b>9</b> ) <b>2</b> , 2.36)	1.2 (0.87, 1.64)	0.98 (0.68, 1.41)	
13exposure	First trimester	21.95 (-16.90, 60.80)	0.14 (-0.03, 0.31)	0.98 (0.57, 1.70)	1 ( <b>ਉ.<u></u> 275</b> 1.77)	0.97 (0.71, 1.33)	0.78 (0.54, 1.13)	
14 15 <sup>to</sup> PM <sub>2.5</sub>	Second trimester	-18.21 (-57.78, 21.37)	0.04 (-0.13, 0.21)	0.85 (0.48, 1.50)	$0.61$ $\frac{1}{8}$ $\frac{1}{8}$ $\frac{1}{8}$ , $1.15$	0.92 (0.67, 1.27)	1.33 (0.94, 1.89)	
	Third trimester	-37.38 (-81.43, 6.68)	-0.32 (-0.51, -0.13)	1.35 (0.74, 2.47)	1.51 (2) (2) (2.85)	1.08 (0.76, 1.54)	1.12 (0.76, 1.66)	
16 17	Total pregnancy	-20.02 (-55.69, 15.65)	-0.1 (-0.26, 0.05)	0.81 (0.49, 1.33)	0.69 <b>a (<u>5</u>∕∄</b> , 1.16)	1 (0.75, 1.34)	1.2 (0.87, 1.66)	
1 <b>E</b> stimated	90 days prior to conception	-13.23 (-45.50, 19.03)	-0.12 (-0.26, 0.02)	1.2 (0.76, 1.89)	1.62 (75)	1.21 (0.93, 1.57)	1.24 (0.92, 1.66)	
19 exposure	First trimester	0.3 (-32.36, 32.96)	0.08 (-0.06, 0.22)	1.01 (0.64, 1.60)	1.15 (0.7), 1.86)	1.17 (0.90, 1.52)	1.27 (0.94, 1.71)	
20 to NO <sub>2</sub>	Second trimester	-22.85 (-53.39, 7.70)	-0.04 (-0.17, 0.09)	1.11 (0.72, 1.70)	1.08 <b>2</b> 0. <b>6</b> , 1.69)	1.06 (0.83, 1.36)	1.46 (1.10, 1.93)	
21 to NO <sub>2</sub> 22	Third trimester	-32.72 (-67.16, 1.72)	-0.16 (-0.32, -0.01)	1.13 (0.69, 1.85)	1.35 (0.80), 2.28)	1.24 (0.94, 1.65)	1.58 (1.14, 2.18)	
23	Total pregnancy	-16.58 (-43.35, 10.20)	-0.03 (-0.15, 0.09)	1.03 (0.71, 1.50)	1.13 (0.75, 1.69)	1.16 (0.93, 1.44)	1.44 (1.13, 1.85)	
All signific	cant findings in the table are be	old.			an Co			
26					n∕on d sim			
27				mii J				
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		pyrigh	ben-20	
eTable 6 Associations between PM <sub>2.5</sub> and NO <sub>2</sub> exposure in diffe	erent pregnancy periods and advers	it, ince	irgh outcom	nes (co-exposure models)
	<del> </del>	ding	75 <b>9</b> 0.11	_

6	Mean difference			o Odd ratios				
7	Per IQR increase in	Birth weight, grams	Birth length, cm	PTB (case: 33)	LBW (case: 30)	LGA (case: 108)	SGA (case: 84)	
8	i ei iQK merease m	(95% CI)	(95% CI)	(95% CI)	(3 <b>5 2</b> I)	(95% CI)	(95% CI)	
9 10		(N=941)	(N=927)	(N=945)	(1 <mark>2 (1</mark> 2)	(N=945)	(N=945)	
1 Estimated	90 days prior to conception	75.00 (-9.86, 159.86)	0.23 (-0.14, 0.60)	0.98 (0.56, 1.89)	0.41 (2)	1.14 (0.55, 2.40)	1.18 (0.48, 2.92)	
12 exposure	First trimester	19.59 (-71.23, 110.41)	0.00 (-0.40, 0.39)	0.97 (0.26, 3.65)	0.66 ( <b>3</b> . <b>7</b> 4 <u>5</u> 3.05)	0.67 (0.32, 1.42)	0.73 (0.28, 1.93)	
13 <sup>1</sup> 14to PM <sub>2.5</sub>	Second trimester	-25.62 (-104.32, 53.09)	0.08 (-0.26, 0.42)	1.34 (0.37, 4.86)	0.94 (3) <b>5</b> (4.21)	0.83 (0.42, 1.62)	0.69 (0.34, 1.40)	
15	Third trimester	13.77 (-72.33, 99.86)	-0.2 (-0.58, 0.17)	1.12 (0.31, 4.07)	0.94 (2.3.35)	1.00 (0.48, 2.12)	0.57 (0.26, 1.23)	
16	Total pregnancy	21.13 (-36.41, 78.67)	0.02 (-0.23, 0.27)	0.73 (0.33, 1.61)	0.52 <b>(a) (5</b> /2 <b>4g</b> 1.15)	0.98 (0.60, 1.61)	0.55 (0.32, 0.96)	
1Estimated	90 days prior to conception	-18.63 (-64.02, 26.76)	-0.09 (-0.29, 0.10)	1.24 (0.61, 2.49)	1.3 (2.64)	1.27 (0.84, 1.90)	1.39 (0.90, 2.15)	
19exposure	First trimester	-14.53 (-61.15, 32.09)	0.05 (-0.16, 0.25)	0.91 (0.45, 1.83)	1.14 🕳 🚾 2.37)	1.33 (0.89, 2.00)	1.70 (1.07, 2.69)	
20	Second trimester	-14.46 (-57.45, 28.54)	-0.08 (-0.26, 0.11)	1.22 (0.63, 2.36)	1.36 (0.68 2.71)	1.27 (0.87, 1.87)	1.50 (1.00, 2.24)	
20 21 to NO <sub>2</sub>	Third trimester	-13.13 (-64.87, 38.62)	0.04 (-0.18, 0.27)	0.77 (0.35, 1.67)	0.97 ( 4.2 2.16)	1.41 (0.90, 2.23)	1.77 (1.08, 2.91)	
22	Total pregnancy	-15.02 (-49.33, 19.30)	0.00 (-0.15, 0.15)	1.08 (0.64, 1.80)	1.28 🙇 .7 🕏 2.18)	1.21 (0.90, 1.63)	1.60 (1.15, 2.23)	
A 11 - : : C:		.1.1			<u> </u>			

2All significant findings in the table are bold.

2 All significant findings in the table are bold.

2 Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births, and adjusted for the other air pollutant similar technologies.

eTable 7 Comparison of major confounders and outcomes for those with missing and non-missing outcome data and non-missing confounder data

	Excluded from analyses*	Included in analyses	p-value
	(n=256)	(n = 1,017)	
Maternal age, mean ± SD	$28.47 \pm 0.22$	$28.78 \pm 0.11$	0.22
Gestational week, mean ± SD	$39.34 \pm 0.12$	$39.39 \pm 0.04$	0.69
Maternal BMI (kg/m <sup>2</sup> ), mean $\pm$ SD	$21.13 \pm 0.17$	$21.56 \pm 0.09$	0.03
Infant's sex			
Female, n (%)	136 (53.33)	527 (51.87)	0.676
Male, n (%)	119 (46.67)	489 (48.13)	
Primiparity			0.583
Yes, n (%)	226 (22.22)	791 (77.78)	
No, n (%)	61 (23.83)	195 (76.17)	
Season			0.00
Spring (Mar-May), n (%)	74 (32.46)	337 (35.62)	
Summer (Jun-Aug), n (%)	74 (32.46)	189 (19.98)	
Autumn (Sep-Nov), n (%)	55 (24.12)	143 (15.12)	
Winter (Dec-Feb), n (%)	25 (10.96)	277 (29.28)	
*Excluded from analyses due to missing co	gnitive outcome data		

## BMJ Open

Associations of air pollution exposures in preconception and pregnancy with birth outcomes and infant neurocognitive development: analysis of the Complex Lipids in Mothers and Babies (CLIMB) prospective cohort in Chongqing, China

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- 2 outcomes and infant neurocognitive development: analysis of the Complex Lipids in
- 3 Mothers and Babies (CLIMB) prospective cohort in Chongqing, China
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- 27 Abstract
- Objectives: To investigate the associations of traffic-related air pollution exposures in early
- 29 pregnancy with birth outcomes and infant neurocognitive development.
- **Design:** Cohort study.
- **Setting:** Eligible women attended six visits in the maternity clinics of two centres, the First
- 32 Affiliated Hospital of Chongqing Medical University and Chongqing Health Centre for
- Women and Children.
- Participants: Women who were between 20 and 40 years of age and were at 11–14 weeks
- 35 gestation with a singleton pregnancy were eligible for participation. Women were excluded if
- they had a history of premature delivery before 32 weeks of gestation, maternal milk allergy
- or aversion, or severe lactose intolerance. 1,273 pregnant women enrolled in 2015-2016 and
- 38 1,174 live births were included in this analysis.
- **Exposures:** Air pollution concentrations at their home addresses, including particulate matter
- 40 (PM) with diameter  $\leq 2.5 \mu m$  (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>), during pre-conception and
- 41 each trimester period were estimated using land-use regression models.
- **Outcome measures:** Birth outcomes (i.e., birthweight, birth length, preterm birth (PTB), low
- birth weight (LBW), large for gestational age (LGA) and small for gestational age (SGA)
- status) and neurodevelopment outcomes measured by the Chinese version of Bayley Scales of
- 45 Infant Development (CBSID).
- **Results:** An association between SGA and per-interquartile range (IQR) increases in NO<sub>2</sub> was
- found in the first trimester (odds ratio (OR): 1.57, 95% confidence interval (CI): 1.06, 2.32)
- and during the whole pregnancy (OR: 1.33, 99% CI: 1.01, 1.75). Both PM<sub>2.5</sub> and NO<sub>2</sub> exposure
- in the 90 days prior to conception were associated with lower Psychomotor Development Index
- 50 (PDI) scores (β: -6.15, 95% CI: -8.84, -3.46; β: -2.83, 95% CI: -4.27, -1.39, respectively).
- 51 Increased NO<sub>2</sub> exposure was associated with an increased risk of psychomotor development
- delay (PDD) during different trimesters of pregnancy.
- 53 Conclusions: Increased exposures to NO<sub>2</sub> during pregnancy were associated with increased
- risks of SGA and psychomotor development delay, while increased exposures to both PM<sub>2.5</sub>
- and NO<sub>2</sub> pre-conception were associated with adverse psychomotor development outcomes at
- 56 12 months of age.

**Keywords:** Air pollution; birth outcomes; child cognition



#### 59 Article summary

#### Strengths and limitations of this study

- This study uniquely explored the impacts of both pre-conception and prenatal exposure to PM<sub>2.5</sub> and NO<sub>2</sub> on neurodevelopmental outcomes in young infants, within an urban environment characterized by relatively high air pollution levels.
  - We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification.
- Our sample size was relatively small, limiting the statistical power to assess several outcomes.
- We defined exposure windows for clinically-defined trimesters; sensitive periods may be shorter or longer than 3 months, or may exist in the overlap of multiple trimesters.

#### INTRODUCTION

 Air pollution is a major environmental factor that has been linked to a range of adverse health outcomes in children. Maternal exposure to air pollutants during pregnancy, especially particulate matter (PM) with diameter  $\leq 2.5 \mu m$  (PM<sub>2.5</sub>) and nitrogen dioxide (NO<sub>2</sub>), has been found to be associated with adverse birth outcomes, including pre-term birth (PTB)(1), term low birth weight (TLBW) (2), and small for gestational age (SGA) status (3). According to the developmental origins of health and disease (DOHaD) hypothesis, prenatal exposures to air pollution may lead to adverse birth outcomes and subsequently increase the susceptibility to the development of certain diseases later in life (4). A number of epidemiological studies have linked prenatal air pollution exposure with neurodevelopmental disorders such as autism spectrum disorder (ASD), attention deficit hyperactivity disorder (ADHD), and cognitive impairment (5). Although the underlying biological mechanisms are still unclear, some studies indicated that prenatal air pollution exposure may induce systemic oxidative stress that triggers intrauterine inflammation, leading to damage to several fetal organs, including the brain (6, 7). It is also unclear that whether the adverse effects of air pollution may start earlier before conception. Three months before conception was considered as a critical developmental window for gametogenesis. Air pollution exposure during the three-month preconception period may have adverse effects on gametogenesis of sperm (8, 9) and ova cells (10). Exposures to PM<sub>2.5</sub> in preconception period have been associated with various neurodevelopmental outcomes, such as neural tube defects (11), lower psychomotor development scores (12), higher risk of ASD (13, 14), and higher risk of intellectual disability (15). Further research is required due to inconsistencies across studies in terms of studied health outcomes and exposure levels of air pollution (12). Additionally, while there is growing evidence for the effects of preconception PM<sub>2.5</sub> exposure on the risk of adverse neurodevelopmental outcomes, no study to date has examined the effects of preconception NO<sub>2</sub> exposure. Exposure to NO<sub>2</sub> during pregnancy may be linked to compromised neural development in children, particularly affecting fine psychomotor skills (16). Studying PM<sub>2.5</sub> along with NO<sub>2</sub> may allow us to explore how multiple pollutants affect birth outcomes and infant neurocognitive development independently and jointly. Moreover, both PM<sub>2.5</sub> and NO<sub>2</sub> are regulated traffic-related air pollutants in many countries. Understanding their impacts on birth and infant neurocognitive development can provide valuable insights for policymakers and public health authorities to develop effective air quality regulations and interventions.

Many studies have reported the effects of prenatal exposure to air pollution on neurodevelopmental function in children. However, the reported associations vary, due to the heterogeneous assessments of air pollution and neurodevelopmental outcomes (5, 17).

The current study leveraged the Complex Lipids in Mothers and Babies (CLIMB) cohort, a prospective birth cohort recruited in Chongqing, China (18), with trimester-specific maternal PM<sub>2.5</sub> and NO<sub>2</sub> air pollution exposure derived from a spatio-temporal land use regression (LUR) model (19). The aim of this analysis was to examine the associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposures during pre- and during pregnancy, with birth and infant neurocognitive development outcomes at 12 months of age.

A key aspect in all studies like this one is the accuracy of documenting exposure; a recent Chinese study determined air pollution exposure based on data from the nearest monitoring station (20) may not reflect the fine temporal and spatial variability of pollutant exposures among participants. Our study employed common air pollution exposure models based on advanced geographic information systems (GIS), to address some of the limitations of previous studies (21).

In addition, the timing of exposure is also critical in determining the effects of exposure on developmental outcomes. Indeed, the evidence from previous studies on the sensitive time windows for exposure pre- and during pregnancy remains inconclusive. Some studies have indicated that the early-to-mid pregnancy phase may be a critical period in terms of the impact of air pollution on neurodevelopment(22, 23). Early pregnancy is particularly important for neurogenesis and neuromigration, making it a susceptible period (24). However, some studies reported stronger associations for middle or late pregnancy (20, 25, 26). More studies identifying critical periods are needed to enhance our understanding of how pre-conception and prenatal air pollution exposure affect neurodevelopment. With this cohort, we are able to examine the effects of exposure pre-conception, at each trimester, and the entire pregnancy.

#### **METHODS**

#### **Study population**

- Participant recruitment in the CLIMB cohort has been described previously (27). In brief,
- women who were between 20 and 40 years of age and were at 11–14 weeks gestation with a
- singleton pregnancy were eligible for participation. Women were excluded if they had a self-
- stated history of premature delivery before 32 weeks of gestation, maternal milk allergy or
- aversion, or severe lactose intolerance.
- From September 2015 to November 2016, a total of 1,500 women were recruited into the cohort.
- Participants attended six visits at the First Affiliated Hospital of Chongqing Medical University
- and Chongqing Health Centre for Women and Children: 11–14 weeks' gestation (visit 1), 22–
- 28 week's gestation (visit 2), 32-34 week's gestation (visit 3), at birth (visit 4), 6 weeks
- postnatal (visit 5), and 12 months postnatal (visit 6).
- Women who withdrew from the study (n = 146), terminated their pregnancy (n = 29),
- miscarried (n = 12) or were lost to follow-up (n = 40) were excluded from the analysis, leaving
- a sample size of 1,273 women. Analyses were restricted to mothers whose detailed residential
- addresses during pregnancy were known (**Figure 1**). A total of 1,174 live births were thus
- included in the pregnancy and neonatal outcomes analysis. Subsequently, at 1 year follow-up,
- 146 946 children were included in the analysis of neurodevelopment outcomes.

#### Study setting

- The study area focused on the urban center of the Chinese municipality of Chongqing (**Figure**)
- 2). The terrain of Chongqing is predominantly hilly and mountainous, with the core area
- located in a synclinal valley at the confluence of the Yangtze River and the Jialing River (28).
- 151 The urban core of Chongqing, our study area, has a population of approximately 6.52 million
- people, a land area of 5,472 square kilometers, and 4.62 million vehicles (29). It shows a higher
- population density of approximately 1,191 people per square kilometer and a lower number of
- motor vehicles of 0.71 per capita. The urban core of Chongqing used to have multiple old
- industries with higher NO<sub>2</sub> and PM<sub>2.5</sub> emissions, including the Chongqing Iron and Steel
- 156 Company in Dadukou District and the Chongqing Thermal Power Plant in Jiulongpo District,
- both of which have been relocated to rural areas in Chongging. The main sources of pollution
- in the area now include traffic-related emissions, construction activities, and anthropogenic
- sources such as outdoor grilling and emissions from food establishments (30). The coverage

 rate of urban population with access to gas in Chongqing was 95.34% (29), suggesting a low reliance on biomass cookstoves in urban areas.

#### **Exposure assessment**

The address of participants was collected at the first visit. Exposure assessment based on spatiotemporal land use regression (LUR) models for PM<sub>2.5</sub> and NO<sub>2</sub> were developed for the study region. A description of the methodology of exposure modelling has been reported previously (19). Briefly, the models included both spatial and temporal components of exposure. PM<sub>2.5</sub> and NO<sub>2</sub> concentration data were collected from 17 routine monitoring sites operated by the Chongqing Environmental Monitoring Center in 2015-2016. For the spatial component of models, we calculated annual average concentrations of each pollutant in 2015, and fit linear regression models using five groups of geographic data (road network, land use, topography, vegetation, and population density) as spatial predictor variables. For the temporal component of models, we calculated the residuals from the spatial component at each monitoring site on a daily basis by subtracting the predicted annual average concentration from the observed daily average concentrations measured in 2015 and 2016, and then fitted generalised additive models (GAM) using seven groups of meteorological data (temperature, amount of rainfall, rainfall events, relative humidity, horizontal visibility, wind direction and wind speed) as temporal predictor variables. The meteorological variables were used to account for the influence of weather on the change in air pollution concentration over time. To account for the remaining spatial autocorrelation, the smoothed terms of longitude and latitude were fit to spatiotemporal residuals which were calculated by subtracting the sum of the spatial temporal predictions from the measured daily average concentrations in 2015 and 2016. The performance of the PM<sub>2.5</sub> spatiotemporal models was good (Correlation (COR)-R<sup>2</sup>: 0.72) and the NO<sub>2</sub> spatiotemporal model was low (COR-R<sup>2</sup>: 0.39) when providing concentration estimates in absolute terms.

Combining the family address coordinates of each pregnant woman and the gestation period of the pregnancy (calculated from the date of last menstrual period to the date of delivery), we used this spatiotemporal model to estimate the average exposure of each pregnant woman in 90 days prior to pregnancy (90D), first trimester (T1), second trimester (T2), third trimester (T3) and whole pregnancy period (WP), respectively.

#### **Outcomes**

#### Birth outcomes

#### Neurodevelopment outcomes

The Chinese version of Bayley Scales of Infant Development (CBSID) was used to assess mental and psychomotor development for infants in this study. The CBSID is appropriate for evaluation of infants from 2–30 months old (33) and takes into consideration each infant's age in days. Infants were assessed at around 12 months (range from 11 months and 15 days to 12 months and 15 days) by a trained examiner, with ages corrected for preterm birth. These scales have been formally adapted to the Chinese language and locally standardized to become culturally appropriate, with two main indexes: the Mental Development Index (MDI) and the Psychomotor Development Index (PDI). The MDI component comprised 163 items and assessed age-appropriate items related to cognitive functioning, personal and social development, and language development (see eTable 1 in the Supplement). The PDI component comprised 81 items and assessed age-appropriate fine and gross motor skills (see eTable 2 in the Supplement). The test provided raw scores for mental and psychomotor development that were converted to standardized (in terms of age in days) MDI and PDI scores, based on norms for the Chinese population. As with other forms of the Bayley test these index scores have a mean of 100, and a standard deviation of 15, with a lower score reflecting poorer performance (34). If an infant refused to cooperate with the examiners to finish the task, a second assessment was arranged within two weeks. If the infant could not cooperate at the second BSID assessment, their data were classified as missing. In addition to the continuous scores, we define mental developmental delay (MDD) and psychomotor developmental delay (PDD) if the score is less than 85 (35).

#### **Covariates**

Socio-demographic data were collected through interviews by trained nurses. The following potential confounders were identified: maternal age at enrolment (in years), infant sex (male/female), maternal BMI at 11–14 weeks' gestation (kg/m²), parity (Yes/No), monthly

 household income level (categorized as: <2,000 yuan, 2,000–7,000 yuan, 7,000–10,000 yuan, or >10,000 yuan), season of birth (categorized as: Spring (Mar-May), Summer (Jun-Aug), Autumn (Sep-Nov) or Winter (Dec-Feb)). Season of birth was taken into consideration because air pollution and related environmental factors, such as temperature and humidity, may vary across different seasons (i.e., air pollution levels tend to be higher during winter). Some studies suggest that the season of birth may indirectly influence cognitive function through factors such as seasonal differences in food availability affecting maternal nutrition during pregnancy, sunlight exposure impacting maternal vitamin D levels, and children's early-life indoor and outdoor activities. Marital status (Single/married) and smoking or drinking during pregnancy (Yes/no) were not taken into account in this analysis because of the homogeneity of the study population (i.e., 98.6% women were married and 99.6% women reported not smoking or drinking alcohol during pregnancy). We did not adjust dietary supplements during pregnancy because all pregnant women routinely take folic acid in this cohort.

#### Statistical analyses

Data were described in terms of mean ± SD or median (IQR) for continuous variables, or as percentages for categorical variables. Modelled PM<sub>2.5</sub> and NO<sub>2</sub> exposure levels in 90D, T1, T2, T3 and WP were considered separately. We examined the Spearman correlation between each of the exposures in the different pregnancy periods. For birth outcomes, multivariable linear regression was used for continuous outcomes (e.g., birth weight and birth length) to estimate β coefficient and their 95% confidence intervals (CIs) and multivariable logistic regression for binary outcomes (e.g., PTB, LBW, LGA and SGA status) to estimate odds ratio (OR) and 95% CIs. For mental and psychomotor development (e.g., MDI and PDI scores), multivariable linear regression models were fit to estimate β coefficient and their 95% CIs. We also conducted multivariable logistic regression analysis for binary neurodevelopment outcomes (i.e., MDD and PDD). Models were adjusted for maternal age at enrolment, infant sex, maternal BMI at 11–14 weeks gestation, primiparity, monthly household income level, and season of birth. We also ran co-exposure models to estimate associations of one air pollutant whilst additionally adjusting for the other air pollutant (i.e., PM<sub>2.5</sub> effects in T1 adjusted for NO<sub>2</sub> in T1). Effect estimates are reported for each IQR increase of PM<sub>2.5</sub> and NO<sub>2</sub>. All analyses were performed using STATA version 17. A p-value of <0.05 was considered statistically significant to address multiple comparisons in the analyses.

#### Patient and public involvement

 None.

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RESUL	TS
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#### Study participants

Participant characteristics are presented in **Table 1**. Of those participating women, the mean age was 28.7 years and mean BMI was 21.5 kg/m². 98.0% of women were of Han ethnicity, 77.9% were primiparous, and 67.6% had completed tertiary education. 33 (2.8%), 30 (2.6%), 108 (9.2%), 84 (7.2%) of the 1,174 births considered in this analysis were classified as PTB, LBW, LGA and SGA, respectively. For those 946 children who completed the BSID test, the mean MDI score was 94.7 (SD: 17.7) and the mean PDI score was 87.4 (SD: 14.9). The proportions of participants with MDD (MDI <85) and PDD (PDI < 85) were 27.1% and 42.4%, respectively.

#### **Exposure assessment**

Median  $PM_{2.5}$  exposure concentrations were 57.31  $\mu$ g/m³ (IQR: 5.76) and median  $NO_2$  exposure levels were 50.46  $\mu$ g/m³ (IQR: 5.51) during the whole pregnancy period (eTable 3 in the Supplement). For  $PM_{2.5}$ , the concentration in the pre-conception and T1 were considerably lower than other periods, close to  $10~\mu$ g/m³. The between-trimester and 90D values for  $NO_2$  were generally moderately correlated (Pearson's r > 0.5). The correlation coefficients of  $PM_{2.5}$  were more variable between time periods reflecting the high variability of  $PM_{2.5}$  concentrations, with values ranging from -0.78 to +0.68. Correlations between  $PM_{2.5}$  and  $NO_2$  in the same pregnancy period were moderately correlated (Pearson's  $r \sim 0.6$ , eTable 4 in the Supplement).

Table 1. Characteristics of study sample in the CLIMB cohort (N = 1,174)

6Characteristic of mother	N	n (%) / mean ± SD	Characteristic of child	N	$n$ (%) /mean $\pm$ SD
7Maternal age (Years)	1,174	$28.7 \pm 3.5$	Gestational week (week)	1,174	$39.4 \pm 1.5$
8BMI (kg/m²)	1,174	$21.5 \pm 2.9$	Birth weight (g)	1,165	$3314.4 \pm 428.8$
<sup>9</sup> Han ethnicity (%)	1,174		Birth length (cm)	1,149	$49.7 \pm 1.9$
<sup>10</sup> Yes		1,151 (98.0%)	New born sex	1,172	
1No		23 (2.0%)	Female	•	561 (47.9%)
Marital status (%)	1,174	, ,	Male		611 (52.1%)
15ingle		16 (1.4%)	Birth outcomes		
Married		1,158 (98.6%)	Preterm birth (PTB)	1,174	-
18rimiparity (%)	1,174	,	Yes		33 (2.8%) 1,141 (97.2%) 30 (2.6%) 1,141 (97.2%) 108 (9.2%) 1,066 (90.8%) 84 (7.2%) 1,090 (92.8%) 94.7 ± 17.7 87.4 ± 14.9 276 (27.1%) 741 (72.9%) 431 (42.4%)
1¾es		914 (77.9%)	No		1,141 (97.2%)
1\$\otimes_0		260 (22.1%)	Low birth weight (LBW)	1,174	3
<sup>19</sup> History of miscarriage or	1,174		Yes	•	30 (2.6%)
<sup>20</sup> abortion (%)					Ì
$2V_{es}$		553 (47.1%)	No		1,141 (97.2%)
22 No 23		621 (52.9%)	Large for gestational age	1,174	, , , , , , , , , , , , , , , , , , ,
			(LGA)	•	
24 28moking/drinking during	1,174		Yes		108 (9.2%)
28 regnancy (%)	,				
2¥es		5 (0.4%)	No		1,066 (90.8%)
2 <b>%</b> Io		1,169 (99.6%)	Small for gestational age	1,174	
29		, , ,	(SGA)	,	
3€ducation level	946		Yes		84 (7.2%)
<sup>3</sup> Low: High school or below		306 (32.3%)	No		1,090 (92.8%)
<sup>3</sup> High: College/uni or above		640 (67.6%)	BSID test	946	
33 ob 34 osewife 35 observed as the state of	946		MDI score		$94.7 \pm 17.7$
34 Full-time		762 (80.5%)	PDI score		$87.4 \pm 14.9$
35 Housewife		82 (8.7%)	Mental development	946	
39thers		102 (10.8%)	Delay (MDI < 85)		276 (27.1%)
3 Household income (Monthly)	946	,	Normal (MDI $\geq$ 85)		741 (72.9%)
392000 RMB		186 (19.7%)	Psychomotor Development	946	(1 11 1)
4 <b>2</b> 000-4000 RMB		329 (34.8%)	Delay (PDI < 85)	, .0	431 (42.4%)
44000-7000 RMB		292 (30.9%)	Normal (PDI $\geq$ 85)		586 (57.6%)
4 <del>2</del> 7000-10000 RMB		139 (14.7%)	Season of birth	1,174	
43		100 (11.770)	Spring (Mar-May)	-,-/ '	411 (35.01%) 263 (22.40%)
44			Summer (Jun-Aug)		263 (22.40%)
45			Autumn (Sep-Nov)		198 (16.87%)
46			Winter (Dec-Feb)		302 (25.72%)
<del>47</del> 48 280			( )		302 (25.72%)
70					2
49					5

#### Association with birth outcomes

In the unadjusted models (eTable 5 in the Supplement), higher exposure concentrations of PM<sub>2.5</sub> in T3 were significantly associated with lower birth length (β: -0.32, 95% CI: -0.51, -0.13; per IQR increase). We also observed increased NO<sub>2</sub> in T3 were significantly associated with lower birth length (β: -0.16, 95% CI: -0.32, -0.01; per IQR). A risk between SGA and increases in NO<sub>2</sub> (per IQR) was found in T2 (OR: 1.46, 95% CI: 1.10, 1.93), T3 (OR: 1.58,

201	93% C1. 1.14, 2.18) and in the whole pregnancy period (OK. 1.44, 93% C1. 1.13, 1.83). We
288	observed no evidence of associations of NO2 with overall birth weight, birth length and other
289	adverse birth outcomes (e.g., PTB, LBW, and LGA).
290	In the adjusted models ( <b>Table 2</b> ), we found increased effect size for NO <sub>2</sub> and SGA in T2
291	(OR: 1.57, 95% CI: 1.06, 2.32), and slightly reduced effects size for NO <sub>2</sub> and SGA in the
292	whole pregnancy period (OR: 1.33, 95% CI: 1.01, 1.75) compared with the unadjusted
293	model. We observed no evidence of associations with birth length in the adjusted models.
294	After co-adjustment for PM <sub>2.5</sub> (see eTable 6 in the Supplement), the association of NO <sub>2</sub>

with SGA was also found in T1 (OR: 1.70, 95% CI: 1.07, 2.69), T3 (OR: 1.77, 95% CI: 1.08,

Γ1 (C. nancy periou 2.91) and in the whole pregnancy period (OR: 1.60, 95% CI: 1.15, 2.23). 

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6		fference	o Odds ratios				
7	Per IQR increase in	Birth weight, grams	Birth length, cm	PTB (case: 33)	LBW <b>및</b> (유물: 30)	LGA (case: 108)	SGA (case: 84)
8	rei iQK iliciease ili	(95% CI)	(95% CI)	(95% CI)	(9 <b>8</b> % <u>(\$</u> I)	(95% CI)	(95% CI)
9 10		(N=941)	(N=927)	(N=945)	(1 <del>) (1)</del>	(N=945)	(N=945)
1Estimated	90 days prior to conception	59.73 (-16.52, 135.98)	0.15 (-0.176, 0.48)	0.24 (0.06, 1.00)	0.49 (8.38) 1.29)	1.40 (0.72, 2.71)	1.66 (0.75, 3.68)
12 exposure	First trimester	6.21 (-73.79, 86.20)	0.04 (-0.308, 0.388)	0.88 (0.28, 2.80)	0.76 (9.315 2.81)	0.86 (0.45, 1.67)	1.33 (0.58, 3.04)
13 <sup>1</sup> 14to PM <sub>2.5</sub>	Second trimester	-37.64 (-107.73, 32.44)	0.02 (-0.283, 0.326)	1.62 (0.53, 4.96)	1.34 (\$ 58 4.68)	1.00 (0.55, 1.83)	0.94 (0.50, 1.76)
15	Third trimester	4.20 (-73.17, 81.57)	-0.17 (-0.509, 0.162)	0.92 (0.29, 2.90)	0.92 (2.85)	1.29 (0.65, 2.53)	0.83 (0.42, 1.66)
16	Total pregnancy	8.01 (-41.10, 57.11)	0.02 (-0.198, 0.230)	0.77 (0.38, 1.54)	0.62 ( 1.25)	1.15 (0.75, 1.77)	0.84 (0.52, 1.35)
1Estimated	90 days prior to conception	-1.03 (-41.88, 39.81)	-0.04 (-0.215, 0.139)	0.84 (0.45, 1.57)	1.04 ( 1.98)	1.31 (0.91, 1.88)	1.45 (0.99, 2.12)
18 19exposure	First trimester	-9.78 (-50.84, 31.28)	0.04 (-0.133, 0.223)	0.90 (0.49, 1.65)	1.03 ( <b>‡.9</b> 6 1.91)	1.21 (0.85, 1.72)	1.57 (1.06, 2.32)
_	Second trimester	-20.82 (-59.11, 17.47)	-0.06 (-0.222, 0.112)	1.31 (0.73, 2.34)	1.34 (1.75 2.40)	1.21 (0.86, 1.70)	1.36 (0.95, 1.95)
20 to NO <sub>2</sub>	Third trimester	-9.50 (-56.00, 36.99)	-0.01 (-0.213, 0.191)	0.79 (0.40, 1.59)	0.95 ( <del>§</del> .4 <del>%</del> 1.94)	1.42 (0.94, 2.13)	1.51 (0.97, 2.36)
22 23	Total pregnancy	-8.45 (-37.73, 20.83)	0.00 (-0.125, 0.130)	0.97 (0.62, 1.51)	1.04 ( <b>2</b> .66 1.64)	1.20 (0.93, 1.56)	1.33 (1.01, 1.75)

2½ Il significant findings in the table are bold.
2½ Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

#### Association with infant neurodevelopment outcomes

In unadjusted models, PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower MDI and PDI scores in offspring (β: -3.54, 95% CI: -5.37, -1.71; β: -3.42, 95% CI: -4.96, -1.89) (**Table 3**). We also observed an unexpected positive association between PM<sub>2.5</sub> exposures in second trimester with MDI (β: 4.21, 95% CI: 2.43, 6.00) and PDI (β: 2.63, 95% CI: 1.12, 4.14). Exposure to NO<sub>2</sub> was associated with lower MDI (-1.90, 95% CI: -3.36, -0.44) and PDI in the 90 days prior to conception (-2.86, 95% CI: -4.08, -1.65). NO<sub>2</sub> exposure was also associated with lower PDI scores in T3 (-1.97, 95% CI: -3.29, -0.65) and in the whole pregnancy periods (-1.08, 95% CI: -2.11, -0.05). We did not observe any association between NO<sub>2</sub> and MDI in any pregnancy periods.

In the adjusted models (Table 3), we found PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower PDI scores (β: -6.15, 95% CI: -8.84, -3.46). Similarly, there was also a significant association of increased NO<sub>2</sub> exposure and lower PDI score in the 90 days prior to conception (β: -2.83, 95% CI: -4.27, -1.39), T1 (β: -1.91, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -2.57, -0.26) and whole pregnancy period (β: 1.15, 95% CI: -3.37, -0.46), T3 (β: 1.92, 95% CI: -3.37, -0.46), T3 (β: -1.91, 95% CI: -3.37

- also a significant association of increased NO<sub>2</sub> exposure and lower PDI score in the 90 days prior to conception (β: -2.83, 95% CI: -4.27, -1.39), T1 (β: -1.91, 95% CI: -3.37, -0.46), T3 (β: -1.92, 95% CI: -3.57, -0.26) and whole pregnancy period (β: -1.15, 95% CI: -2.19, -0.11). The positive association between PM<sub>2.5</sub> exposures in second trimester with PDI (β: 3.76, 95% CI: 1.27, 6.24) remained. We did not observe any association with MDI in any pregnancy periods.
- In the co-exposure models (**Table 3**), PM<sub>2.5</sub> exposure in the 90 days prior to conception was associated with lower PDI scores (β: -4.74, 95% CI: -7.73, -1.75). We also observed a positive association between PM<sub>2.5</sub> exposures in second trimester with PDI (β: 5.51, 95% CI:2.73, 8.28).
- Exposure to NO<sub>2</sub> was significantly associated with lower PDI in 90D (β: -1.72, 95% CI: -3.31,
- 320 -0.12), T1 (β: -1.80, 95% CI: -3.46, -0.15), T2 (β: -2.11, 95% CI: -3.63, -0.60), T3 (β: -1.92,
- 95% CI: -3.76, -0.09) and whole pregnancy period (β: -1.68, 95% CI: -2.89, -0.46).

Table 3. Associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposure in different pregnancy periods and continuous SSID scores

PDI 95% CI

(N=946)

-3.42 (-4.96, -1.89)

0.04 (-1.52, 1.61)

2.63 (1.12, 4.14)

-1.76 (-3.42, -0.10)

0.5 (-0.82, 1.83)

-2.86 (-4.08, -1.65)

-1.17 (-2.43, 0.08)

0.00 (-1.18, 1.18)

-1.97 (-3.29, -0.65)

-1.08 (-2.11, -0.05)

Crude models

MDI (95% CI)

(N=946)

-3.54 (-5.37, -1.71)

-1.07 (-2.93, 0.79)

4.21 (2.43, 6.00)

-1.43 (-3.41, 0.55)

1.64 (0.06, 3.21)

-1.90 (-3.36, -0.44)

-0.08 (-1.57, 1.42)

1.81 (0.41, 3.22)

0.04 (-1.54, 1.62)

0.67 (-0.56, 1.89)

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(N=945) in se

-6.15 (-8.84, \$\frac{\text{m}\_{10}}{25} \frac{\text{R}}{6}

-2.11 (-4.95, **3**)

3.76 (1.27, 62)

-1.37 (-4.12, **a b** 

0.23 (-1.52, 7.28)

-2.83 (-4.27, 513**3**5)

-1.91 (-3.37, ₺₧₺

-0.75 (-2.11,**3**).6

-1.92 (-3.57, **≥**0.2€)

 $-1.15 (-2.19, \frac{1}{2}).12)$ 

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Adjusted models\*

MDI (95% CI)

(N=945)

-1.98 (-5.19, 1.23)

-1.66 (-5.02, 1.70)

3.79 (0.85, 6.73)

-2.73 (-5.99, 0.53)

-0.27 (-2.34, 1.80)

-0.72 (-2.43, 1.00)

0.59 (-1.14, 2.32)

0.56 (-1.05, 2.17)

0.51 (-1.45, 2.47)

0.41 (-0.83, 1.64)

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Co-exposure models\*\*

PDI 95% CI

(N=945)

-4.74 (-7.73, -1.75)

-0.45 (-3.67, 2.76)

5.51 (2.73, 8.28)

0.04(-3.02, 3.09)

1.69 (-0.35, 3.73)

-1.72 (-3.31, -0.12)

-1.80 (-3.46, -0.15)

-2.11 (-3.63, -0.60)

-1.92 (-3.76, -0.09)

-1.68 (-2.89, -0.46)

MDI (95% CI)

(N=945)

-1.73 (-5.30, 1.85)

-2.84 (-6.65, 0.97)

4.19 (0.89, 7.49)

-3.84 (-7.46, -0.22)

-0.85 (-3.28, 1.57)

-0.31 (-2.22, 1.60)

1.28 (-0.68, 3.24)

-0.48 (-2.28, 1.33)

1.52 (-0.66, 3.69)

0.67(-0.77, 2.12)

2

6

Per IQR increase in

First trimester

Second trimester

Third trimester

Total pregnancy

First trimester

Second trimester

Third trimester

Total pregnancy

90 days prior to conception

90 days prior to conception

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Estimated

1<sub>exposure</sub>

13to PM<sub>2.5</sub>

14 15

16 17 15 15 15

1&exposure

20 to NO2

23All significant findings in the table are bold.

24 Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births.

\*Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly house gold income level, and season of births, and adjusted for the other air pollutant.

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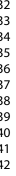
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In the adjusted model, the risk of PDD was found to increase by 112% and 42% with each per-IQR increase in PM<sub>2.5</sub> (OR: 2.12, 95% CI: 1.45, 3.11) and NO<sub>2</sub> (OR: 1.42, 95% CI: 1.16, 1.75) in the 90 days prior to conception (**Table 4**). There was also a significant association between increased NO<sub>2</sub> exposure and the risk of PDD in T1 (OR: 1.29, 95% CI: 1.05, 1.58), T3 (OR: 1.27, 95% CI: 1.01, 1.60), and the whole pregnancy period (OR: 1.17, 95% CI: 1.02, 1.36). We did not observe any association with MDD in any pregnancy periods.



1 2 3 4	330 331	Table 4. Associations	s between PM <sub>2.5</sub> and NO <sub>2</sub>		MJ Open  nt pregnancy periods	by copyright, incading and mental	psychomotor develop	mental
5 6	331	delay				g for	S S	
<del>7</del> 8	3		Crude	Crude models		nodels* 💆 📆	£	ıre models**
9		Per IQR increase in	MDD (95% CI)	PDD (95% CI)	MDD (95% CI)	PDD (95% @1)	MDD (95% CI)	PDD (
10	10 Fei iQK increase in		(N=946)	(N=946)	(N=945)	(N=94 <b>)</b>	(N=945)	(N=

8		Crude models		Aujusteu i	110dc12 6 3.4	Co-caposure	inoucis
9	Per IQR increase in	MDD (95% CI)	PDD (95% CI)	MDD (95% CI)	PDD (95% 60)	MDD (95% CI)	PDD (95% CI)
10	rei iQK increase in	(N=946)	(N=946)	(N=945)	(N=94 <b>)</b> 12 12	(N=945)	(N=945)
Estimated	90 days prior to conception	1.45 (1.16, 1.83)	1.49 (1.20, 1.83)	0.95 (0.64, 1.42)	2.12 (1.45, 3.3)	0.97 (0.63, 1.51)	1.78 (1.17, 2.71)
13exposure	First trimester	1.05 (0.83, 1.33)	1.04 (0.84, 1.28)	1.14 (0.73, 1.79)	1.42 (0.96, <b>ឌ្គី</b> :ជុំត្តិ	1.35 (0.80, 2.25)	1.16 (0.74, 1.82)
14 15 <sup>to</sup> PM <sub>2.5</sub>	Second trimester	0.63 (0.49, 0.80)	0.77 (0.63, 0.95)	0.81 (0.54, 1.22)	0.72 (0.51, <b>a a a</b>	0.83 (0.52, 1.31)	0.57 (0.38, 0.85)
	Third trimester	1.23 (0.96, 1.58)	1.19 (0.95, 1.49)	1.25 (0.82, 1.90)	1.17 (0.80, <b>2 2 3</b>	1.39 (0.87, 2.23)	0.98 (0.64, 1.49)
16 17	Total pregnancy	0.84 (0.69, 1.03)	0.98 (0.82, 1.18)	1.07 (0.82, 1.39)	1.07 (0.84, 🗓 🌋 💆	1.17 (0.86, 1.59)	0.90 (0.68, 1.20)
1 <b>E</b> stimated	90 days prior to conception	1.20 (0.99, 1.45)	1.41 (1.19, 1.67)	0.97 (0.77, 1.21)	1.42 (1.16, ] 📆	0.97 (0.76, 1.25)	1.24 (0.99, 1.56)
19 exposure	First trimester	0.97 (0.80, 1.17)	1.18 (0.99, 1.40)	0.91 (0.72, 1.13)	1.29 (1.05, <b>ā</b> .5 <b>8</b>	0.84 (0.65, 1.09)	1.24 (0.99, 1.57)
20 P 21 to NO <sub>2</sub>	Second trimester	0.79 (0.66, 0.95)	1.04 (0.88, 1.22)	0.94 (0.76, 1.15)	1.14 (0.95, <b>¾</b> .3	0.98 (0.77, 1.24)	1.31 (1.06, 1.63)
22	Third trimester	1.04 (0.85, 1.28)	1.25 (1.04, 1.50)	0.94 (0.73, 1.21)	1.27 (1.01, 🖺 .6🤁	0.86 (0.65, 1.14)	1.28 (0.99, 1.65)
23	Total pregnancy	0.92 (0.79, 1.07)	1.16 (1.01, 1.33)	0.94 (0.80, 1.11)	1.17 (1.02, <b>\bar{3</b> .3 <mark>\bar{6}</mark>	0.9 (0.75, 1.08)	1.21 (1.02, 1.43)

All significant findings in the table are bold.

25 Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births. 27\*Models adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly has been been and season of births, and

2adjusted for the other air pollutant.

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#### **DISCUSSION**

We analyzed associations between modelled PM<sub>2.5</sub> and NO<sub>2</sub> pre- and during pregnancy with birth and neurodevelopment outcomes in singleton children born in a south-western metropolis of China in 2015-16. We found the likelihood of SGA increased by 33% per IQR higher exposure to NO<sub>2</sub> in the whole pregnancy periods after adjusted for maternal age at enrolment, infant's sex, maternal BMI at 11–14 weeks' gestation, primiparity, monthly household income level, and season of births and PM<sub>2.5</sub>. For childhood cognitive development, increased exposure to PM<sub>2.5</sub> and NO<sub>2</sub> in the 90 days prior to conception were both associated with lower PDI scores, with the effect size per IQR being higher for PM<sub>2.5</sub> than for NO<sub>2</sub>. Increased NO<sub>2</sub> exposure was associated with an increased risk of PDD during different trimesters of pregnancy.

Many studies from other geographic areas, including Europe (36-38), the United States (22, 26), and Asia (23, 39-41), have found significant associations between prenatal air pollution exposure and a variety of adverse neurodevelopmental outcomes. Our finding of a negative association between prenatal NO<sub>2</sub> air pollution exposure and infant neurocognitive development is consistent with these reports. A recent Chinese birth cohort study of 15,778 child-mother pairs in Foshan reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with an increased risk of suspected developmental delay (OR: 1.06, 95% CI: 0.94, 1.19) measured by a five-domain scale and developmental quotient (DQ) (23). A birth cohort study of 520 mother-child pairs in South Korea reported that maternal NO<sub>2</sub> exposure during pregnancy was associated with impairment of psychomotor development ( $\beta = -1.30$ , p = 0.05) but – as in the present study - not with cognitive function ( $\beta = -0.84$ , p = 0.20) (39). However, results from previous research varied by air pollutants. For example, a Chinese study of 1193 mother-newborn pairs in Changsha found significant associations between PM<sub>2.5</sub> exposure in trimester two and lower neurobehavioral developmental scores, while other air pollutants such as PM<sub>10</sub>, carbon monoxide (CO), and Sulfur dioxide (SO<sub>2</sub>) had null or even reverse associations. In this study, we observed that the negative effect of NO<sub>2</sub> exposure during pregnancy on PDI is significant at 5% level; this negative effect of NO<sub>2</sub> still remained after adjustment for PM<sub>2.5</sub>. This heterogeneity may relate to the temporality of exposure assessment, types of outcome assessment instruments or evaluators, and the levels of air pollution. In addition, air pollution mixtures may have differed among the study regions, thus there are several potential explanations for the heterogeneity of the findings. We also observed negative correlations between certain exposures, indicating the need to consider potential collinearity in our twopollutant models. In Chongqing, a major industrial city in southwest China, air pollution may

come from industrial and traffic emissions, construction activities, and dust, and negative correlations may occur if different sources contribute disproportionately to each pollutant. Their correlations may also be affected by seasonal changes and variations in weather patterns.

Future research should also explore the impact of source-specific air pollution on children's

cognitive health.

To date, most studies on prenatal air pollution exposure and child neurodevelopment have been conducted in developed countries with relatively low levels of air pollution. In this study, the level of air pollution was higher (median  $PM_{2.5}$ : 57.31 µg/m³, IQR: 5.76; median  $NO_2$ : 50.46 µg/m³, IQR: 5.51) compared to studies in developed countries such as Europe and the United States. In a multi-centre European cohort, the mean  $PM_{2.5}$  and  $NO_2$  exposure concentration during pregnancy were 13.4 µg/m³ and 11.5 µg/m³ (36). Researchers found that the psychomotor development score significantly decreased by 0.68 points (95% CI: -1.25, -0.11) for every 10 µg/m³ increase in  $NO_2$ , and there was also a non-significant decrease of 1.64 points (95% CI: -3.47, 0.18) for every 5 µg/m³ increase in  $PM_{2.5}$  during pregnancy (36). Factors such as the types of pollutants and concentrations may differ between China and other regions with a lower air pollution level, leading to variations in the observed effects.

Contrary to expectations, we found significant positive associations between prenatal exposure to PM<sub>2.5</sub> air pollution in the second trimester and PDI. However, no association was observed between PM<sub>2.5</sub> exposures in the second trimester and the risk of PDD. Given the existing literature and the conflicted observation here, we believe that this is likely to be spurious/sample specific. Some plausible explanations include the uneven distribution of PDI scores, the potentially inappropriate selection of the cut-off value of 85 (which may not effectively discriminate between groups), or the possibility that the observed outcome occurred by chance. Several epidemiological studies have reported associations between prenatal exposure to high levels of PM<sub>2.5</sub> and lower neurodevelopment in children ranging in age from 6 months to 6 years (12, 37, 42-44). In agreement with our findings, a multi-centre cohort study from six European countries investigated the effects of prenatal exposure to multiple air pollutants including PM<sub>2.5</sub>, PM<sub>10</sub>, coarse particles, NO<sub>2</sub> and nitrogen oxides (NOx) among 9482 children between 1 and 6 years; the authors found nonsignificant positive associations between prenatal  $PM_{2.5}$  exposure and normal neurodevelopment ( $\beta$ : 1.64, 95% CI: -3.47, 0.18; per 5 μg/m<sup>3</sup> increase in PM<sub>2.5</sub>) (36). Similarly, another study examining the effects of multiple pollutant exposures on early childhood cognition at 40 days of age in a highly exposed area of Spain also found PM<sub>10</sub>, PM<sub>coarse</sub>, PM<sub>2.5absorbance</sub>, NO<sub>2</sub>, NO<sub>x</sub>, and Ozone (O<sub>3</sub>) were linked to lower

 motor function in children, except for PM<sub>2.5</sub> (45). The inconsistent findings could be because of heterogeneity between studies in terms of exposure (e.g., exposure assessment methods used, PM<sub>2.5</sub> exposure levels, or composition of PM<sub>2.5</sub>).

The prevalence of MDD and PDD in our study is higher than in other studies that also used the CBSID to report developmental delay rates, which were at 17% (46), 15.78% (47), and 13.68% (48). This may be attributed to the younger age of infants in our study, which were assessed at around 12 months, compared to most studies assessed at around 24 months. A Chinese study and a South Korean study also found lower scores on the MDI and PDI in 1-year-old children (49, 50). Aside from the conflicting findings regarding prenatal PM<sub>2.5</sub> exposure and neurodevelopmental outcomes, results regarding the most potential sensitive time windows before and during pregnancy are also inconclusive. Some studies suggested that early-to-mid pregnancy might be a potential sensitive period (22, 23), while other studies found stronger associations for middle-to- late pregnancy, thus results are equivocal (20, 25, 26).

The potential biological mechanisms by which air pollution could affect neurodevelopment are not yet clearly understood. There is evidence suggesting that exposure to prenatal PM<sub>2.5</sub> could potentially induce maternal immune activation during pregnancy (51). Higher levels of cytokines or reactive oxygen species may potentially interfere fetal neurodevelopment through three mechanisms: crossing the placental barrier into the fetal body, inducing fetal immune dysregulation, and contributing to inadequate placental perfusion that affects nutritional processes and oxygenation of maternal blood (52). More research is needed to investigate trimester effects of air pollution on neurodevelopment and provide better understanding on the underlying biological mechanisms. Our study is the first to consider an exposure window 90 days prior to conception for NO<sub>2</sub>. A novel observation is that effects of NO<sub>2</sub> or PM<sub>2.5</sub> air pollution on child cognition can be seen at least 90 days prior to conception, representing a potentially vulnerable periods in relation to air pollution on neurodevelopment. Similar results were found in previous study recruited 1329 mother-child pairs in Wuhan, China (12). This study reported a higher level of PM<sub>2.5</sub> during preconception (Median: 76.1 µg/m<sup>3</sup>) and in the first trimester (Median: 82.3 μg/m<sup>3</sup>). This study found for each doubling of PM<sub>2.5</sub> exposure during preconception, children's PDI scores was reduced by 6.15 (95% CI: -8.84, -3.46) points. A potential explanation is that preconception air pollution exposures induce genetic and epigenetic alterations in sperm, that increase the risk of adverse health outcomes in offspring (53, 54). To date, all studies examined the effect of maternal preconception exposure while

omitting paternal exposures (17). Future studies should consider the effect of preconception paternal exposure in relation to childhood health outcomes.

This study has several strengths. We developed an LUR model to capture spatial and temporal variations of air pollution at individual level to reduce exposure misclassification if using monitoring stations. This is a novel study to investigate both pre-conception and prenatal PM<sub>2.5</sub> and NO<sub>2</sub> exposure with neurodevelopment outcomes among young infants, in the context of a relatively high air pollution urban environment. The exposure levels in our study were similar as those in comparable urban areas in Chinese cities. A study in Shanghai, China reported an average NO<sub>2</sub> exposure during pregnancy from 2014 to 2015, predicted by the LUR model, of 48.23 μg/m³ (Mean PM<sub>2.5</sub> in our study: 50.52 μg/m³) (55). Similarly, a study in Tianjin found the annual average PM<sub>2.5</sub> exposure to be 62 μg/m³ in 2017 (Mean NO<sub>2</sub> in our study: 57.48 μg/m³) (56). Wu et al. developed a LUR model for PM<sub>2.5</sub> in the main urban area of Chongqing (57). This model predicted an annual average PM<sub>2.5</sub> concentration of 40.6 μg/m³ (57), whereas our prediction is higher at 55.9 μg/m³ (19). This difference can be attributed to the temporal variations. Wu et al. used monitoring data from 2013, while we utilized data from 2015. It could be considered that our GAM model, with its temporal component, could explain temporal variations and is more suitable for pregnancy-specific exposure estimates.

A major limitation of this study was that our sample size was relatively small, limiting the statistical power to assess several outcomes, although the higher exposures in Chongqing than in some other studies may increase probability of detecting effects. In terms of limitations, due to a lack of information on participant time-activity patterns, exposure estimates in this study refer only to ambient concentrations at home addresses, and no other activity spaces (e.g., indoor, workplace, commuting) were considered. We may have thus underestimated total air pollution exposure. Second, we defined exposure windows for clinically-defined trimesters; sensitive periods may be shorter or longer than 3 months, or they may exist in the overlap of multiple trimesters. However, we were unable to investigate the sensitive time windows using established methods such as distributed lag non-linear models due to the lack of highly timeresolved air pollution estimates. Third, the performance of the NO<sub>2</sub> spatiotemporal model was low (COR-R<sup>2</sup>: 0.39), which may introduce exposure misclassification and therefore bias in the coefficients. It may lead to underestimation of the association if the NO<sub>2</sub> spatiotemporal model inadequately represents the true variability in NO<sub>2</sub> levels. Or conversely, it could overestimate the association between NO<sub>2</sub> exposure and the outcome if the model fails to account for certain factors or inaccurately estimates NO<sub>2</sub> levels. Finally, we were unable to include some other air

 pollutants such as polycyclic aromatic hydrocarbons (PAH), black carbon (BC) and Ozone, which have bene found particular harmful to neurodevelopment in children (58). Although we have accounted for most of the important confounders in this study, unfortunately, we did not collect information on the feeding patterns of infants. This may undermine the validity and reliability of our findings.

#### CONCLUSION

- This study provides evidence for an association between NO<sub>2</sub> exposure prior to- and during pregnancy with birth and neurodevelopmental outcomes in a birth cohort in Chongqing, China.

  Fyposure to NO<sub>2</sub> and PM<sub>3</sub>, exposure before pregnancy was associated with a lower
- Exposure to NO<sub>2</sub> and PM<sub>2.5</sub> exposure before pregnancy was associated with a lower psychomotor development score. Increased NO<sub>2</sub> exposure was linked to a risk of psychomotor development delay during various pregnancy trimesters.

#### **Declarations**

#### Ethics approval and consent to participate

- Ethical approval for this study was granted by the Ethics Committee of Chongqing Medical
  University (#2014034). The participants provided their written informed consent to
- participate in this study. Written informed consent was obtained from the individual(s) for the
- publication of any potentially identifiable images or data included in this article.

### **Data availability statement**

- The data that support the findings of this study are available from Chongqing Medical
- 485 University but restrictions apply to the availability of these data, which were used under
- license for the current study, and so are not publicly available. Data are however available
- from the authors upon reasonable request and with permission of Chongqing Medical
- 488 University.

#### 489 Competing interests

The authors declare that they have no conflicts of interests.

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- The views expressed are those of the author(s) and not necessarily those of the NIHR, the
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#### 503 Contributors

- Y. X., T.L.H., H.Z. and PNB. conceived and designed research; T.Z., Y.X. and H.Z. recruited
- the patients and collected the samples; T.K., A.H., and J.G. constructed the air pollution
- model; Y.C. analyzed, interpreted the data and prepared the figures; Y.C and T.K were major
- 507 contributors in writing the manuscript text; YSC, JC, TLH, YX, ALH, and PNB substantively
- revised the manuscript; All authors read and approved the final manuscript.

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- 690 Figure 1. Flow chart of the study population in CLIMB
- Figure 2. Study area and location of monitoring sites (OpenStreetMap contributors,
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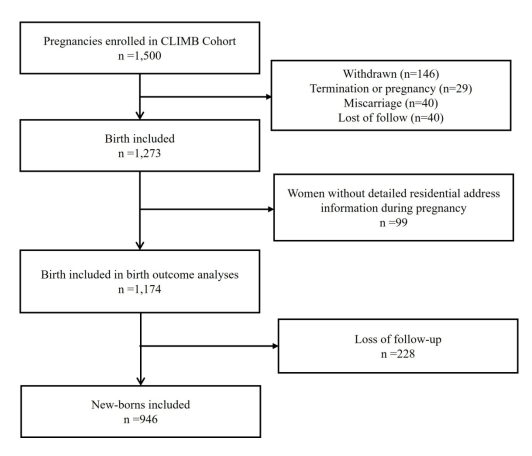


Figure 1 Flow chart of the study population in CLIMB 173x144mm (300 x 300 DPI)

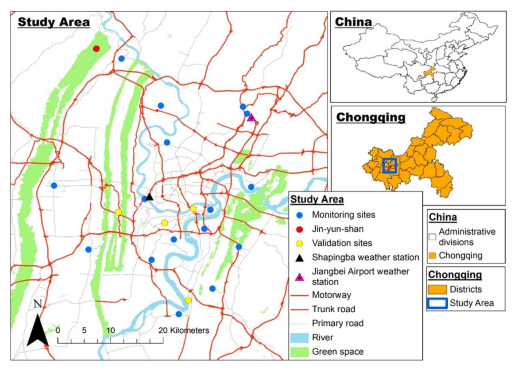


Figure 2 Study area and location of monitoring sites (OpenStreetMap contributors, 2015; https://data.nextgis.com/en/region/CN-50/).

159x112mm (220 x 220 DPI)

Associations of air pollution exposures in preconception and pregnancy with birth outcomes and infant neurocognitive development: analysis of the Complex Lipids in Mothers and Babies (CLIMB) prospective cohort in Chongqing, China

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## **Supplement**

## eTable 1 Mental Development Index (Chinese version)

## 智力量表

(\*可偶尔观察到)

序号	月龄	条目	计分
1	0.1	对铃声反应	
2	0.1	抱起时安静	
3	0.1	对摇鼓声反应	
4	0.1	对尖声反应: (电灯开关)	
5	0.1	短暂地注视红环	
6	0.2	短暂地注视人	
7	0.4	稍长时间地注视红环	
8	0.5	眼的水平协调活动(红环)	
9	0.7	眼的水平向天活动 (光)	
10	0.7	眼睛追随移动的人	
11	0.7	对说话声反应	
12	0.8	眼的垂直协调活动 (光)	
13	0.9	发声一至两次	
14	1	眼的垂直协调活动(红环)	
15	1.2	眼的旋转协调活动(光)	
16	1.2	眼的旋转细条活动(光环)	
17	1.3	*自由环视周围	
18	1.5	社交笑: 测试者谈话与微笑时	
19	1.6	眼转向红环	
20	1.6	眼转向光	
21	1.6	*发声至少四次	
22	1.7	期待性兴奋	
23	1.7	对面部的纸有反应	
24	1.9	能用视觉辨认母亲	
25	1.9	社交笑: 测试者微笑与安静时	
26	2	*对测试者的微笑和说话有发声反应	
27	2.1	*用眼睛寻找声源(详细说明)	
28	2.2	*发出两种不同的声音	
29	2.2	对手的遮蔽眨眼	
30	2.2	对面孔的消失有反应	
31	2.4	注视方木	
32	2.6	从一物转看另一物	
33	2.6	眼睛追随铅笔	
34	2.7	对抱起有预感性的调节反应	
35	2.9	目光追随横过桌面的球	
36	2.9	头追随悬摆的环	

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37	3.1	头追随逐渐消失的勺子
38	3.2	操作红环
39	3.3	简单地玩摇鼓
40	3.4	*轻轻地抚摸桌沿
41	3.4	*意识到陌生环境
42	3.5	头转向铃声
43	3.6	头转向摇鼓声
44	3.6	*手碰手的玩耍
45	3.6	将红环送入口中
46	3.7	伸手够悬环
47	3.8	看自己的手
48	4.2	接近悬环 (优势手)
49	4.4	*发声时的姿态(描述)
50	4.4	*主动抚摸桌沿
51	4.4	接近镜像
52	4.4	注意小糖丸
53	4.6	伸手取方木
54	4.7	喜欢嬉戏
55	4.9	伸手时眼手协调
56	4.9	拾起方木 (优势手)
57	5	保持两块方木
58	5	持久地看红环
59	5	头部跟着掉下的勺转动
60	5	探索性地玩纸
61	5	对镜像微笑
62	5	坚持够东西
63	5.1	在小床内重新找到摇鼓
64	5.1	*辨别生人
65	5.4	举起倒扣的茶杯
66	5.5	*敲打玩耍
67	5.5	探索性地玩细绳
68	5.5	伸手取第二块方木
69	5.6	*由一手向另一手传递物体
70	5.8	*对产生响声感兴趣
71	5.9	灵巧而直接地拾起方木
72	6	*对镜像开玩笑
73	6	用把柄举起茶杯
74	6	寻找掉落的勺子
75	6.1	牵拉细绳获取红环
76	6.1	保留三块方木中的两块
77	6.6	*发出四个不同的音节
78	6.8	能配合玩游戏
79	7	恰当地牵拉细绳获取红环
80	7.1	玩摇铃, 对细节感兴趣
81	7.4	企图获得三块方木
31	1	

Ī	I	1
82	7.4	有目的地摇铃
83	7.5	*选择性地倾听熟悉的词语
84	8	*对 da-da 或类同词
85	8.1	暴露玩具
86	8.2	注意测试者的乱写
87	8.3	将手指插入桩板洞中
88	8.6	观看书中图画
89	8.9	对他人的言语要求有反应
90	9.1	拿起茶杯获得方木
91	9.8	寻找盒子里面的东西
92	10.3	遵照命令将方木放入茶杯 (放入数)
93	10.7	企图模仿乱写
94	10.8	模仿用勺子搅拌
95	10.9	遵照命令停止
96	10.9	推动小汽车
97	11	模仿地拍打哨娃
98	11.1	*重复引入发笑的把戏
99	11.2	解开裹着的方木
100	11.2	将三块方木放入杯中
101	11.4	*快速而不清的表达
102	11.4	揭开兰盒子的盖
103	11.5	翻开书页
104	11.5	摇晃悬环的
105	11.8	将骰子放入盒中(6个)
106	12	恰当地握持画笔
107	12.2	模仿说单词(记录用过的词)
108	12.4	重复地插一根桩钉
109	12.5	用手势表达想要的东西
110	12.9	自动乱写
111	12.9	能说两个词
112	13	搭两层塔
113	13.1	出示鞋子或其他衣服或自己的玩具
114	13.2	从瓶中移出小糖丸
115	13.3	掺九块方木放入杯中
116	14.3	*盖上圆盒
117	14.4	
118	14.8	用棍子够取玩具
119	15.4	搭三层塔
120	15.7	在 70 秒钟内插完桩钉
121	16.1	指出娃娃身体的各部分:三个部位以上
122	16.3	粉红模板:放置圆形模块
123	16.6	<ul><li></li></ul>
124	17.2	用笔模仿画一划
125	17.5	在 42 秒钟内插完桩钉
126	17.6	说出一物名
	1	

127	177	对桩桩执行论会(方语计划位行约、技术在)
	17.7	对娃娃执行指令(在通过部位打钩:椅、杯、鼻)
128	18.1	用语言表达要求
129	18.6	不用于一划的乱写?
130	18.8	兰色模板: 放置两个圆快和方块
131	18.8	指出三幅画
132	19.1	能说两个单词的句子
133	19.2	说出一副画名
134	19.2	
135	19.3	找出两物
136	19.8	在 30 秒钟内插完桩钉
137	20.4	粉红模板: 完成
138	20.4	搭六层塔
139	20.5	兰色模板,放置六个模块
140	21	指出五副画
141	21.1	说出三物名
142	21.2	勉强合格地安装破娃娃
143	21.2	区别两物:杯、盘、盒
144	22.8	辨认钟表: 第四张图 1, 2, 3, 4, 5
145	22.9	说出三幅画名
146	23.8	粉红模板(反转)
147	24.3	近似地安装破娃娃
148	24.6	区别三物:杯、盘、盒
149	24.7	兰色模板,在 150 秒钟内完成
150	25	搭八层塔
151	25.1	指出七副画
152	25.1	用方木搭火车
153	25.7	说出五副画名
154	26.3	模仿笔划:垂直线和水平线
155	27.1	辨认钟表: 第2张图
156	27.6	理解两个方位词
157	28	在 22 秒钟内插完桩钉
158	28.5	兰色模板: 90 秒钟内完成
159	29.5	折纸
160	29.6	
161	30+	正确安装破娃娃
162	30+	"—"的概念
163	30+	理解三个方位词

## 运动量表

(\*可偶尔观察到,△可在施测智力量表时观察到)

<b>ċ</b> □	□ #A	A 17	>1 A
序号	月龄	条目	计分
1	0.1	抱起靠肩时抬头	
2	0.1	抱起靠肩时调整姿势	
3	0.1	侧头	
4	0.1	爬起	
5	0.8	△保留红环	
6	0.8	*伸臂玩耍	
7	0.8	*踢腿玩耍	
8	0.8	头起竖起: 垂直位	
9	1.6	头部稳定地竖起	
10	1.7	抬头 (背悬位)	
11	1.8	由侧卧转向仰卧	
12	2.2	在俯卧位时用双臂撑起自己	
13	2.2	支撑下坐起	
14	2.5	保持头部稳定	
15	2.6	*双手张开占优势	
16	3.3	头平衡	
17	3.4	*尺侧一手掌抓握方木	
18	3.5	轻度支撑坐位	
19	4.3	*由仰卧转向侧卧	
20	4.7	努力想坐起	
21	5.0	部分的拇指相对(桡侧一手掌)拾起方木	
22	5.1	独坐片刻	
23	5.1	*单手抽取	
24	5.2	*转腕	
25	5.2	牵拉坐起	
26	5.6	△试图获取小糖丸	
27	5.7	独立 30 秒钟或以上	
28	5.8	由仰卧转向俯卧	
29	6.2	稳定地独坐	
30	6.5	独坐时协调好	
31	6.6	*舀起小糖丸	
32	6.6	△完全的拇指相对拾起方木	
33	7	早期跨步运动	
34	7.5	牵拉站起	
35	7.6	*不完全的拇指相对抓糖丸	
36	7.6	走路之前的行进方式(俯卧、手膝、手足、其他)	
37	8.3	使两个勺子或方木在中线相碰	
38	8.5	跨步运动	

39	8.6	自己坐起
40	8.6	借助家具站起
41	8.9	精细地抓糖丸 (灵巧地钳夹)
42	9.6	拍手(中线技巧)
43	9.8	坐下
44	10	扶助下行走
45	11.1	独站
46	12	投球
47	12.1	独走
48	12.4	起立丨
49	13.2	扶助下右足独站
50	13.7	扶助下左足独站
51	14.1	侧身走
52	14.5	扶助上楼梯
53	14.7	倒退走
54	15.1	扶助下楼梯
55	17.6	试图站在行木上
56	18.7	左足独站
57	19.3	单足踏在行木上走
58	19.9	起立Ⅱ
59	20.1	右足独站
60	21.1	走直线: 大致方向
61	23.1	行木: 双足站立
62	24	踮脚走几步
63	24.3	独自上楼梯: 双足
64	24.4	双足跳离地面
65	25.3	独自下楼梯
66	25.6	行木: 企图跨步
67	25.6	倒行两米半
68	25.7	自第一级台阶下跳下
69	29.2	自第二级台阶下跳下
70	29.8	踮脚走两米半
71	29.9	跳远: 10 至 35cm (记录距离)
72	30+	起立:川
73	30+	上楼梯:双足交替向前
74	30+	行木: 交替步伐走部分路程
75	30+	保持双足走在直线上 (两米半)
76	30+	跳远: 35cm 至 60cm
77	30+	跳过: 5cm 高的绳子
78	30+	跳远: 60cm 至 85cm
79	30+	独脚跳两次以上
80	30+	下楼梯: 双足交替向前
81	30+	跳过 20cm 高的绳子



ge 43 of 45			ВМЈ Ор	en	/bmjopen-202:		
eTable 3 Distributions across whole pregnance			evel in 90 days pric		th trimester (\$1, T2, 175 on 2 July En	and T3) and comb	ined
				Estimated ex	kposure (မုန္တိ/ရွိက်ပွဲ		
	N	Minimum	25 <sup>th</sup> percentile	Mean ± SD	Mediang + JQR	75 <sup>th</sup> percentile	Maximum
Estimated exposure to PM <sub>2.5</sub>					ont S to te		
90 days prior to conception	1,174	38.17	44.00	$52.91 \pm 10.99$	48.43 5 8.07	62.06	80.53
First trimester	1,174	37.26	43.77	$52.07 \pm 10.98$	47.2 <b>% 5</b> 5 .31	61.08	82.41
Second trimester	1,174	38.46	47.57	$58.64 \pm 12.21$	版分式 57.9選 <del>即算</del> .62	67.19	90.02
Third trimester	1,174	37.03	47.25	$61.83 \pm 16.04$	58.8 <b>2</b> ± <b>3</b> 8.7	75.95	96.48
Total pregnancy	1,174	46.69	54.85	$57.48 \pm 3.97$	57.3 ± ± 5.76	60.61	66.98
Estimated exposure to NO <sub>2</sub>				V/0	ainin		
90 days prior to conception	1,174	25.86	45.49	$49.59 \pm 6.34$	بق ع. 49.944 ± <b>8</b> .27	53.76	70.48
First trimester	1,174	20.81	44.60	$48.8 \pm 6.43$	48.9 <b>2</b> ± <b>8</b> .51	53.10	69.31
Second trimester	1,174	28.93	47.18	$50.98 \pm 6.23$	51.2 ± \(\frac{1}{2}\).72	54.90	70.42
Third trimester	1,174	20.57	47.20	$51.79 \pm 6.78$	52.4 <b>§</b> ± <b>£</b> 47	56.67	75.12
Total pregnancy	1,174	27.50	47.89	$50.52 \pm 5.08$	50.4 <b>6</b> ± <b>8</b> 51	53.40	67.53

eTable 4 Pearson's correlations of PM<sub>2.5</sub> and NO<sub>2</sub> between each of the five different pregnancy time periods (N = 1,174)

	Estimated exposure to		PM <sub>2.5</sub>			or NO2					
		90 days prior	First	Second	Third	Total	90 days print	First	Second	Third	Total
		to conception	trimester	trimester	trimester	pregnancy	to conception 202	trimester	trimester	trimester	pregnancy
PM <sub>2.5</sub>	90 days prior to conception	1					4. Donemo				
	First trimester	-0.065	1				ent S to te				
	Second trimester	-0.779	-0.2012	1			baded uperi xt and				
	Third trimester	0.288	-0.7613	-0.1688	1		l from eur ( <i>t</i> d data				
	Total pregnancy	-0.534	-0.2709	0.6838	0.3858	1	A min				
NO <sub>2</sub>	90 days prior to conception	0.6383	0.0684	-0.4588	0.3714	0.0376	1 <b>ing</b> :				
	First trimester	0.1537	0.6352	-0.0159	-0.4927	-0.0633	0.5545	1			
	Second trimester	-0.431	0.0714	0.6269	-0.0133	0.7251	0.3345	0.5399	1		
	Third trimester	0.3027	-0.5213	0.0528	0.6817	0.4432	0.7149	0.2159	0.5145	1	
	Total pregnancy	0.0057	0.0781	0.2862	0.0737	0.4779	0.678 <b>cm</b> ii	0.7435	0.8755	0.7331	1

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eTable 5 Associations between PM<sub>2.5</sub> and NO<sub>2</sub> exposure in different pregnancy periods and adverse birgh outcomes (unadjusted models)

<u> </u>					g or				
6 7		Mean d	ifference		Odd ratios				
, 8 г	Per IQR increase in	Birth weight, grams (95%	Birth length, cm	PTB (case: 33)	LB <b>W</b> 55 (50)	LGA (case: 108)	SGA (case: 84)		
9	rei iQK increase in	CI)	(95% CI)	(95% CI)	( <b>9 5 2</b> CI)	(95% CI)	(95% CI)		
10		(N=941)	(N=927)	(N=945)	<b>4</b> 5)	(N=945)	(N=945)		
Estimated	90 days prior to conception	9.28 (-31.26, 49.83)	-0.09 (-0.27, 0.09)	0.98 (0.56, 1.74)	1.35 <b>4</b> ( <b>2</b> ) <b>2</b> , 2.36)	1.2 (0.87, 1.64)	0.98 (0.68, 1.41)		
13exposure	First trimester	21.95 (-16.90, 60.80)	0.14 (-0.03, 0.31)	0.98 (0.57, 1.70)	1 ( <b>ਉ.<u></u> 275</b> 1.77)	0.97 (0.71, 1.33)	0.78 (0.54, 1.13)		
14 15 <sup>to</sup> PM <sub>2.5</sub>	Second trimester	-18.21 (-57.78, 21.37)	0.04 (-0.13, 0.21)	0.85 (0.48, 1.50)	$0.61$ $\frac{1}{8}$ $\frac{1}{8}$ $\frac{1}{8}$ , $1.15$	0.92 (0.67, 1.27)	1.33 (0.94, 1.89)		
	Third trimester	-37.38 (-81.43, 6.68)	-0.32 (-0.51, -0.13)	1.35 (0.74, 2.47)	1.51 (2) (2) (2.85)	1.08 (0.76, 1.54)	1.12 (0.76, 1.66)		
16 17	Total pregnancy	-20.02 (-55.69, 15.65)	-0.1 (-0.26, 0.05)	0.81 (0.49, 1.33)	0.69 <b>a (<u>5</u>∕∄</b> , 1.16)	1 (0.75, 1.34)	1.2 (0.87, 1.66)		
1 <b>E</b> stimated	90 days prior to conception	-13.23 (-45.50, 19.03)	-0.12 (-0.26, 0.02)	1.2 (0.76, 1.89)	1.62 (75)	1.21 (0.93, 1.57)	1.24 (0.92, 1.66)		
19 exposure	First trimester	0.3 (-32.36, 32.96)	0.08 (-0.06, 0.22)	1.01 (0.64, 1.60)	1.15 (0.7), 1.86)	1.17 (0.90, 1.52)	1.27 (0.94, 1.71)		
20 to NO <sub>2</sub>	Second trimester	-22.85 (-53.39, 7.70)	-0.04 (-0.17, 0.09)	1.11 (0.72, 1.70)	1.08 <b>2</b> 0. <b>6</b> , 1.69)	1.06 (0.83, 1.36)	1.46 (1.10, 1.93)		
22	Third trimester	-32.72 (-67.16, 1.72)	-0.16 (-0.32, -0.01)	1.13 (0.69, 1.85)	1.35 $(0.8)$ , $2.28$ )	1.24 (0.94, 1.65)	1.58 (1.14, 2.18)		
23	Total pregnancy	-16.58 (-43.35, 10.20)	-0.03 (-0.15, 0.09)	1.03 (0.71, 1.50)	1.13 (0.75, 1.69)	1.16 (0.93, 1.44)	1.44 (1.13, 1.85)		
All signific	cant findings in the table are be	old.			an Co				
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eTable 6 Associations between PM2.5 and NO2 exposure in different pregnancy periods and adverse birth outcomes (co-exposure models)

<del>3</del> 6		Mean d	Odd ratios					
7	Per IQR increase in	Birth weight, grams	Birth length, cm	PTB (case: 33)	LBW (유물: 30)	LGA (case: 108)	SGA (case: 84)	
8	i ei iQik ilicicase ili	(95% CI)	(95% CI)	(95% CI)	(9 <b>§</b> 💆 (ČI)	(95% CI)	(95% CI)	
9 10		(N=941)	(N=927)	(N=945)	(1 <mark>% (9</mark> ) (45)	(N=945)	(N=945)	
1Estimated	90 days prior to conception	75.00 (-9.86, 159.86)	0.23 (-0.14, 0.60)	0.98 (0.56, 1.89)	0.41 (2)	1.14 (0.55, 2.40)	1.18 (0.48, 2.92)	
12 exposure	First trimester	19.59 (-71.23, 110.41)	0.00 (-0.40, 0.39)	0.97 (0.26, 3.65)	0.66  (2.1)  3.05	0.67 (0.32, 1.42)	0.73 (0.28, 1.93)	
13 <sup>1</sup> 14to PM <sub>2.5</sub>	Second trimester	-25.62 (-104.32, 53.09)	0.08 (-0.26, 0.42)	1.34 (0.37, 4.86)	0.94 ( 4.21)	0.83 (0.42, 1.62)	0.69 (0.34, 1.40)	
15 15 1VI <sub>2.5</sub>	Third trimester	13.77 (-72.33, 99.86)	-0.2 (-0.58, 0.17)	1.12 (0.31, 4.07)	0.94 ( <b>a</b> ) <b>3</b> ( <b>a</b> ) 3.35)	1.00 (0.48, 2.12)	0.57 (0.26, 1.23)	
16	Total pregnancy	21.13 (-36.41, 78.67)	0.02 (-0.23, 0.27)	0.73 (0.33, 1.61)	0.52 🙀 🔁 1.15)	0.98 (0.60, 1.61)	0.55 (0.32, 0.96)	
1Estimated	90 days prior to conception	-18.63 (-64.02, 26.76)	-0.09 (-0.29, 0.10)	1.24 (0.61, 2.49)	1.3 (6.64)	1.27 (0.84, 1.90)	1.39 (0.90, 2.15)	
19exposure	First trimester	-14.53 (-61.15, 32.09)	0.05 (-0.16, 0.25)	0.91 (0.45, 1.83)	1.14 🕳 🥰 2.37)	1.33 (0.89, 2.00)	1.70 (1.07, 2.69)	
•	Second trimester	-14.46 (-57.45, 28.54)	-0.08 (-0.26, 0.11)	1.22 (0.63, 2.36)	1.36 (2.71)	1.27 (0.87, 1.87)	1.50 (1.00, 2.24)	
20 21 to NO <sub>2</sub>	Third trimester	-13.13 (-64.87, 38.62)	0.04 (-0.18, 0.27)	0.77 (0.35, 1.67)	0.97 ( 4 2.16)	1.41 (0.90, 2.23)	1.77 (1.08, 2.91)	
22 23	Total pregnancy	-15.02 (-49.33, 19.30)	0.00 (-0.15, 0.15)	1.08 (0.64, 1.80)	1.28 🙇 .7 🗲 2.18)	1.21 (0.90, 1.63)	1.60 (1.15, 2.23)	

2\[All significant findings in the table are bold.\[2\]
2\[All significant findings in for the other air pollutant similar technologies.