Combined Effects of Prenatal Polycyclic Aromatic Hydrocarbons and Material Hardship on Child IQ

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Running/Short Title: Prenatal Air Pollution, Hardship and Child IQ

Conflicts of Interest: None
Abstract:

Importance: Polycyclic aromatic hydrocarbons are common carcinogenic and neurotoxic urban air pollutants. Toxic exposures, including air pollution, are disproportionately high in communities of color and frequently co-occur with chronic economic deprivation.

Objectives: We examined whether the association between child IQ and prenatal exposure to polycyclic aromatic hydrocarbons differed between groups of children whose mothers reported high vs. low material hardship during their pregnancy and through child age 5. We tested statistical interactions between hardships and polycyclic aromatic hydrocarbons, as measured by DNA adducts in cord blood, to determine whether material hardship exacerbated the association between adducts and IQ scores.

Design: Prospective cohort. Participants were recruited from 1998 to 2006 and followed from gestation through age 7 years.

Setting: Urban community (New York City)

Participants: A community-based sample of 276 minority urban youth

Exposure measure: Polycyclic aromatic hydrocarbon-DNA adducts in cord blood as an individual biomarker of prenatal polycyclic aromatic hydrocarbon exposure. Maternal material hardship self-reported prenatally and at multiple timepoints through early childhood.

Main outcome measure: Child IQ at 7 years assessed using the Wechsler Intelligence Scale for Children.

Results: Significant inverse effects of high cord PAH-DNA adducts on full scale IQ, perceptual reasoning and working memory scores were observed in the groups whose mothers reported a high level of material hardship during pregnancy or recurring high hardship into the child’s early years, and not in those without reported high hardship. Significant interactions were observed between high cord adducts and prenatal hardship on working memory scores (β=-8.07, 95% CI (-14.48, -1.66)) and between high cord adducts and recurrent material hardship (β=-9.82, 95% CI (-16.22, -3.42)).
**Conclusion**: The findings add to other evidence that socioeconomic disadvantage can increase the adverse effects of toxic physical "stressors" like air pollutants. Observed associations between high cord adducts and reduced IQ were significant only among the group of children whose mothers reported high material hardship. These results indicate the need for a multifaceted approach to prevention.

**Keywords**: air pollution, child IQ, prenatal exposure, PAH, adducts
1: Introduction

Exposure to polycyclic aromatic hydrocarbons (PAH) is prevalent in urban populations as a result of incomplete combustion of fossil fuels and other organic material. Specific sources of PAH include combustion of diesel, gasoline, coal, residential heating oil, tobacco smoke, and chargrilled or broiled foods. There is growing evidence that exposures to ambient and indoor air pollutants have adverse effects on neurodevelopment and that such toxic exposures are disproportionately high in lower income communities of color. These minority populations are also more likely to experience material hardship, an indicator of chronic economic stress, to live in lower quality housing, and to have inadequate educational and nutritional resources compared to higher income communities. Such socioeconomic stressors potentially compound or increase the effect of toxic environmental exposures.

In addition to exerting epigenetic effects, PAH bind covalently to DNA to form adducts, a widely used biomarker that reflects inter-individual variation in exposure, absorption, metabolic activation, and DNA repair, thereby providing an individual biologic dosimeter of an individual’s exposure to PAH. PAH-DNA adducts have previously associated with cancer in adults and with adverse neurodevelopmental outcomes in children.

The fetus is considered particularly susceptible to the effects of PAH exposure due to slower clearance of chemicals, underdeveloped detoxification and repair mechanisms, high rate of metabolic activity, and rapid growth during fetal development. PAH are readily transferred across the placenta and the fetal blood brain barrier. Here we tested the hypothesis that the adverse effect of prenatal exposure to PAH, measured by PAH-DNA adducts in cord blood, would be greater among children whose mothers experienced material hardship during pregnancy and in the children’s early years compared to children whose mothers did not experience material hardship. Material hardship is a measure used to assess an individual’s unmet basic needs in the areas of food, housing, and clothing. As has been shown with lead, even in the absence of significant main effects, combined
exposures to environmental toxicants and social stress can have significant impacts on neurodevelopment and therefore are of concern\textsuperscript{23-25}. We focused on prenatal PAH exposure because of the extensive structural and cellular-level changes that occur during the prenatal period and prior studies that have suggested that prenatal exposure to PAH adversely affects cognitive development\textsuperscript{26,27}. We evaluated both prenatal material hardship and that experienced continuously from pregnancy through childhood because prior studies have reported adverse effects of economic disadvantage and stress experienced during both developmental periods\textsuperscript{28,29}.

2: Methods

2.1: The Columbia Center for Children’s Environmental Health (CCCEH) cohort study:

A complete description of the CCCEH cohort and study design appears elsewhere\textsuperscript{30}. Briefly, African-American and Dominican women who resided in Washington Heights, Harlem, or the South Bronx in New York City (NYC) were recruited into a prospective cohort study between 1998 and 2006 through the local prenatal care clinics. To reduce the potential for confounding, enrollment was restricted to women who were non-active cigarette smokers in the age range of 18-35 years; non-users of other tobacco products or illicit drugs; free of diabetes, hypertension, or known HIV; and had initiated prenatal care by the 20\textsuperscript{th} week of pregnancy. The Institutional Review Board of the New York Presbyterian Medical Center approved the study. The mothers provided informed consent for themselves and their children and children provided assent at age 7.

2.2: Personal interviews, home caretaking environment, and maternal intelligence:

2.2.1: Prenatal interview:

A 45-minute questionnaire administered by a trained bilingual interviewer during the last trimester of pregnancy elicited demographic information, residential history, health and
environmental data such as active smoking (to confirm non-active smoking status as reported on the screening questionnaire) and exposure to environmental tobacco smoke (ETS). In the cohort, self-reported ETS exposure was positively correlated with cotinine measured in cord blood ($r=0.44$, $p$-value$<0.0001$). The questionnaire also elicited information on dietary PAH (consumption of broiled, fried, grilled or smoked meat) and information related to income and education.

2.2.2: Postnatal interviews and assessments:

Postnatal interviews were administered in-person at 6 months and annually thereafter to determine changes in residence, ETS exposure, and health and environmental conditions. The PERI-D was also re-administered at those interviews. At child age 3, the quality of the proximal caretaking environment was assessed using Caldwell and Bradley’s Home Observation for Measurement of the Environment (HOME)\(^{32}\). Maternal nonverbal intelligence was measured at child age 3 by the Test of Non-Verbal Intelligence-Third Edition (TONI-3)\(^{33}\), a 15-minute, language-free measure of general intelligence, relatively stable and free of cultural bias.

2.3: Material hardship:

A measure of material hardship\(^{21}\), assessing the level of unmet basic needs in the areas of food, housing, and clothing, was obtained prenatally and at child age 6 months, and 1, 2, 3, and 5 years by asking: “In the past year has there been a time when you: 1. couldn’t afford to buy food?; 2. couldn’t afford a place to stay?; 3. couldn’t afford gas/electricity?; or 4. couldn’t afford clothing?” Each answer was dichotomized (yes/no). High prenatal hardship was defined by a positive answer to at least one of the four questions at the prenatal assessment, and recurrent hardship by a positive response to at least one of the questions at $\geq50\%$ of the prenatal period and postnatal visits.
2.4 Biomarker measurement and prenatal monitoring: At the time of delivery, umbilical cord blood and maternal blood were collected and transported within several hours of collection to the CCCEH Molecular Epidemiology Laboratory. The buffy coat, packed red blood cells, and plasma were separated and stored at −70°C. DNA adducts of the representative PAH, benzo[a]pyrene (B[a]P), were analyzed in extracted white blood cell DNA using a high performance chromatography (HPLC)/fluorescence method which detects B[a]P tetraols. The adducts were dichotomized into detectable/nondetectable (high/low) because 57% of cord blood DNA samples had levels below the limit of detection (0.25 adducts per $10^8$ adducts). Some children lacked data on cord DNA adducts due to inadequate quantity or quality of DNA (n=145).

PAH metabolites were measured in spot urine collected at child age 5 at the Centers for Disease Control and Prevention (CDC) using automated liquid-liquid extraction and gas chromatography/isotope dilution high-resolution mass spectrometry, as previously described. Although a short-term biomarker (half-life of 6-35 hours), in conditions of chronic exposure the metabolites are considered a useful measure of exposure to PAH from all exposure sources and pathways. To adjust for urinary dilution of the samples, specific gravity (SG) measurements were obtained using a handheld refractometer (PAL-10-S-P14643C0; TAGO, Bellevue, WA). Metabolite levels were adjusted for SG using the formula: freshweight metabolites for the subject*(mean SG-1)/(SG for that subject-1).

2.5: Outcomes:

At child age 7 years, trained research workers administered the Wechsler Intelligence Scale for Children-Fourth Edition (WISC-IV), which consists of 15 subtests, 10 of which are core subtests used for the present study. Raw scores were converted into scaled scores based on a metric with a mean of 10 and standard deviation of 3 specific for the child’s age group as described in the WISC-IV manual. The scaled scores were used to derive four composite scores
(working memory, perceptual reasoning, processing speed, and verbal comprehension). These four indices were summed to derive the composite score for full scale IQ. The average expected performance on the full scale IQ score on the WISC-IV is 100, with a standard deviation of 15.40.

2.6: Statistical analysis:

A total of 394 children had available data on cord adducts. Of these, 276 also had available data on material hardship and WISC outcomes. 27 subjects were missing data on one or more covariates identified as predictors of IQ in the present cohort (at p<0.1) or as reported in the literature. These covariates included maternal self-report of ETS exposure during pregnancy, maternal education (<high school; ≥high school), ethnicity, maternal intelligence (treated as a continuous variable), child sex, and quality of the early home caretaking environment assessed at child age 3 (treated as a continuous variable). We conducted multiple imputation for missing data on those covariates.

Adducts were treated as high/low (detectable/nondetectable). Stratified analyses assessed the effect of high cord adducts in the different strata of material hardship (high vs. low prenatal and recurrent vs. non-recurrent hardship, respectively). To determine if the effects of high cord adducts in different strata of material hardship are statistically different or not, we tested statistical interactions, a product term between adducts and material hardship (prenatal or recurrent), in a linear regression model, with the terms for adducts, material hardship and key covariates regressed on each WISC outcome. The interaction term betas and 95% confidence intervals (CI) were then examined to determine whether the associations between PAH and IQ within the low and high hardship strata were statistically different.

In separate models with a smaller number of subjects (n=230), we adjusted for postnatal PAH exposure, using the SG-adjusted level of PAH metabolites in child urine at age 5. In further analyses (n=226), we adjusted for levels of the pesticide chlorpyrifos (CPF) and lead (n=156) in cord blood because these toxicants have been associated with lower working
memory scores in our cohort. Our analyses involved multiple comparisons; however to reduce the possibility of making a type II error, we did not perform Bonferroni adjustment.

As in previous analyses, to account for potential bias due to selection and loss to follow up, in post-hoc sensitivity analyses, we applied the inverse probability weighting (IPW) technique. As before, to model probability of staying in the study for each subject, we used a logistic model that included baseline variables for race/ethnicity, receipt of public assistance during pregnancy, high school education, college education, reported satisfaction with living conditions, cord adduct level, neighborhood poverty rate, Spanish language linguistic isolation, and indicator variables for missing data on these variables. Note that the missing data on covariates in the 27 subjects were first filled in with single imputation.

3: Results

Socio-demographic and exposure characteristics of the sample included in the analysis are presented in Table 1. Within our fully enrolled cohort, there were 394 women with available cord adduct data. A consort diagram showing how we arrived at our final analyzed sample is shown in Figure 1. There were no significant differences between the children included in the analysis and those not included due to missing WISC outcome or cord adduct data (n=118), except that there was a higher percentage of African Americans in the included group (38% vs. 26%) (Supplemental Table S1). In the sample analyzed, the correlation between prenatal material hardship and cord adducts was not significant (r=-0.03, p=0.67) and high cord adducts was not correlated with PAH metabolites at age 5 years (r=0.04, p=0.52) (Supplemental Table S2).

As we had previously hypothesized, stratified analyses showed that the association between PAH-DNA adducts in cord blood and IQ measures were significant only among the children whose mothers reported high prenatal or recurrent material hardship (Table 2). Among the group with high prenatal hardship, children who had high levels of adducts in cord blood had
a 5.81 point lower full scale IQ score, a 5.44 point lower perceptual reasoning score and a 6.67 point lower working memory score compared to children whose cord adducts were low (Table 2). The same significant relationships between adducts and IQ were not observed in the low material hardship group.

Similarly, after stratifying on recurrent hardship, adducts were significantly associated with full scale IQ, perceptual reasoning and working memory only within the high hardship group (Table 2, Figure 2). Among the group with recurrent hardship, children who had high levels of adducts in cord blood had a 6.63 point lower full scale IQ score, a 5.66 point lower perceptual reasoning score and a 8.06 point lower working memory score compared to children with low cord adducts. Statistically significant interactions between both prenatal (β=-8.07, 95% CI (-14.48, -1.66)) and recurrent (β=-9.82, 95% CI (-16.22,-3.42)) material hardship and cord adducts were observed on working memory scores (Table 2). Regarding the overall main effects of high cord adducts on cognitive outcomes, associations were uniformly inverse as hypothesized and significant for full scale IQ (β=-3.45, 95% CI (-6.35, -0.55)) and processing speed (β=-3.72, 95% CI (-7.28, -0.17)) and borderline for perceptual reasoning (β=-3.02, 95% CI (-6.30, 0.26)) (data not shown). We also conducted the stratified and interaction analyses in the subset with postnatal PAH exposure data available (n=230) (Supplemental Table S3). There was no difference in results before and after adjusting for postnatal PAH exposure, and therefore we did not include this variable in our final model.

In the subset with available cord plasma CPF data (n=226), CPF was significantly associated with working memory scores, after adjusting for cord adducts and stratified on prenatal and recurrent material hardship (data not shown). When the interaction model for cord adducts and recurrent hardship was adjusted for CPF, the interaction on working memory remained significant (β= -10.93, 95% CI (-17.94, -3.93)) and the interaction on full scale IQ and verbal comprehension became significant (β= -6.48, 95% CI (-12.76, -0.20) and β=-6.19, 95%
CI (-10.83, -1.57), respectively. The same interactions on working memory and full scale IQ were seen when considering prenatal hardship (β= -13.01, 95% CI (-19.97, -6.04) and β= -7.23, 95% CI (-13.52, -0.94), respectively) (Supplemental Table S4). Cord adduct and CPF levels were not significantly correlated (r= -0.02, p=0.81); neither were cord adduct and lead levels (r= -0.06, p=0.48). Furthermore, after adjusting for prenatal lead exposure in a smaller subsample with available data (n=156), high cord adducts were significantly associated with working memory scores in both the high prenatal (β= -10.45, 95% CI (-16.87, -4.02) and recurrent material hardship (β= -9.46, 95% CI (-16.08, -2.83) groups. The corresponding interaction terms were also significant (Supplemental Table S5).

After repeating all analyses with IPW, the direction and magnitude of associations did not materially change, indicating that the results are not influenced by sample selection and loss to follow-up (Supplemental Table S6).

4: Discussion

This is the first report of an interaction between chronic socioeconomic stress and prenatal exposure to PAH, represented by PAH-DNA adducts in cord blood, on children’s IQ. Cord PAH-DNA adducts are a direct measure of the individual fetal dose of PAH integrating exposure over the past 3-4 months47.

The findings are of concern because, as has been shown with lead, even a modest decrease in IQ can impact lifetime earnings48,49. They are also consistent with studies showing modification of the neurotoxic effect of lead by social class22.

PAH are common urban pollutants and include known carcinogens and neurotoxicants such as B[a]P. B[a]P, considered a representative PAH, is highly correlated with the other 7 genotoxic PAH measured in prenatal air (r = 0.80–0.96, p = 0.001 except for dibenz[a,h]anthracene, r = 0.53, p < 0.001)14. Although PAH is ubiquitous in the urban environment, low-income communities are disproportionately exposed due to greater siting of
heavily trafficked roadways, bus and truck depots, power plants and industrial boilers, and the higher prevalence of smokers in low-income households\textsuperscript{8,50}.

This report adds to the growing literature on the vulnerability of the developing fetus and young child to the toxic effects of environmental pollutants\textsuperscript{17,18} as well as to socioeconomic disadvantage \textsuperscript{51-53} (for review see \textsuperscript{54}). Additional studies have observed similar decreases in IQ score as measured by the WISC-IV in response to environmental toxicants experienced prenatally and during early childhood as our present study\textsuperscript{55-58}. Economic deprivation and related stress early in life have been linked to behavioral problems and lower IQ scores in children\textsuperscript{59}. Cumulative poverty and hardship in the first year of life were associated with negative effects on cognitive function in childhood\textsuperscript{58}.

The observed interaction between PAH (cord adducts) and material hardship is consistent with other reports that adverse social conditions can modify the neurotoxicity of environmental pollutants such as lead, traffic-related pollutants, and ETS\textsuperscript{22,55,60-62} (see\textsuperscript{63} for review).

A number of prior reports in other populations have indicated adverse neurodevelopmental effects of air pollution\textsuperscript{26,27,64,65}. In the present NYC cohort, prenatal exposure to PAH, as measured by 48-hour prenatal air monitoring, was associated with delayed mental development at age 3 years\textsuperscript{30} and was associated with lower intelligence at age 5 years in both the NYC cohort and in a parallel Polish cohort\textsuperscript{26,27}. In those reports, the association with PAH-DNA adducts was not examined.

Mechanisms underlying interactions between toxic pollutants and psychosocial factors are not well understood. However, chronic psychosocial stress is known to increase allostatic load that can impair individual resilience and ability to recover from toxic insults by interfering with normal functioning of protective toxicokinetic and toxicodynamic processes, resulting in elevated inflammatory tone\textsuperscript{63,66,67}. Production of inflammatory mediators is also stimulated by physical toxicants\textsuperscript{68,69}. Thus the two types of exposures could potentiate each other through
common physiological pathways such as inflammation\textsuperscript{63,68,70,71}. Research on the combined effect of maternal stress during pregnancy and prenatal air pollution in mice showed that these stressors act synergistically to induce neuroinflammation, leading to future neurobehavioral disorders\textsuperscript{29}.

With respect to the harmful effects of prenatal PAH exposure, a number of additional pathways have been suggested including endocrine disruption\textsuperscript{72-74}, binding to receptors for placental growth factors resulting in decreased exchange of oxygen and nutrients\textsuperscript{75}, binding to the human Ah receptor to induce P450 enzymes\textsuperscript{76}, DNA damage resulting in activation of apoptotic pathways\textsuperscript{77-79}, oxidative stress due to inhibition of the brain antioxidant scavenging system\textsuperscript{80}, epigenetic alterations affecting gene expression\textsuperscript{81,82} and/or altered expression of nuclear transcription factors that mediate the onset of neuronal cell differentiation\textsuperscript{19}.

Our findings are consistent with those from other animal and human studies. Although the exposure in experimental laboratory studies is considerably higher than those in the NYC cohort, impaired memory has been observed in animals exposed gestationally to PAH at doses below those causing overt toxicological effects\textsuperscript{83,84}. Differences in the medial temporal/memory composite have been significantly related to children’s socioeconomic status\textsuperscript{85}.

Regarding the interpretation of our results, it is possible that PAH are equally toxic under conditions of low and high hardship, but that low hardship families have unmeasured resources positively affecting the health and development of children and buffering the adverse impact of PAH exposure\textsuperscript{86}. Furthermore, those individuals experience material hardship may also have had a poor prenatal diet\textsuperscript{87,88} and exposure to other chemical toxicants, such as lead\textsuperscript{57,89}, ETS\textsuperscript{55} and other air pollutants\textsuperscript{90-92}, that also may account for the lower IQ seen in the children.

In addition, it is possible that the WISC-IV scores at age 7 are capturing manifestations other than child IQ. For example, Oliveras-Rentas et al., found that among a sample of high-functioning children with autism-spectrum disorder, scores on the WISC-IV were correlated with adaptive behavior as measured on the Vineland instrument and autism spectrum symptomology.
scores as measured on the Autism Diagnostic Observation Schedule (ADOS). This study did not see a significant correlation between WISC-IV scores and ADHD symptomology\textsuperscript{93}. However, an additional study investigated clusters of IQ profiles as defined by the WISC-IV and found an association between processing speed and inattention among children diagnosed with ADHD. They concluded that WISC-IV scores may be helpful in predicting symptomology of children with ADHD\textsuperscript{94}. In particular, working memory scores on the WISC may be affected by clinical behavior problems\textsuperscript{40}.

We acknowledge a number of additional limitations. Although we have adjusted for the possible confounding effects of ETS, there is always the possibility that some residual confounding remains; and we did not have data on a measure of psychological stress. In addition, we excluded active smokers, illicit drug users, and women with preexisting disease, thereby limiting generalizability.

The strengths of the study include the longitudinal design and ability to account for a number of factors other than PAH exposure that are known to affect child neurobehavioral development via available biomarker and questionnaire data.

There is growing recognition of the continuing need to document interactions between adverse social conditions/psychosocial stressors and environmental toxicants and to understand the mechanisms involved for effective intervention\textsuperscript{63,95}. The present results suggest the need for a multifaceted approach to reduce PAH exposure and alleviate material hardship in order to protect the developing fetus and young child. The approach could combine screening women early in pregnancy to identify those needing material support and policy interventions to reduce air pollution exposure in urban areas, especially low-income communities.

5. Conclusion

This study provides evidence that material hardship influences the effect of prenatal exposure to environmental PAH, measured by PAH-DNA adducts in cord blood, on child IQ.
The associations between PAH on full scale IQ and working memory were seen mainly among the group of children whose mothers experienced material hardship during pregnancy and in the children’s early years.

PAH are widespread in urban environments worldwide, largely as a result of fossil fuel combustion. Their concentrations can be reduced using currently available pollution controls, greater energy efficiency, alternative energy sources, and regulatory intervention to remove highly polluting sources. These results add to prior data linking PAH to cognitive and behavioral problems in children and suggest the need for a multifaceted approach to prevention.
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References


93. Oliveras-Rentas RE, Kenworthy L, Roberson RB, 3rd, Martin A, Wallace GL. WISC-IV profile in high-functioning autism spectrum disorders: impaired processing speed is associated with increased autism communication symptoms and decreased adaptive


### TABLES

Table 1. Socio-demographic and exposure characteristics of subjects (n=276)$^a$

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean</th>
<th>SD</th>
<th>range</th>
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<tr>
<td>Cord blood PAH-DNA adducts (% detectable)</td>
<td>43.8</td>
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<td></td>
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<tr>
<td>PAH Metabolites at Age 5 [% &gt; median]$^b$</td>
<td>47.0</td>
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<td>Sex (% female)</td>
<td>53.6</td>
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<td>Maternal ETS (% reporting smoker in the home)</td>
<td>33.3</td>
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<td>WISC Full Scale</td>
<td>98.6</td>
<td>13.2</td>
<td>(48, 133)</td>
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<td>WISC Verbal Comprehension</td>
<td>96.3</td>
<td>12.2</td>
<td>(45, 134)</td>
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<tr>
<td>WISC Processing Speed</td>
<td>101.1</td>
<td>15.2</td>
<td>(62, 138)</td>
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<tr>
<td>WISC Perceptual Reasoning</td>
<td>100.8</td>
<td>14.3</td>
<td>(63, 137)</td>
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<td>WISC Working Memory</td>
<td>97.4</td>
<td>14.2</td>
<td>(54, 135)</td>
</tr>
<tr>
<td>Maternal education (% &gt; high school education)</td>
<td>63.4</td>
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<tr>
<td>Ethnicity (% AA)</td>
<td>38.0</td>
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<td>Maternal IQ (TONI)</td>
<td>20.7</td>
<td>8.7</td>
<td>(0, 43)</td>
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<tr>
<td>Prenatal hardship (% ≥1 hardships)</td>
<td>42.8</td>
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<td>Recurrent up to 5 years (% ≥ 1 hardship, 50% of the time)</td>
<td>39.9</td>
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<td>HOME inventory</td>
<td>39.8</td>
<td>6.0</td>
<td>(22, 52)</td>
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</table>

$^a$ Subjects have data on cord adducts and IQ test results at age 7.

$^b$ Variable dichotomized at the median level for the entire population (8223.41 ng/m$^3$)
Table 2. Association between cord blood PAH-DNA adducts on IQ at age 7, stratified by prenatal and recurrent material hardship (N=276)\(^a\)

<table>
<thead>
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<td>(\beta_{adducts}) (95% CI)</td>
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<td>(\beta_{interaction}) (95% CI)</td>
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<td>(\beta_{adducts}) (95% CI)</td>
<td>(\beta_{interaction}) (95% CI)</td>
</tr>
<tr>
<td>Full Scale</td>
<td>-1.79 (-5.50, 1.93)</td>
<td>-5.81 (-10.35, 1.26)*</td>
<td>-4.66 (-10.43, 1.11)</td>
<td>-1.32 (-4.97, 2.33)</td>
<td>-6.63 (-11.28, -1.98)*</td>
<td>-5.59 (-11.37, 0.20)</td>
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<td>Verbal Comprehension</td>
<td>-1.08 (-4.29, 2.14)</td>
<td>-3.36 (-7.61, 1.00)</td>
<td>-2.39 (-7.57, 2.80)</td>
<td>-0.79 (-3.98, 2.40)</td>
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<td>Processing Speed</td>
<td>-3.59 (-8.21, 1.02)</td>
<td>-4.17 (-9.75, 1.41)</td>
<td>-0.97 (-8.09, 6.16)</td>
<td>-3.46 (-7.71, 0.79)</td>
<td>-4.02 (-10.13, 2.09)</td>
<td>-0.83 (-7.97, 6.30)</td>
</tr>
<tr>
<td>Perceptual Reasoning</td>
<td>-1.45 (-5.86, 2.95)</td>
<td>-5.44 (-10.27, -0.61)*</td>
<td>-4.66 (-11.20, 1.89)</td>
<td>-1.29 (-5.55, 2.96)</td>
<td>-5.66 (-10.71, -0.61)*</td>
<td>-4.74 (-11.32, 1.85)</td>
</tr>
<tr>
<td>Working Memory</td>
<td>0.57 (-3.70, 4.85)</td>
<td>-6.67 (-11.38, -1.95)*</td>
<td>-8.07 (-14.48, -1.66)*</td>
<td>1.24 (-3.13, 5.60)</td>
<td>-8.06 (-12.49, -3.63)*</td>
<td>-9.82 (-16.22, -3.42)*</td>
</tr>
</tbody>
</table>

\(^*\)p-value<0.05

\(^a\) Adjusting for ETS, sex, maternal education, maternal intelligence, ethnicity, and the home caretaking environment.
Figure Legends:

Figure 1: Subject selection for present analysis

Figure 2: Full Scale IQ and Working Memory Scores in the low and high cord PAH-DNA adduct groups stratified by recurrent hardship (n=276)
PAH-DNA cord adduct data available (N=394)

Child WISC scores at age 7 years (N=287)

Prenatal and at least one postnatal hardship measure available:
Sample included in final model (N=276)

Missing child WISC scores at age 7 years (N=107)

Missing prenatal and/or at least one postnatal hardship measure
(N=11)

Figure 1
Figure 2

A box plot showing the distribution of working memory and full scale IQ scores across different levels of adducts (Low, High) and hardship (Low, High). The x-axis represents the combinations of adducts and hardship levels, while the y-axis represents the average estimated score.
Highlights:

- PAH-DNA adducts in cord blood provided an individual measure of prenatal exposure.
- Material hardship in pregnancy and child’s early life proxied economic deprivation.
- Adverse effects on child IQ at age 7 were seen only among mothers with hardship.
- Interaction between high adducts and hardship on working memory was significant.
- These results indicate the need for a multifaceted approach to prevention.