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RESEARCH ARTICLE

Risk of lead exposure, subcortical brain structure, and cognition in a large cohort of 9-to 10-year-old children

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Abstract

Background

Lead, a toxic metal, affects cognitive development at the lowest measurable concentrations found in children, but little is known about its direct impact on brain development. Recently, we reported widespread decreases in cortical surface area and volume with increased risks of lead exposure, primarily in children of low-income families.

Methods and findings

We examined associations of neighborhood-level risk of lead exposure with cognitive test performance and subcortical brain volumes. We also examined whether subcortical structure mediated associations between lead risk and cognitive performance. Our analyses employed a cross-sectional analysis of baseline data from the observational Adolescent Brain Cognitive Development (ABCD) Study. The multi-center ABCD Study used school-based enrollment to recruit a demographically diverse cohort of almost 11,900 9- and 10-year-old children from an initial 22 study sites. The analyzed sample included data from 8,524 typically developing child participants and their parents or caregivers. The primary outcomes and measures were subcortical brain structure, cognitive performance using the National Institutes of Health Toolbox, and geocoded risk of lead exposure.

Children who lived in neighborhoods with greater risks of environmental lead exposure exhibited smaller volumes of the mid-anterior (partial correlation coefficient $[r_p] = -0.040$), central $(r_p = -0.038)$, and mid-posterior corpus callosum $(r_p = -0.035)$. Smaller volumes of these three callosal regions were associated with poorer performance on cognitive tests measuring language and processing speed. The association of lead exposure risk with cognitive performance was partially mediated through callosal volume, particularly the mid-posterior corpus callosum. In contrast, neighborhood-level indicators of disadvantage were not associated with smaller volumes of these brain structures.

Data Archive. The ABCD data used in this report came from https://doi.org/10.15154/1523347.

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Conclusions

Environmental factors related to the risk of lead exposure may be associated with certain aspects of cognitive functioning via diminished subcortical brain structure, including the anterior splenium (i.e., mid-posterior corpus callosum).

Introduction

Elevated blood-lead concentrations during childhood, which are more common among low-income children [1], are associated with intellectual deficits and behavioral problems [2–13]. While it has been estimated that 40% of the world's children have elevated blood-lead levels, 90% of these children live in low- and middle-income countries [14]. Lead-based paint and leaded gasoline reflect primary transnational sources of exposure, but there may be many more sources of lead exposure in developing countries, such as lead-glazed ceramics, lead mining and smelting, flour mills, lead-battery recycling plants, and lead-containing medicines and cosmetics [15,16]. The annual cost of childhood lead exposure in these low- and middle-income countries is nearly \$1 trillion [17].

Lead undoubtedly has effects on brain development [18], but most studies investigating its effects on brain structure and function involve heavily exposed children and adults [19–24]. We recently reported that greater *risk* of lead exposure [25,26]—based on age of housing and poverty levels in participants' residential neighborhoods [1,27]—was negatively associated with cognition and whole-brain cortical structure in 9- and 10-year-old children of low-income families [28]. Cortical vertex maps showed that reductions in total cortical surface area and volume were associated with increasing lead risk in children of low- but not high-income families.

Lead exposure is also associated with subcortical brain morphology. Occupational lead exposure is inversely associated with posterior corpus callosum and periventricular white matter volumes [19], and childhood lead exposure is associated with altered white-matter connectivity in young adults' corpus callosum and corona radiata [21]. Given different spatiotemporal trajectories of white- and gray-matter development [29,30], the low-socioeconomic-status (SES)-specific inverse associations that we observed between lead risk and cortical gray matter [28] may be differentially manifested within subcortical brain regions. If the cognitive effects of lead exposure are mediated by effects on brain structures that mature at different rates and times during development (e.g., in-utero, infancy, childhood, adolescence) [18], then identifying such regional biomarkers and their associations with neighborhood-level risk of exposure may increase our understanding of the temporally-dependent impact on developmental outcomes.

No studies have investigated associations between children's low-level lead exposure and subcortical brain structure. We employed neighborhood-level risk scores, which are validated proxies for census-tract-level prevalence of elevated blood-lead levels and mean blood-lead levels [28,31], to evaluate associations between brain structure and lead risk at high and low levels of exposure. We analyzed Adolescent Brain Cognitive Development SM Study (ABCD Study attaction data to determine relationships between lead-exposure risk, subcortical volumes, and performance on NIH Toolbox cognitive tests [32,33]. Our goal was to explore (1) associations between subcortical brain regions and lead-exposure risk, (2) whether family income moderated these associations, (3) subcortical volumes' associations with cognitive performance, and (4) whether the volumes of those subcortical regions mediated relationships between lead risk and cognition.

Methods

Participants

ABCD is a 10-year longitudinal study involving 21 U.S. study sites [34]. Using school-based enrollment [35], the consortium enrolled nearly 11,900 9- and 10-year-old children from an initial 22 sites. The ABCD Study's data collection sites all used the same protocol for recruitment, testing, and neuroimaging [36]. ABCD demographics correspond well with the American Community Survey (ACS) [37]. Our data came from the July 2019 ABCD 2.0.1 data release [38], which included baseline data for 11,875 children. Currently, there are no bloodlead data for ABCD participants.

Centralized IRB approval was obtained from the University of California, San Diego. Study sites obtained approval from their local IRBs. Parents provided written informed consent; children provided written assent. Data collection and analysis complied with all ethical regulations.

Lead risk and area deprivation index (ADI)

We used a high-resolution nationwide map (https://www.vox.com/a/lead-exposure-risk-map) to obtain geocoded lead-risk scores for participants' census tracts [25,26]. These scores reflect national deciles of a weighted sum of two validated correlates of lead exposure (https://github.com/voxmedia/data-projects/blob/master/vox-lead-exposure-risk/calculate-lead-risk.py): the ages of homes (weight = 0.58) and poverty (0.42) rates [1,27], both derived from ACS data. Lead-risk scores were previously shown to be highly associated with childhood lead exposure in children [28,31]. For instance, we previously showed that these lead-risk scores were positively associated with both the rate of elevated blood-lead levels across 13 states and 2 cities at the census-tract level and the geometric mean of census-tract-level blood-lead levels in the state of Maryland [28]. ABCD site-by-site lead-risk-score distributions have been published previously [28]. Briefly, for some sites, the lead-risk scores were uniformly distributed, while other sites showed modalities at either lower, intermediate, and/or higher lead-risk scores, thereby reflecting geographic differences in risk of exposure [39].

ADI is a composite metric of neighborhood deprivation (e.g., low education, poor plumbing) [40,41] and, like lead-exposure risk, incorporates poverty rates. Unlike lead-exposure risk, ADI does not include data related to age of housing. Census-tract-level ADI was computed based on coefficient values of past research [40] and discretized into national percentiles. The R code for computing and merging ADI (and its national percentile) with ABCD data is available: https://github.com/ABCD-STUDY/geocoding/blob/master/Gen_data_proc.R.

ABCD data

We analyzed uncorrected baseline performance on seven NIH Toolbox tests [32] and baseline structural brain measures (volumes of 21 subcortical regions) [42]. Data collection procedures are described in detail elsewhere [32,42,43].

NIH Toolbox tests show good convergent validity compared with gold standards of cognitive testing [44]: (1) the Picture Vocabulary Test (a measure of language; Version 2.0, Ages 3 +), (2) the Flanker Inhibitory Control and Attention Test (attention, executive function; Version 2.0, Ages 8–11), (3) the List Sorting Working Memory Test (working memory; Version 2.0, Ages 7+), (4) the Dimensional Change Card Sort Test (executive function; Version 2.0, Ages 8–11), (5) the Pattern Comparison Processing Speed Test (processing speed; Version 2.0, Ages 7+), (6) the Picture Sequence Memory Test (episodic memory; Version 2.0, Form A, Ages 8+), and (7) the Oral Reading Recognition Test (language; Version 2.0, Ages 3+). Briefly,

the NIH Toolbox was administered on an iPad (~25–30 min to complete), with all task's instructions being read by the examiner, except for the Pattern Comparison Processing Speed Test (i.e., presented by an audio recording). All tasks were administered with the iPad in an upright position (~45° degree angle), with some tasks incorporating a home-base "button" in front of the iPad for the participants on which the participants would place their index finger between the task's trials (Flanker Inhibitory Control and Attention Test, Dimensional Change Card Sort Test).

We obtained measures of subcortical brain structure using FreeSurfer v5.3.0 on acquired T₁w MRI volumes; ABCD neuroimaging data collection and processing procedures have been described [45]. Depending on the study site, Siemens, Philips, and GE scanners were used, with T₁ acquisition times (min:s) being 7:12, 5:38, and 6:09, respectively. Accordingly, a random effect of scanner serial number was included in analyses (as described below). All neuroimaging parameters are available: https://abcdstudy.org/images/Protocol_Imaging_Sequences.pdf. Structural magnetic resonance imaging (MRI) T₁ images were acquired first, followed by functional MRI and diffusion MRI images (data not shown). T₁w images were corrected for gradient nonlinearity distortions per scanner-manufacturer guidelines, with further details comprehensively described previously [45]. ABCD data are publicly available on the NIMH Data Archive (https://data-archive.nimh.nih.gov/abcd).

Statistical analyses

Analyses included 8,524 children with complete data for the variables of interest (Table 1 in S1 Appendix). Participants' data were excluded if the primary residential address was invalid (remaining n = 11,175) or unable to be geocoded into a 1–10 lead-risk score (remaining n = 11,169), a valid household/family income was not provided (remaining n = 10,234), there were missing data for sex, age, parental education, race, ethnicity, NIH toolbox test or composite scores, or structural imaging measures (remaining n = 9,519), if the ADI score was missing or invalid (weighted sum = 0) (remaining n = 9,331), or if the neuroimaging data did not pass all quality-control measures or there were neuroanatomical variants (incidental findings) judged to be of possible clinical significance (final n = 8,524) [46].

We employed linear mixed-effects models to determine lead-risk associations and family income × lead-risk interactions on 21 subcortical volumes. We averaged bilateral data across both hemispheres (Table 1 in S1 Appendix). In accordance with previous research analyzing associations between brain structure and socioeconomic and/or environmental conditions [28,47,48], we controlled for children's age, sex, race, ethnicity, maximum parental education, and family income. Analyses of subcortical data also controlled for intracranial volume (ICV), which was inversely associated with both lead risk, Spearman's rho (ρ) = -0.11, p < .001, and ADI, ρ = -0.11, p < .001. Random-effects structures included random intercepts for MRI serial number (i.e., some study sites have multiple machines) and family identification number (i.e., many ABCD participants were siblings). We used the Benjamini-Hochberg false-discovery-rate (FDR) algorithm to correct for multiple comparisons [49].

Lead risk, age (in months), and ICV were continuous factors. Maximum parental education was a continuous factor with seven levels ($1 = \le 6^{th}$ grade; $2 = 7^{th} - 9^{th}$ grade; $3 = 10^{th} - 12^{th}$ grade, no diploma; 4 = high-school graduate, GED or equivalent; 5 = Some college with no degree, Associate's degree; 6 = bachelor's degree; 7 = master's degree, professional degree, or doctorate). Children's race and ethnicity were categorical factors derived from parent reports on the child. Race had 6 = levels: "White", "Black", "Asian", "American Indian or Alaska Native", "Native Hawaiian or Other Pacific Islander", or "Other" (e.g., multiracial). Ethnicity had two levels: "Hispanic" or "Not Hispanic". As within ABCD's Data Exploration and Analysis Portal

(deap.nimhda.org), family income was a categorical factor with 3 levels, per parents' reported household income (Low Income: ≤\$50K; Middle Income: \$50K-\$100K; High Income: ≥\$100K). Categorical factors were effects-coded to facilitate interpretation of main effects given higher-level interactions [50]. Continuous factors were centered (i.e., a constant was subtracted) to make parameter estimates more interpretable [50].

To evaluate whether lead-risk associations could be explained by general neighborhood disadvantage, we conducted sensitivity analyses in which ADI replaced lead risk in the models. The ADI national percentile was converted into deciles and centered to match the lead-risk analyses. To further verify lead-risk associations with subcortical volume, we replaced lead risk with age of housing (i.e., the factor included in computing lead risk but not ADI). These data, which generally reflect census-tract-level housing-age-based estimates of the proportion of homes with lead-based paint hazards [27], were maintained on their original scale (range = 0.05–64.54), Box-Cox transformed to correct for positive skewness, and mean-centered for analysis.

Because our research has shown that cortical structure mediates associations between environmental factors (family income) and cognition [47], we examined whether subcortical structure mediated associations between lead risk and cognitive performance. Subcortical volume was mean-centered when estimating indirect associations. Analyses employed linear mixedeffects models, with children's sex, age, parental education, race, ethnicity, and family income as covariates. Here, the criterion was cognitive test score, so the random-effects structure included random intercepts for study site (rather than MRI serial number) and family identification number. Accordingly, because these analyses ultimately tested whether the association between lead risk and cognitive test score was mediated by subcortical volume, in which the criterion was cognitive test score and not subcortical volume, ICV was not included as a covariate in these analyses of cognitive performance. The presence of statistically significant indirect associations (i.e., products of unstandardized regression coefficients: lead-risk→brain-structure, brain-structure→cognition, controlling for lead risk) was evaluated via construction of bias-corrected percentile bootstrapping 95% confidence intervals (CI; 10,000 bootstrapped samples) [51], irrespective of the statistical significance of the corresponding total effect [52]. To minimize issues of multicollinearity of multiple mediators (i.e., collinear relationships between subcortical volumes of different regions) [53], only subcortical volumes associated with cognitive test performance were tested as mediators.

We conducted analyses using MATLAB's Statistics and Machine Learning Toolbox 11.7 (R2020a; MathWorks). Model output and model-fit characteristics are provided in Tables 2–116 in the S1 Appendix. Statistical reporting in the main text is in the form of t-tests except when involving categorical factors (e.g., family income), in which results are in the form of t tests computed using MATLAB's *anova* function (i.e., the combined statistical significance of all coefficients of the corresponding factor). Effect sizes of continuous factors are represented by partial correlation coefficients (t_p), which control for all model covariates and are calculated using the corresponding t-statistic and degrees of freedom [54]. The 95% CIs of the effect sizes were derived from the sample variance of the partial correlation [55].

Results

Sample characteristics

Children included in our analyses did not appreciably differ in key sociodemographic indicators compared with the entire ABCD cohort (Table 1). Lead-risk scores were bimodally distributed: 40.5% of participants lived in neighborhoods with low lead-risk scores (lead risk \leq 3; n = 3,450); 32.3%, intermediate lead-risk scores ($4 \leq \text{lead risk} \leq 7$; n = 2,749); and 27.3%, high lead-risk scores (lead risk \geq 8; n = 2,325) (Fig 1A in S1 Appendix). Participants also tended to

	Release 2.0.1 (%)	Sample with Complete Data Used in This Study (%)			
Sex					
Male	6,188 (52.1%)	4,469 (52.4%)			
Female	5,681 (47.8%)	4,055 (47.6%)			
Missing/Undefined	6 (0.1%)	0 (0%)			
Income Bracket					
<\$50K (Low)	3,222 (27.1%)	2,415 (28.3%)			
\$50-100K (Mid)	3,070 (25.9%)	2,482 (29.1%)			
>\$100K (High)	4,565 (38.4%)	3,627 (42.6%)			
Missing/Undefined	1,018 (8.6%)	0 (0%)			
Race					
American Indian/Alaska Native	62 (0.5%)	42 (0.5%)			
Asian	276 (2.3%)	188 (2.2%)			
Black	1,867 (15.7%)	1,168 (13.7%)			
Native Hawaiian/Pacific Islander	16 (0.1%)	10 (0.1%)			
Other	1,958 (16.5%)	1,364 (16.0%)			
White	7,523 (63.4%)	5,752 (67.5%)			
Missing/Undefined	173 (1.5%)	0 (0%)			
Ethnicity					
Hispanic	2,409 (20.3%)	1,640 (19.2%)			
Not Hispanic	9,308 (78.4%)	6,884 (80.8%)			
Missing/Undefined	158 (1.3%)	0 (0%)			
Total	11,875	8,524			

Table 1. Demographics for the Adolescent Brain Cognitive Development (ABCD) study.

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live in less-deprived neighborhoods (i.e., \leq 5th decile; Fig 1B in S1 Appendix). On average, based on housing age, ~20.5% of houses in participants' neighborhoods were estimated to contain lead-based paint hazards (Fig 1C in S1 Appendix) [26,27]. There were positive associations between ADI and both lead risk, ρ = 0.36, p < .001, and housing age (i.e., estimated percentage of homes in the census tract with lead-based paint hazards based on housing age), ρ = 0.13, p < .001.

Lead risk and subcortical brain structure

Of 21 subcortical regions (Fig 1) [56], lead-exposure risk was associated with smaller volumes (mm³) of the posterior (r_p = -0.022 [-0.033, -0.011]), mid-posterior (r_p = -0.035 [-0.046, -0.024]), central (r_p = -0.038 [-0.048, -0.027]), and mid-anterior corpus callosum (r_p = -0.040 [-0.051, -0.030]) (Figs 2 and 3). The mid-anterior, central, and mid-posterior corpus callosal associations passed FDR correction (q < .05) [49]. Anterior corpus callosum volume was not associated with lead risk (r_p = -0.004 [-0.015, 0.007]). No other main effects of lead risk were significant, $P_s \ge .074$ (uncorrected) (Tables 2–22 in S1 Appendix).

For posterior, mid-posterior, central, and mid-anterior corpus callosal volumes, there were no main effects of family income, $F(2, 8508)s \le 1.10$, $ps \ge .334$ (Tables 13–16 in S1 Appendix), Family Income × Lead Risk interactions (Fig 3), $F(2, 8508)s \le 0.98$, $ps \ge .376$, or Sex × Lead Risk interactions, $F(1, 8509)s \le 0.32$, $ps \ge .570$ (Tables 23–26 in S1 Appendix).

Accounting for neighborhood disadvantage

Correcting for FDR, ADI was inversely associated with subcortical gray-matter ($r_p = -0.037$ [-0.048, -0.026]), cerebellar cortical ($r_p = -0.032$ [-0.043, -0.021]), accumbens area ($r_p = -0.029$

Subcortical Region	Risk of Lead Exposure			Area Deprivation Index		
	<u>t(</u> 8508)	p	b	<u>t(</u> 8508)	p	b
Corpus Callosum (Mid-Anterior)	-3.73	< .001	-1.52	-0.80	.421	-0.48
Corpus Callosum (Central)	-3.47	.001	-1.25	-0.81	.417	-0.42
Corpus Callosum (Mid- Posterior)	-3.21	.001	-1.03	-0.80	.422	-0.37
Corpus Callosum (Posterior)	-2.03	.043	-1.06	-0.49	.624	-0.37
Putamen	-1.78	.074	-4.07	-2.43	.015	-8.06
Caudate	-1.60	.110	-2.99	-1.93	.053	-5.39
3 rd Ventricle	1.52	.129	1.34	-0.34	.733	-0.44
Accumbens Area	-1.30	.193	-0.43	-2.72	.007	-1.34
Lateral Ventricle	1.26	.209	13.37	-0.45	.652	-7.09
Subcortical Gray Volume	-1.23	.220	-14.77	-3.39	.001	-60.70
Hippocampus	1.18	.240	1.54	-1.03	.303	-1.91
Inferior Lateral Ventricle	1.03	.304	0.57	-0.18	.860	-0.15
Ventral Diencephalon	0.93	.355	1.03	-1.54	.123	-2.57
Pallidum	-0.91	.365	-0.63	-2.20	.028	-2.27
Thalamus Proper	-0.41	.678	-0.78	-2.31	.021	-6.34
Corpus Callosum (Anterior)	-0.38	.702	-0.21	1.20	.231	0.94
Cerebellum Cortex	-0.37	.712	-6.41	-2.92	.003	-75.54
4 th Ventricle	0.24	.812	0.55	-2.17	.030	-7.13
Cerebellum White Matter	-0.22	.829	-1.59	-0.43	.669	-4.58
Brain Stem	0.21	.835	1.28	-0.61	.544	-5.55
Amygdala	-0.03	.977	-0.02	-2.69	.007	-2.68

Fig 1. Associations of risk of lead exposure and area deprivation index with subcortical volume. Subcortical regions are sorted vertically by p-value (uncorrected) for risk of lead exposure. For each predictor (risk of lead exposure, area deprivation index), the shade of each cell reflects the strength of the association, with redder colors reflecting more positive associations and bluer colors reflecting more negative associations, in accordance with Fig 2. Analyses controlled for age, sex, parental education, race, ethnicity, family income, intracranial volume, and the interaction between family income and either lead risk or ADI. t(8508) = t-statistic, 8508 degrees of freedom. b = t-statistic regression coefficient (i.e., change in mm 3 regional volume with decile of lead risk or area deprivation index).

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[-0.040, -0.019]), and amygdalar volumes (r_p = -0.029 [-0.040, -0.018]) (Figs 1 and 2). In contrast, corpus callosal subregion volumes did not covary with ADI, $ps \ge .231$ (Tables 27–47 in S1 Appendix). To confirm these dissociations between ADI and lead risk, we reanalyzed these data with the housing-age metric as the predictor of interest. As in the lead-risk analyses, age of housing (i.e., housing-age-based estimates of the proportion of homes with lead-paint hazards) in children's neighborhoods was inversely associated with posterior (r_p = -0.026 [-0.037, -0.015]), mid-posterior (r_p = -0.029 [-0.040, -0.019]), central (r_p = -0.036 [-0.047, -0.025]), and mid-anterior corpus callosal volumes (r_p = -0.034 [-0.045, -0.023]) (Tables 48–68 in S1 Appendix), with the latter three passing FDR correction. Further, when ADI and lead risk (and their interactions with family income) were included in the same model, the FDR-corrected

Area Deprivation Index Cewm Cewm Cewm Cecmp Comp Comp

Partial Correlation Coefficient (r_p)

Fig 2. Regional associations between subcortical volume and risk of lead exposure (top) or area deprivation index (ADI) (bottom). For lead risk, these associations, correcting for false-discovery rate (FDR), were significant for midposterior, central, and mid-anterior corpus callosum. For ADI, subcortical gray matter, cerebellum cortex, accumbens area (not shown in this image), and amygdala. Regions are color-coded in correspondence to the effect size (i.e., partial correlation coefficient) of lead risk (top) and ADI (bottom), controlling for age, sex, parental education, race, ethnicity, family income, intracranial volume, and the interaction between family income and either lead risk or ADI. Regions with bolded outlines passed FDR correction. Blue-shaded regions indicate inverse associations between lead risk (or ADI) and volume (e.g., greater lead risk, lesser volume), while red-shaded regions indicate positive correlations (e.g., greater lead risk, greater volume). These images were generated in MATLAB using data from the *ggseg* toolbox in R [57]. $3v = 3^{rd}$ ventricle; $4v = 4^{th}$ ventricle; 4m = mamygdala; 4m = mam

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associations between lead risk and mid-anterior (r_p = -0.036 [-0.047, -0.025]), central (r_p = -0.034 [-0.045, -0.023]), and mid-posterior corpus callosal volumes (r_p = -0.030 [-0.040, -0.019]) were maintained (q < .05), as were the FDR-corrected associations between ADI and subcortical gray-matter (r_p = -0.034 [-0.045, -0.023]), amygdalar (r_p = -0.033 [-0.043, -0.022]), and cerebellar cortical volumes (r_p = -0.031 [-0.042, -0.020]) (q < .05) (Tables 117–137 in S1 Appendix). These sensitivity analyses suggest that lead-exposure risk, not income or ADI, was associated with diminished callosal subregion volumes.

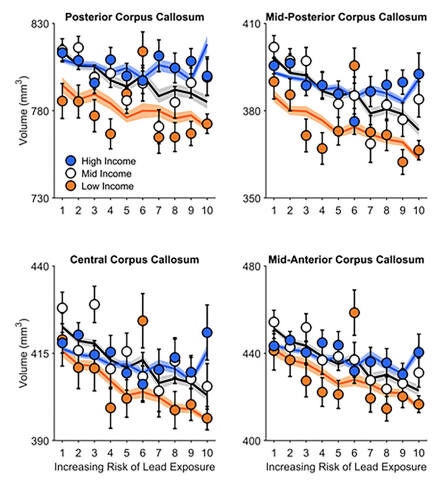


Fig 3. Posterior, mid-posterior, central, and mid-anterior corpus callosal volume significantly decreased with increasing risk of environmental lead exposure. These inverse associations were not significantly different between children from low-, mid-, and high-income families. Error bars represent ±1 between-subjects standard error of the observed means. The solid lines represent means of the marginal fitted values of the model; the shaded area surrounding the solid lines represent ±1 between-subjects standard error of those means. Age, sex, parental education, race, ethnicity, and intracranial volume were included as covariates in this analysis.

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Corpus callosum volume and cognition

Next, we analyzed how mid-posterior, central, and mid-anterior corpus callosal volumes were associated with NIH Toolbox performance (Tables 69–103 in S1 Appendix). Mid-posterior corpus callosal volume was positively associated with performance on the Dimensional Change Card Sort Test ($r_p = 0.042$ [0.031, 0.053]), the Flanker Inhibitory Control and Attention Test ($r_p = 0.038$ [0.027, 0.049]), the Pattern Comparison Processing Speed Test ($r_p = 0.035$ [0.024, 0.046]), the Oral Reading Recognition Test ($r_p = 0.034$ [0.024, 0.045]), and the Picture Vocabulary Test ($r_p = 0.056$ [0.045, 0.067]). Central corpus callosal volume was positively associated with Dimensional Change Card Sort Test ($r_p = 0.027$ [0.017, 0.038]) and Picture Vocabulary Test performance ($r_p = 0.033$ [0.022, 0.044]). Pattern Comparison Processing Speed Test performance was also positively associated with mid-anterior corpus callosal volume ($r_p = 0.022$ [0.011, 0.033]). Secondary analyses showed that anterior corpus callosal volume was positively associated with performance on the Dimensional Change Card Sort Test ($r_p = 0.022$ [0.012, 0.033]) and Picture Vocabulary Test ($r_p = 0.031$ [0.020, 0.042]), and posterior (like

mid-posterior) corpus callosal volume was positively associated with performance on the Dimensional Change Card Sort Test ($r_p = 0.045$ [0.034, 0.056]), the Flanker Inhibitory Control and Attention Test ($r_p = 0.029$ [0.019, 0.040]), the Pattern Comparison Processing Speed Test ($r_p = 0.032$ [0.022, 0.043]), the Oral Reading Recognition Test ($r_p = 0.035$ [0.024, 0.046]), and the Picture Vocabulary Test ($r_p = 0.058$ [0.047, 0.069]).

Neurostructurally-mediated associations between lead risk and cognition

We then examined whether mid-posterior, central, and mid-anterior callosal volumes mediated lead-risk relationships with cognitive performance (Tables 104–116 in S1 Appendix). Our results were consistent with mid-posterior corpus callosal volume mediating associations between lead risk and five cognitive tests (i.e., one test of executive functioning, two of language functioning, one of processing speed, and one of attention/inhibitory control) (Fig 4). Mediation was not present for central or mid-anterior corpus callosal volumes.

Discussion

Higher neighborhood-level risks of childhood lead exposure were associated with smaller mid-anterior, central, and mid-posterior corpus callosal volumes (i.e., the genu, truncus/body, and anterior splenium, respectively); to a lesser extent, lead-exposure risk was associated with smaller posterior corpus callosal volumes (i.e., posterior splenium) [56,58,59]. These associations were absent for ADI, but present for housing age. Thus, the lead-risk associations were likely attributable to factors associated with elevated lead-exposure risk (e.g., residual lead paint in older homes) [1,27]. Because ABCD enrolled 9- to 10-year-old participants at baseline, we cannot know whether these associations existed prior to baseline (i.e., pre- vs. postnatal insults) or disentangle them from other unmeasured confounding factors. However, the relationships between lead risk and callosal volume in mid-anterior to posterior, but not anterior, callosal regions are consistent with research showing earlier development of anterior than posterior corpus callosum, with the latter continuing to grow through adolescence (i.e., anteriorto-posterior maturation) [60-63]. While we recently reported the strongest inverse associations between lead-exposure risk and cortical structure in children of low-income families [28], the nonspecific inverse associations in callosal structure observed here suggest differential socioeconomic and environmental modulation of cortical and subcortical developmental trajectories, which is consistent with known spatially and temporally variable structural brain maturation [30,64].

Previous research on lead's impact on corpus callosal structure has been inconsistent. Stewart and colleagues [19] reported inverse correlations between bone-lead levels and posterior corpus callosal volume in occupationally exposed lead workers. Brubaker et al. [21] reported increased white matter integrity in the callosal genu, body, and splenium in adults exposed to lead during childhood, but Hsieh et al. [65] found nonsignificant differences in corpus callosal white-matter integrity in lead-exposed workers. Lasky et al. [66] reported no significant differences in callosal volume in lead-exposed (pre- or postnatal) versus non-lead-exposed rhesus monkeys, while Rai et al. [67] suggested that exposure to metal mixtures (arsenic, cadmium, and lead) in developing rats may thin the corpus callosum. There are no endogenous lead-exposure data yet in ABCD, but our study suggests that lead-exposure risk and its related environmental factors may be associated with corpus callosal morphology.

Exposures to other neurotoxicants are similarly associated with callosal morphology [68–72]. Prenatal alcohol exposure was most consistently associated with smaller areas of more posterior callosal regions in 8- to 22-year-olds [68,69], and prenatal particulate matter air pollution exposure, especially during the 3rd trimester, was most strongly associated with smaller

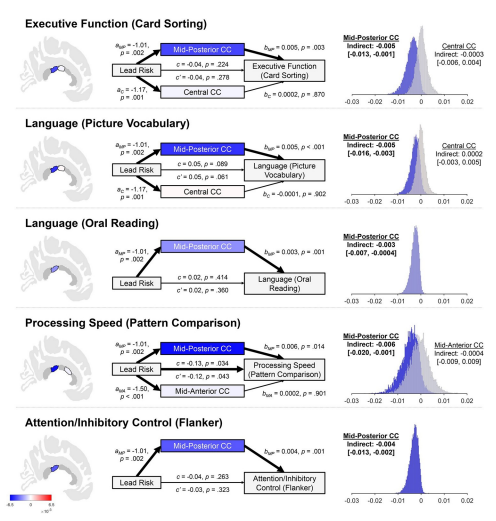


Fig 4. Mediational analyses showing indirect associations of lead risk on cognitive performance through midposterior corpus callosal (CC) volume. The title of each row [e.g., "Executive Function (Card Sorting)"] refers to the construct and task used to assess that construct. The shading of the callosal regions, the mediator boxes (e.g., Mid-Posterior CC, Central CC), and histograms reflect the strength of the indirect association (per the color bar in the lower left corner). Callosal regions in dark gray were not incorporated as mediators in the corresponding analysis. Total effects of lead risk are represented by c, direct effects of lead risk are represented by c', and a and b values refer to the associations of lead risk on callosal volume and callosal volume on cognitive performance, respectively. The subscript of the a and b values refer to the callosal subregion (i.e., C = central, MA = mid-anterior, MP = midposterior). All a, b, c, and c'values are unstandardized regression coefficients. Thick arrows designate significant associations. Thin arrows designate non-significant associations. The presence of the statistical significance of indirect associations was determined by construction of bias-corrected percentile bootstrapping 95% confidence intervals (i.e., 10,000 bootstrapped samples), irrespective of the statistical significance of the corresponding total effect. The distributions of these bootstrapped indirect associations are shown, along with the original-sample indirect associations and corresponding confidence intervals. Bolded font indicates statistically significant indirect associations. Age, sex, parental education, race, ethnicity, and family income were included as covariates in analyses. Card Sorting = Dimensional Change Card Sort Task. Picture Vocabulary = Picture Vocabulary Test. Oral Reading = Oral Reading Recognition Test. Pattern Comparison = Pattern Comparison Processing Speed Test. Flanker = Flanker Inhibitory Control and Attention Test.

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volumes of the callosal body in 8- to 12-year-olds (mid-anterior + central + mid-posterior) [70]. Posterior corpus callosal size was 3.1% (7.9%) smaller in 12- to 18-year-old males (females) prenatally exposed to cigarette smoking [71], and splenial (posterior corpus callosal) size in 4.5-year-olds was 1.4% smaller given high versus low levels of prenatal polychlorinated-

biphenyl exposure [72]. Here, we found a 5.2% mean decrease in mid-posterior, central, and mid-anterior corpus callosal volume in individuals living in census tracts with the highest versus lowest lead-risk score.

Lead's well-established effects on cognition and potential associations with corpus callosal structure are complemented by relationships between corpus callosal structure, cognition, and intelligence [73]. In our sample, anterior callosal regions were weakly associated with cognitive function whereas posterior regions were more strongly associated with cognitive performance. The association between mid-posterior callosal structure and processing speed (i.e., the Pattern Comparison Processing Speed Test) corroborates research on callosal volume and processing speed in occupationally lead-exposed and non-exposed adult men [18]. Our reported associations between mid-posterior callosal volume and performances on the Picture Vocabulary Test and the Oral Reading Recognition Test are also consistent with research suggesting that posterior callosal regions are critical for interhemispheric transfer between temporal-parietal-occipital cortical regions involved in language processing [74–77].

The lead-risk metric was primarily a function of the estimated neighborhood-level prevalence of lead-based paint given the age of houses in that neighborhood [25,26]. Even though lead-based house paint and leaded gasoline (for on-road vehicles) were banned in the US in 1978 [78] and 1996 [79], children remain at risk for ingesting lead via (1) drinking water provided through lead service lines (i.e., lead-containing plumbing) [80], (2) lead-contaminated dust and soil given the prior use of lead-based paint in older buildings [27], and (3) lead-contaminated topsoil from past leaded-gasoline vehicle emissions [81]. Indeed, while average blood lead levels have substantially declined over the past several decades, a 2021 study estimated that nearly 400,000 1-to-11-year-olds in 2011-2016 had blood-lead levels exceeding the CDC's reference level of 5 µg/dL [82]. Here, about one in five houses in our participants' neighborhoods were estimated to contain lead-based paint hazards, which is consistent with national surveys showing that 25% of United States housing stock contains one or more lead hazards [27]. Of 14 risk factors known to impair neurodevelopment (e.g., medical conditions, low SES), lead exposure was reported as 2nd only to preterm birth in its impact on total reduction in IQ [83]. Lead-associated decrements in IQ have been suggested to contribute to annual costs of approximately \$977 billion in low- and middle-income countries [17], many of which do not have regulations as to limits of the concentration of lead in paint [84]. Accordingly, reducing risks of lead exposure (and addressing related environmental factors) will have economic and health benefits that facilitate prosperity in the United States and worldwide.

Lead risk and housing age were distinctly associated with callosal morphology. In contrast, ADI, a metric of neighborhood socioeconomic disadvantage [40,41], was inversely associated with subcortical gray-matter, cerebellar cortical, accumbens area, and amygdalar volumes, but not callosal morphology. Despite research on the effects of neighborhood SES on child development (e.g. ADI) [85] and family-specific SES measures on children's brain structure [86–88], less is known about how neighborhood-level metrics of disadvantage influence neurocognitive development [87,89]. Our results are consistent with previous reports showing (1) smaller cortical and subcortical volumes in 8- to 21-year-olds living in low-SES neighborhoods [90] and (2) greater age-related increases in right amygdalar volume in adolescents living in more disadvantaged neighborhoods (potentially due to the groups' lower baseline amygdalar volumes) [91]. Given associations between amygdalar volume and depressive symptoms [92], it will be critical to evaluate longitudinal trajectories of mental health and brain development in ABCD and whether neighborhood characteristics increase risk or promote resilience to developmental insults such as lead neurotoxicity.

Limitations and future directions

In the current report, the primary residential addresses of 9- to 10-year-old children were used to derive community-based risk estimates of lead exposure, which our past research has shown are valid proxies of exposure [28]. Peak lead exposure during childhood occurs in toddlers (i.e., 2- to 3-years-old) from the confluence of hand-to-mouth ingestion of lead-contaminated floor dust, soil, water, and paint chips [93], but older children are also vulnerable to lead toxicity [28,94,95]. For example, IQ in older children has been shown to be better predicted by concurrent than past blood-lead levels [94,96], and, even at low levels of exposure, IQ was shown to be associated with concurrent blood-lead levels in 7- to 14-year-olds [97,98]. Ultimately, the age of greatest vulnerability to lead neurotoxicity is unclear [95], but recent evidence has suggested that exposure to other neurotoxicants (i.e., air pollution) is also associated with brain structure in 9- to 10-year-olds [99].

As we cannot physically manipulate lead exposure in ABCD participants, the longitudinal design of the ABCD Study and its diverse cohort offer opportunities to evaluate how developmental trajectories of brain and cognitive development are associated with differential risks of lead exposure. The ABCD Study does not yet have endogenous lead-exposure data in its participants, but it has been actively collecting address histories for its participants since birth, which, when completed, will facilitate understanding of the critical developmental periods of lead neurotoxicity vulnerability. Similarly, even though ABCD's data collection sites are primarily in metropolitan areas, the recruitment areas of these 21 sites represent at least 20% of the 9-to-10-year-old US population [35]. Further, while the potential for reverse causation is inherent to cross-sectional studies, thereby limiting causal inference, it is unlikely that poor cognitive performance elicits altered brain structure, or that altered brain structure induces risk factors of lead exposure here, thus supporting the temporal ordering within our cross-sectional mediational analyses (lead risk \rightarrow brain structure \rightarrow cognitive performance) [100].

While ADI and lead risk were strongly correlated, over 85% of the variance in lead risk was not accounted for by ADI, reflecting a possible necessity to analyze multiple factors of environmental health disparities in future research [101]. Specifically, our analyses showed dissociations between the subcortical regions associated with lead risk and ADI, and the associations between lead risk and corpus callosal volumes were maintained when including ADI in another set of models. Thus, our results suggest that subcortical brain structure (i.e., corpus callosal volume) in adolescents may be uniquely associated with factors related to increased risk of lead exposure (i.e., age of housing, with older homes more likely to contain lead-based paint hazards) [27].

Even though the ADI and lead-risk data reflect community- rather than individual-level estimates, past research has argued that using community-level data to evaluate environmental risks may ultimately prevent exposure to the hazards before the individual is actually exposed to them (i.e., screening communities/homes before occupancy) [1,102]. Indeed, individual screening questionnaires of potential lead exposure may not accurately identify the children with elevated blood lead levels [103]. Therefore, while community-level risk estimates inherently over- and underestimate risks of specific individuals in that community, future incorporation of such geocoded data in cognitive neuroscience research may considerably advance understanding of the environmental contextualization of an individual's neurocognitive and brain development, especially given the more common practice of collecting data pertaining to the individual (e.g., SES).

As individuals are rarely exposed to isolated chemicals (but to mixtures of chemicals) [104], the incorporation of multiple data sources reflecting "mixtures" of environmental health disparities may also offer substantial insight into the collective and synergistic factors to target in

environmental remediation interventions. Simply, the risk of lead exposure does not exist in a vacuum but is associated with past and current practices that have differentially subjected children to such risks. For example, the burden of lead-exposure's effects is typically greatest in children in the lowest SES families [105-108], a glaring example of environmental injustice [105,109]. Similarly, Black and Hispanic children tend to have greater mean blood lead levels than white children [5,110-112] and are more likely than white children to live in homes or regions with greater risks of lead exposure [113-115]. Further, lead-poisoning rates (and, thus, children's blood-lead levels) are associated with multiple community-level factors [39], including value and age of houses, poverty rates, population density, and percentage of the population who are Black or Hispanic [1,116], signifying racial residential segregation as a potential explanatory mechanism for lead exposure disparities [117]. While the data in the current manuscript may reflect differences in lead exposure, these differences would then ultimately be due to disparate conditions that initially elicited such differences, thereby focusing any potential intervention efforts on the originating disparities. Indeed, recent research has shown that soillead concentrations tended to be elevated in samples taken from historically redlined neighborhoods compared to those in "best" and "desirable" neighborhoods, per zone designations by the 1930's Home Owners' Loan Corporation [118]. Ultimately, because we do not currently have endogenous lead-exposure data in our participants, the results related to risk of lead exposure may be alternatively explained (at least partially) by these other systemic and environmental factors. Accordingly, upon collection of bodily lead data and additional geocoded data in ABCD, our future research will involve analyses of both chemical and "environmentaldisparity mixtures" to both study developmental trajectories more comprehensively and evaluate the relative strengths of the associations between adolescent development and other sources of disparity (e.g., air pollution, residential segregation).

Conclusion

Lead-induced cognitive deficits are likely governed by how lead exposure influences brain structure [18], and our results, consistent with callosal mediation of lead-risk associations with cognition, offer the testable hypothesis in future studies that "dose" of lead exposure mediates cognitive functioning through changes in mid-posterior corpus callosal structure. ABCD is measuring lead concentrations in shed deciduous (baby) teeth [119–121], but those data will not be available for several years. ABCD is also exploring the collection and analysis of blood lead levels to gauge how well they are correlated with neighborhood-level lead-risk estimates. ABCD is also working to incorporate electronic health records as part of its dataset, which may also help elucidate these relationships via past lead screening results. Until then, this study, which uses neighborhood-level lead-exposure risk, provides potential evidence that cognitive deficits from low-level lead toxicity (and its related environmental factors) may operate by diminishing subcortical brain structure [28].

Supporting information

S1 Appendix. Supplementary figures and statistical output tables. (DOCX)

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ABCD Acknowledgement. Data used in the preparation of this article were obtained from the Adolescent Brain Cognitive Development (ABCD) Study (https://abcdstudy.org/), held in the NIMH Data Archive (NDA). This is a multisite, longitudinal study designed to recruit more

than 10,000 children aged 9–10 and follow them over 10 years into early adulthood. A full list of supporters is available at https://abcdstudy.org/federal-partners/. A listing of participating sites and a complete listing of the study investigators can be found at https://abcdstudy.org/principal-investigators/. ABCD consortium investigators designed and implemented the study and/or provided data but did not necessarily participate in analysis or writing of this report. This manuscript reflects the views of the authors and may not reflect the opinions or views of the NIH or other ABCD consortium investigators. The ABCD data repository grows and changes over time. The ABCD data used in this report came from https://doi.org/10.15154/1523347.

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Writing – review & editing: Andrew T. Marshall, Rob McConnell, Bruce P. Lanphear, Wesley K. Thompson, Megan M. Herting, Elizabeth R. Sowell.

References

- Lanphear BP, Byrd RS, Auinger P, Schaffer SJ. Community characteristics associated with elevated blood lead levels in children. Pediatrics. 1998; 101(2):264–71. https://doi.org/10.1542/peds.101.2.264 PMID: 9445502
- Reuben A, Caspi A, Belsky DW, Broadbent J, Harrington H, Sugden K, et al. Association of childhood blood lead levels with cognitive function and socioeonomic status at age 38 years and with IQ change and socioeconomic mobility between childhood and adulthood. JAMA. 2017; 317(12):1244–51. https://doi.org/10.1001/jama.2017.1712 PMID: 28350927
- Bellinger DC. Very low lead exposures and children's neurodevelopment. Current Opinion in Pediatrics. 2008; 20:172–7. https://doi.org/10.1097/MOP.0b013e3282f4f97b PMID: 18332714.
- Canfield RL, Henderson CR, Cory-Slechta DA, Cox C, Jusko TA, Lanphear BP. Intellectual impairments in children with blood lead concentrations below 10 ug per deciliter. The New England Journal of Medicine. 2003; 348(16):1517–26. https://doi.org/10.1056/NEJMoa022848 PMID: 12700371; PubMed Central PMCID: PMC4046839.
- Lanphear BP, Dietrich K, Auinger P, Cox C. Cognitive deficits associated with blood lead concentrations <10 ug/dL in US children and adolescents. Public Health Reports. 2000; 115:521–9. https://doi.org/10.1093/phr/115.6.521 PMID: 11354334; PubMed Central PMCID: PMC1308622.
- Needleman HL, Gunnoe C, Levinton A, Reed R, Peresie H, Maher C, et al. Deficits in psychologic and classroom performance of children with elevated dentine lead levels. The New England Journal of Medicine. 1979; 300(13). Epub 689. https://doi.org/10.1056/NEJM197903293001301 PMID: 763299.
- Wasserman GA, Factor-Litvak P, Liu X, Todd AC, Kline JK, Slavkovich V, et al. The relationship between blood lead, bone lead and child intelligence. Child Neuropsychology. 2003; 9(1):22–34. https://doi.org/10.1076/chin.9.1.22.14497 PMID: 12815520
- Dietrich KN, Ris MD, Succop PA, Berger OG, Bornschein RL. Early exposure to lead and juvenile delinquency. Neurotoxicology and Teratology. 2001; 23:511–8. https://doi.org/10.1016/s0892-0362 (01)00184-2 PMID: 11792521.
- Wright JP, Dietrich KN, Ris MD, Hornung RW, Wessel SD, Lanphear BP, et al. Association of prenatal and childhood blood lead concentrations with criminal arrests in early adulthood. PLOS Medicine. 2008; 5(5):e101. https://doi.org/10.1371/journal.pmed.0050101 PMID: 18507497; PubMed Central PMCID: PMC2689664.
- Braun JM, Kahn RS, Froehlich T, Auinger P, Lanphear BP. Exposures to environmental toxicants and attention deficit hyperactivity disorder in U.S. children. Environmental Health Perspectives. 2006;

- 114:1904–9. https://doi.org/10.1289/ehp.9478 PMID: 17185283; PubMed Central PMCID: PMC1764142.
- Guilarte TR, Opler M, Pletnikov M. Is lead exposure in early life an environmental risk factor for schizophrenia? Neurobiological connections and testable hypotheses. NeuroToxicology. 2012; 33(3):560– 74. https://doi.org/10.1016/j.neuro.2011.11.008 PMID: 22178136; PubMed Central PMCID: PMC3647679.
- 12. Mazumdar M, Xia W, Hofmann O, Gregas M, Ho Sui S, Hide W, et al. Prenatal lead levels, plasma amyloid β levels, and gene expression in young adulthood. Environmental Health Perspectives. 2012; 120(5):702–7. https://doi.org/10.1289/ehp.1104474 PMID: 22313790; PubMed Central PMCID: PMC3346789.
- 13. Wu J, Basha MR, Brock B, Cox DP, Cardozo-Pelaez F, McPherson CA, et al. Alzheimer's disease (AD)-like pathology in aged monkeys after infantile exposure to environmental metal lead (Pb): Evidence for a developmental origin and environmental link for AD. The Journal of Neuroscience. 2008; 28(1):3–9. https://doi.org/10.1523/JNEUROSCI.4405-07.2008 PMID: 18171917; PubMed Central PMCID: PMC2486412.
- Fewtrell LJ, Pruss-Ustun A, Landrigan P, Ayuso-Mateos JL. Estimating the global burden of disease of mild mental retardation and cardiovascular diseases from environmental lead exposure. Environ Res. 2004; 94(2):120–33. https://doi.org/10.1016/s0013-9351(03)00132-4 PMID: 14757375.
- Falk H. International environmental health for the pediatrician: Case study of lead poisoning. Pediatrics. 2003: 112(1):259

 –64. PMID: 12837919.
- Nriagu JO, Blankson ML, Ocran K. Childhood lead poisoning in Africa: a growing public health problem. The Science of the Total Environment. 1996; 181:93–100. https://doi.org/10.1016/0048-9697(95) 04954-1 PMID: 8820380.
- Attina TM, Trasande L. Economic costs of childhood lead exposure in low- and middle-income countries. Environmental Health Perspectives. 2013; 121(9):1097–102. https://doi.org/10.1289/ehp.1206424 PMID: 23797342
- Schwartz BS, Chen S, Caffo B, Stewart WF, Bolla KI, Yousem D, et al. Relations of brain volumes with cognition function in males 45 years and older with past lead exposure. NeuroImage. 2007; 37:633– 41. https://doi.org/10.1016/j.neuroimage.2007.05.035 PMID: 17600728
- Stewart WF, Schwartz BS, Davatzikos C, Shen D, Liu D, Wu X, et al. Past adult lead exposure is linked to neurodegeneration measured by brain MRI. Neurology. 2006; 66:1476–84. https://doi.org/10.1212/01.wnl.0000216138.69777.15 PMID: 16717205
- 20. Brubaker CJ, Dietrich KN, Lanphear BP, Cecil KM. The influence of age of lead exposure on adult gray matter volume. NeuroToxicology. 2010; 31:259–66. https://doi.org/10.1016/j.neuro.2010.03.004 PMID: 20226811; PubMed Central PMCID: PMC2866835.
- Brubaker CJ, Schmithorst VJ, Haynes EN, Dietrich KN, Egelhoff JC, Lindquist DM, et al. Altered myelination and axonal integrity in adults with childhood lead exposure: A diffusion tensor imaging study. NeuroToxicology. 2009; 30:867–75. https://doi.org/10.1016/j.neuro.2009.07.007 PMID: 19619581; PubMed Central PMCID: PMC2789851.
- 22. Cecil KM, Brubaker CJ, Adler CM, Dietrich K, Altaye M, Egelhoff JC, et al. Decreased brain volume in adults with childhood lead exposure. PLOS Medicine. 2008; 5(5):e112. https://doi.org/10.1371/journal.pmed.0050112 PMID: 18507499; PubMed Central PMCID: PMC2689675.
- Cecil KM. Effects of early low-level lead exposure on human brain structure, organization and functions. Journal of Developmental Origins of Health and Disease. 2011; 2(1):17–24. https://doi.org/10.1017/S2040174410000486
- Trope I, Lopez-Villegas D, Cecil KM, Lenkinski RE. Exposure to lead appears to selectively alter metabolism of cortical gray matter. Pediatrics. 2001; 107(6):1437–42. https://doi.org/10.1542/peds.107.6.1437 PMID: 11389272
- Washington Tracking Network. Childhood Lead Risk Map 2017. Available from: https://fortress.wa.gov/doh/wtn/WTNPortal/.
- **26.** Vox. The risk of lead poisoning isn't just in Flint. So we mapped the risk in every neighborhood in America 2016. Available from: https://www.vox.com/a/lead-exposure-risk-map.
- Jacobs DE, Clickner RP, Zhou JY, Viet SM, Marker DA, Rogers JW, et al. The prevalence of lead-based paint hazards in U.S. housing. Environmental Health Perspectives. 2002; 110:A599–A606. https://doi.org/10.1289/ehp.021100599 PMID: 12361941; PubMed Central PMCID: PMC1241046.
- 28. Marshall AT, Betts S, Kan EC, McConnell R, Lanphear BP, Sowell ER. Association of lead-exposure risk and family income with childhood brain outcomes. Nature Medicine. 2020; 26:91–7. https://doi.org/10.1038/s41591-019-0713-y PMID: 31932788; PubMed Central PMCID: PMC6980739.

- Sowell ER, Trauner DA, Gamst A, Jernigan TL. Development of cortical and subcortical brain structures in childhood and adolescence: A structural MRI study. Developmental Medicine & Child Neurology. 2002; 44(1):4–16. https://doi.org/10.1017/s0012162201001591 PMID: 11811649
- Giedd JN, Blumenthal J, Jeffries NO, Castellanos FX, Liu H, Zijdenbos A, et al. Brain development during childhood and adolescence: A longitudinal MRI study. Nature Neuroscience. 1999; 2(10):861–3. https://doi.org/10.1038/13158 PMID: 10491603
- Wheeler DC, Jones RM, Schootman M, Nelson EJ. Explaining variation in elevated blood lead levels among children in Minnesota using neighborhood socioeconomic variables. Science of the Total Environment. 2019; 650:970–7. https://doi.org/10.1016/j.scitotenv.2018.09.088 PMID: 30308871
- 32. Luciana M, Bjork JM, Nagel BJ, Barch DM, Gonzalez R, Nixon SJ, et al. Adolescent neurocognitive development and impacts of substance use: Overview of the adolescent brain cognitive development (ABCD) baseline neurocognition battery. Developmental Cognitive Neuroscience. 2018; 32:67–79. https://doi.org/10.1016/j.dcn.2018.02.006 PMID: 29525452; PubMed Central PMCID: PMC6039970.
- Gershon RC, Wagster MV, Hendrie HC, Fox NA, Cook KF, Nowinski CJ. NIH toolbox for assessment of neurological and behavioral function. Neurology. 2013; 80:S2–S6. https://doi.org/10.1212/WNL. 0b013e3182872e5f PMID: 23479538; PubMed Central PMCID: PMC3662335.
- 34. Jernigan TL, Brown SA, Dowling GJ. The Adolescent Brain Cognitive Development Study. Journal of Research on Adolescence. 2018; 28(1):154–6. https://doi.org/10.1111/jora.12374 PMID: 29460352.
- Garavan H, Bartsch H, Conway K, Decastro A, Goldstein RZ, Heeringa S, et al. Recruiting the ABCD sample: Design considerations and procedures. Developmental Cognitive Neuroscience. 2018; 32:16–22. https://doi.org/10.1016/j.dcn.2018.04.004 PMID: 29703560; PubMed Central PMCID: PMC6314286.
- Auchter AM, Hernandez Mejia M, Heyser CJ, Shilling PD, Jernigan TL, Brown SA, et al. A description
 of the ABCD organizational structure and communication framework. Dev Cogn Neurosci. 2018;
 32:8–15. Epub 2018/05/01. https://doi.org/10.1016/j.dcn.2018.04.003 PMID: 29706313; PubMed
 Central PMCID: PMC6462277.
- U.S. Census Bureau. American Community Survey. Available at: https://www.census.gov/programssurveys/acs/about.html.
- The ABCD Consortium. Dataset: Release 2.0 and Fix Release 2.0.1. 2019. https://doi.org/10.15154/1503209
- 39. Akkus C, Ozdenerol E. Exploring childhood lead exposure through GIS: A review of the recent literature. International Journal of Environmental Research and Public Health. 2014; 11:6314–34. https://doi.org/10.3390/ijerph110606314 PMID: 24945189
- 40. Kind AJH, Jencks S, Brock J, Yu M, Bartels C, Ehlenbach W, et al. Neighborhood socioeconomic disadvantage and 30-day rehospitalization: A retrospective cohort study. Annals of Internal Medicine. 2014; 161(11):765–74. https://doi.org/10.7326/M13-2946 PMID: 25437404; PubMed Central PMCID: PMC4251560.
- Singh GK. Area deprivation and widening inequalities in US mortality, 1969–1998. American Journal of Public Health. 2003; 93(7):1137–43. https://doi.org/10.2105/ajph.93.7.1137 PMID: 12835199; PubMed Central PMCID: PMC1447923.
- 42. Casey BJ, Cannonier T, Conley MI, Cohen AO, Barch DM, Heitzeg MM, et al. The Adolescent Brain Cognitive Development (ABCD) study: Imaging acquisition across 21 sites. Developmental Cognitive Neuroscience. 2018; 32:43–54. https://doi.org/10.1016/j.dcn.2018.03.001 PMID: 29567376; PubMed Central PMCID: PMC5999559.
- 43. Barch DM, Albaugh MD, Avenevoli S, Chang L, Clark DB, Glantz MD, et al. Demographic, physical and mental health assessments in the adolescent brain and cognitive development study: Rationale and description. Developmental Cognitive Neuroscience. 2018; 32:55–66. https://doi.org/10.1016/j.dcn.2017.10.010 PMID: 29113758
- 44. Weintraub S, Dikmen SS, Heaton RK, Tulsky DS, Zelazo PD, Bauer PJ, et al. Cognition assessment using the NIH Toolbox. Neurology. 2013; 80:S54–S64. https://doi.org/10.1212/WNL. 0b013e3182872ded PMID: 23479546
- 45. Hagler DJ, Hatton SN, Cornejo MD, Makowski C, Fair DA, Dick AS, et al. Image processing and analysis methods for the Adolescent Brain Cognitive Development Study. NeuroImage. 2019; 202:116091. https://doi.org/10.1016/j.neuroimage.2019.116091 PMID: 31415884.
- Li Y, Thompson WK, Reuter C, Brown J, Dougherty RF, Desikan R, et al. Incidental findings on structural brain MRI in the ABCD Study.
- 47. Noble KG, Houston SM, Brito NH, Bartsch H, Kan E, Kuperman JM, et al. Family income, parental education and brain structure in children and adolescents. Nature Neuroscience. 2015; 18(5):773–8. https://doi.org/10.1038/nn.3983 PMID: 25821911; PubMed Central PMCID: PMC4414816.

- 48. Akshoomoff N, Newman E, Thompson WK, McCabe C, Bloss CS, Chang L, et al. The NIH Toolbox cognition battery: Results from a large normative developmental sample (PING). Neuropsychology. 2014; 28(1):1–10. https://doi.org/10.1037/neu0000001 PMID: 24219608
- Benjamini Y, Hochberg Y. Controlling the false discovery rate: A practical and powerful approach to multiple testing. Journal of the Royal Statistical Society Series B (Methodological). 1995; 57(1):289– 300.
- 50. West SG, Aiken LS, Krull JL. Experimental personality designs: Analyzing categorical by continuous variable interactions. Journal of Personality. 1996; 64(1):1–48. https://doi.org/10.1111/j.1467-6494. 1996.tb00813.x PMID: 8656311
- MacKinnon DP, Lockwood CM, Williams J. Confidence limits for the indirect effect: Distribution of the product and resampling methods. Multivariate Behavioral Research. 2004; 39(1):99–128. https://doi.org/10.1207/s15327906mbr3901 4 PMID: 20157642
- **52.** Rucker DD, Preacher KJ, Tormala ZL, Petty RE. Mediation analysis in social psychology: Current practices and new recommendations. Social and Personality Psychology Compass. 2011; 5(6):359–71. https://doi.org/10.1111/j.1751-9004.2011.00355.x
- 53. Preacher KJ, Hayes AF. Asymptotic and resampling strategies for assessing and comparing indirect effects in multiple mediator models. Behavior Research Methods. 2008; 40(3):879–91. https://doi.org/10.3758/brm.40.3.879 PMID: 18697684
- 54. Nakagawa S, Cuthill IC. Effect size, confidence interval and statistical significance: A practical guide for biologists. Biological Reviews. 2007; 82:591–605. https://doi.org/10.1111/j.1469-185X.2007. 00027.x PMID: 17944619
- Aloe AM, Thompson CG. The synthesis of partial effect sizes. journal of the Society for Social Work and Research. 2013; 4(4):390–405. https://doi.org/10.5243/jsswr.2013.24
- 56. Fischl B, Salat DH, Busa E, Albert M, Dieterich M, Haselgrove C, et al. Whole brain segmentation: Automated labeling of neuroanatomical structures in the human brain. Neuron. 2002; 33:341–55. https://doi.org/10.1016/s0896-6273(02)00569-x PMID: 11832223
- Mowinckel AM, Vidal-Piñeiro D. Visualisation of Brain Statistics with R-packages ggseg and ggseg3d. arXiv preprint arXiv:191208200. 2019.
- 58. Collinson SL, Gan SC, Woon PS, Kuswanto C, Sum MY, Yang GL, et al. Corpus callosum morphology in first-episode and chronic schizophrenia: combined magnetic resonance and diffusion tensor imaging study of Chinese Singaporean patients. The British Journal of Psychiatry. 2014; 204:55–60. https://doi.org/10.1192/bjp.bp.113.127886 PMID: 24202961
- **59.** Francis A, Bhojraj TS, Prasad K, Kulkarni S, Montrose D, Eack S, et al. Abnormalities of the corpus callosum in non-psychotic high-risk offspring of schizophrenia patients. Psychiatry Research: Neuroimaging. 2011; 191(1):9–15. https://doi.org/10.1016/j.pscychresns.2010.09.007 PMID: 21145214
- Luders E, Thompson PM, Toga AW. The development of the corpus callosum in the healthy human brain. The Journal of Neuroscience. 2010; 30(33):10985–90. https://doi.org/10.1523/JNEUROSCI.5122-09.2010 PMID: 20720105
- 61. Thompson PM, Giedd JN, Woods RP, MacDonald D, Evans AC, Toga AW. Growth patterns in the developing brain detected by using continuum mechanical tensor maps. Nature. 2000; 404:190–3. https://doi.org/10.1038/35004593 PMID: 10724172
- 62. Giedd JN, Rumsey JM, Castellanos FX, Rajapakse JC, Kaysen D, Vaituzis AC, et al. A quantitative MRI study of the corpus callosum in children and adolescents. Developmental Brain Research. 1996; 91:274–80. https://doi.org/10.1016/0165-3806(95)00193-x PMID: 8852379
- 63. Giedd JN, Blumenthal J, Jeffries NO, Rajapakse JC, Vaituzis AC, Liu H, et al. Development of the human corpus callosum during childhood and adolescence: A longitudinal MRI study. Progress in Neuro-Psychopharmacology and Biological Psychiatry. 1999; 23(4):571–88. https://doi.org/10.1016/s0278-5846(99)00017-2 PMID: 10390717
- 64. Raznahan A, Shaw PW, Lerch JP, Clasen LS, Greenstein D, Berman R, et al. Longitudinal four-dimensional mapping of subcortical anatomy in human development. Proceedings of the National Academy of Sciences. 2014; 111(4):1592–7. https://doi.org/10.1073/pnas.1316911111 PMID: 24474784; PubMed Central PMCID: PMC3910572.
- 65. Hsieh T-J, Chuang H-Y, Chen Y-C, Wang C-L, Lan S-H, Liu GC, et al. Subclinical white matter integrity in subjects with cumulative lead exposure. Radiology. 2009; 252(2):509–17. https://doi.org/10.1148/ radiol.2522080653 PMID: 19546424
- Lasky RE, Luck ML, Parikh NA, Laughlin NK. The effects of early lead exposure on the brains of adult rhesus monkeys: A volumetric MRI study. Toxicological Sciences. 2005; 85(2):963–75. https://doi.org/10.1093/toxsci/kfi153 PMID: 15788724

- 67. Rai NK, Ashok A, Rai A, Tripathi S, Nagar GK, Mitra K, et al. Exposure to As, Cd and Pb-mixture impairs myelin and axon development in rat brain, optic nerve and retina. Toxicology and Applied Pharmacology. 2013; 273(2):242–58. https://doi.org/10.1016/j.taap.2013.05.003 PMID: 23680456
- Riley EP, Mattson SN, Sowell ER, Jernigan TL, Sobel DF, Jones KL. Abnormalities of the corpus callosum in children prenatally exposed to alcohol. Alcoholism: Clinical and Experimental Research. 1995; 19(5):1198–202.
- Sowell ER, Mattson SN, Thompson PM, Jernigan TL, Riley EP, Toga AW. Mapping callosal morphology and cognitive correlates: Effects of heavy prenatal alcohol exposure. Neurology. 2001; 57(2):235–44. https://doi.org/10.1212/wnl.57.2.235 PMID: 11468307.
- Mortamais M, Pujol J, Martínez-Vilavella G, Fenoll R, Reynes C, Sabatier R, et al. Effects of prenatal exposure to particular matter air pollution on corpus callosum and behavioral problems in children. Environmental Research. 2019; 178:108734. https://doi.org/10.1016/j.envres.2019.108734 PMID: 31539824
- Paus T, Nawazkhan I, Leonard G, Perron M, Pike GB, Pitiot A, et al. Corpus callosum in adolescent offspring exposed prenatally to maternal cigarette smoking. NeuroImage. 2008; 40(2):435–41. https:// doi.org/10.1016/j.neuroimage.2007.10.066 PMID: 18221892.
- Stewart P, Fitzgerald S, Reihman J, Gump B, Lonky E, Darvill T, et al. Prenatal PCB exposure, the corpus callosum, and response inhibition. Environmental Health Perspectives. 2003; 111:1670–7. https://doi.org/10.1289/ehp.6173 PMID: 14527849
- Luders E, Narr KL, Thompson PM, Toga AW. Neuroanatomical correlates of intelligence. Intelligence. 2009; 37(2):156–63. https://doi.org/10.1016/j.intell.2008.07.002 PMID: 20160919
- 74. Hines M, Chiu L, McAdams LA, Bentler PM, Lipcamon J. Cognition and the corpus callosum: Verbal fluency, visuospatial ability, and language lateralization related to midsagittal surface areas of callosal subregions. Behavioral Neuroscience. 1992; 106(1):3–14. https://doi.org/10.1037//0735-7044.106.1.3 PMID: 1554435
- **75.** Hofer S, Frahm J. Topography of the human corpus callosum revisited—Comprehensive fiber tractography using diffusion tensor magnetic resonance imaging. NeuroImage. 2006; 32:989–94. https://doi.org/10.1016/j.neuroimage.2006.05.044 PMID: 16854598
- 76. de Lacoste MC, Kirkpatrick JB, Ross ED. Topography of the human corpus callosum. Journal of Neuropathology & Experimental Neurology. 1985; 44(6):578–91. https://doi.org/10.1097/00005072-198511000-00004 PMID: 4056827
- 77. Fryer SL, Frank LR, Spadoni AD, Theilmann RJ, Nagel BJ, Schweinsburg AD, et al. Microstructural integrity of the corpus callosum linked with neuropsychological performance in adolescents. Brain and Cognition. 2008; 67(2):225–33. https://doi.org/10.1016/j.bandc.2008.01.009 PMID: 18346830
- Gilbert SG, Weiss B. A rationale for lowering the blood lead action level from 10 to 2 ug/dL. NeuroToxicology. 2006; 27:693–701. https://doi.org/10.1016/j.neuro.2006.06.008 PMID: 16889836; PubMed Central PMCID: PMC2212280.
- **79.** EPA Takes Final Step in Phaseout of Leaded Gasoline 1996 [cited 2020]. Available from: https://archive.epa.gov/epa/aboutepa/epa-takes-final-step-phaseout-leaded-gasoline.html.
- **80.** Triantafyllidou S, Edwards M. Lead (Pb) in tap water and in blood: Implications for lead exposure in the United States. Critical Reviews in Environmental Science and Technology. 2012; 42(13):1297–352.
- 81. Mielke HW, Gonzales CR, Powell ET, Laidlaw MAS, Berry KJ, Mielke PW, et al. The concurrent decline of soil lead and children's blood lead in New Orleans. Proceedings of the National Academy of Sciences. 2019; 116(44):22058–64. https://doi.org/10.1073/pnas.1906092116 PMID: 31611401; PubMed Central PMCID: PMC6825258.
- 82. Egan KB, Cornwell CR, Courtney JG, Ettinger AS. Blood lead levels in U.S. children ages 1–11 years, 1976–2016. Environmental Health Perspectives. 2021; 129(3):037003. https://doi.org/10.1289/EHP7932 PMID: 33730866
- 83. Bellinger DC. A strategy for comparing the contributions of environmental chemicals and other risk factors to neurodevelopment of children. Environmental Health Perspectives. 2012; 120(4):501–7. https://doi.org/10.1289/ehp.1104170 PMID: 22182676; PubMed Central PMCID: PMC3339460.
- **84.** United Nations Environment Programme. Global Report on the Status of Legal Limits on Lead in Paint. 2016.
- 85. Leventhal T, Brooks-Gunn J. The neighborhoods they live in: The effects of neighborhood residence on child and adolescent outcomes. Psychological Bulletin. 2000; 126(2):309–37. https://doi.org/10.1037/0033-2909.126.2.309 PMID: 10748645
- Brito NH, Noble KG. Socioeconomic status and structural brain development. Frontiers in Neuroscience. 2014; 8:276. https://doi.org/10.3389/fnins.2014.00276 PMID: 25249931; PubMed Central PMCID: PMC4155174.

- 87. Buckley L, Broadley M, Cascio CN. Socio-economic status and the developing brain in adolescence: A systematic review. Child Neuropsychology. 2019; 25(7):859–84. https://doi.org/10.1080/09297049. 2018.1549209 PMID: 30466359
- Farah MJ. The neuroscience of socioeconomic status: Correlates, causes, and consequences. Neuron. 2017; 96:56–71. https://doi.org/10.1016/j.neuron.2017.08.034 PMID: 28957676
- Hackman DA, Farah MJ. Socioeconomic status and the developing brain. Trends in Cognitive Sciences. 2009; 13(2):65–73. https://doi.org/10.1016/j.tics.2008.11.003 PMID: 19135405
- 90. Gur RE, Moore TM, Rosen AFG, Barzilay R, Roalf DR, Calkins ME, et al. Burden of environmental adversity associated with psychopathology, maturation, and brain behavior parameters in youths. JAMA Psychiatry. 2019; 76(9):966–75. https://doi.org/10.1001/jamapsychiatry.2019.0943 PMID: 31141099
- Whittle S, Vijayakumar N, Simmons JG, Dennison M, Schwartz O, Pantelis C, et al. Role of positive parenting in the association between neighborhood social disadvantage and brain development across adolescence. JAMA Psychiatry. 2017; 74(8):824–32. https://doi.org/10.1001/jamapsychiatry. 2017.1558 PMID: 28636697
- 92. Hamilton JP, Siemer M, Gotlib IH. Amygdala volume in major depressive disorder: A meta-analysis of magnetic resonance imaging studies. Molecular Psychiatry. 2008; 13:993–1000. https://doi.org/10.1038/mp.2008.57 PMID: 18504424
- 93. Lanphear BP, Roghmann KJ. Pathways of lead exposure in urban children. Environmental Research. 1997; 74:67–73. https://doi.org/10.1006/enrs.1997.3726 PMID: 9339217.
- 94. Lanphear BP, Hornung R, Khoury J, Yolton K, Baghurst P, Bellinger DC, et al. Low-level environmental lead exposure and children's intellectual function: An international pooled analysis. Environmental Health Perspectives. 2005; 113(7):894–9. https://doi.org/10.1289/ehp.7688 PMID: 16002379; PubMed Central PMCID: PMC1257652.
- 95. Hornung RW, Lanphear BP, Dietrich KN. Age of greatest susceptibility to childhood lead exposure: A new statistical approach. Environmental Health Perspectives. 2009; 117(8):1309–12. https://doi.org/10.1289/ehp.0800426 PMID: 19672413; PubMed Central PMCID: PMC2721877.
- 96. Chen A, Dietrich KN, Ware JH, Radcliffe J, Rogan WJ. IQ and blood lead from 2 to 7 years of age: Are the effects in older children the residual of high blood lead concentration in 2-year-olds? Environmental Health Perspectives. 2005; 113:597–601. https://doi.org/10.1289/ehp.7625 PMID: 15866769
- 97. Menezes-Filho JA, Carvalho CF, Rodrigues JLG, Araújo CFS, dos Santos NR, Lima CS, et al. Environmental co-exposure to lead and manganese and intellectual deficit in school-aged children. International Journal of Environmental Research and Public Health. 2018; 15:2418. https://doi.org/10.3390/ijerph15112418 PMID: 30384464
- 98. Lucchini RG, Zoni S, Guazzetti S, Bontempi E, Micheletti S, Broberg K, et al. Inverse association of intellectual function with very low blood lead but not with manganese exposure in Italian adolescents. Environmental Research. 2012; 118:65–71. https://doi.org/10.1016/j.envres.2012.08.003 PMID: 22925625
- Cserbik D, Chen J-C, McConnell R, Berhane K, Sowell ER, Schwartz J, et al. Fine particulate matter exposure during childhood relates to hemispheric-specific differences in brain structure. Environment International. 2020; 143:105933. https://doi.org/10.1016/j.envint.2020.105933 PMID: 32659528.
- Fairchild AJ, McDaniel HL. Best (but oft-forgotten) practices: mediation analysis. American Journal of Clinical Nutrition. 2017; 105:1259–71. https://doi.org/10.3945/ajcn.117.152546 PMID: 28446497
- Soobader M, Cubbin C, Gee GC, Rosenbaum A, Laurenson J. Levels of analysis for the study of environmental health disparities. Environmental Research. 2006; 102:172–80. https://doi.org/10.1016/j.envres.2006.05.001 PMID: 16781704
- 102. Lanphear BP, Hornung R, Ho M. Screening housing to prevent lead toxicity in children. Public Health Reports. 2005; 120:305–10. https://doi.org/10.1177/003335490512000315 PMID: 16134573
- 103. Cantor AG, Hendrickson R, Blazina I, Griffin J, Grusing S, McDonagh MS. Screening for elevated blood lead levels in childhood and pregnancy: Updated evidence report and systematic review for the US Preventive Services Task Force. Journal of the American Medical Association. 2019; 321 (15):1510–26. https://doi.org/10.1001/jama.2019.1004 PMID: 30990555
- 104. Cory-Slechta DA. Studying toxicants as single chemicals: Does this strategy adequately identify neurotoxic risk? NeuroToxicology. 2005; 26(4):491–510. https://doi.org/10.1016/j.neuro.2004.12.007
 PMID: 16112317
- 105. Brulle RJ, Pellow DN. Environmental justice: Human health and environmental inequalities. Annual Review of Public Health. 2006; 27:103–24. https://doi.org/10.1146/annurev.publhealth.27.021405. 102124 PMID: 16533111.

- 106. Rauh VA, Margolis AE. Research Review: Environmental exposures, neurodevelopment, and child mental health—new paradigms for the study of brain and behavioral effects. The Journal of Child Psychology and Psychiatry. 2016; 57(7):775–93. https://doi.org/10.1111/jcpp.12537 PMID: 26987761; PubMed Central PMCID: PMC4914412.
- 107. Moody HA, Darden JT, Pigozzi BW. The relationship between neighborhood socioeconomic differences and racial residential segregation to childhood blood lead levels in metropolitan Detroit. Journal of Urban Health. 2016; 93(5):820–39. https://doi.org/10.1007/s11524-016-0071-8 PMID: 27538746; PubMed Central PMCID: PMC5052146.
- 108. Lidsky TI, Schneider JS. Lead neurotoxicity in children: basic mechanisms and clinical correlates. Brain. 2003; 126:5–19. https://doi.org/10.1093/brain/awg014 PMID: 12477693.
- **109.** Pellow DN. Environmental inequality formation: Toward a theory of environmental injustice. American Behavioral Scientist. 2000; 43(4):581–601.
- 110. Pirkle JL, Kaufmann RB, Brody DJ, Hickman T, Gunter EW, Paschal DC. Exposure of the U.S. population to lead, 1991–1994. Environmental Health Perspectives. 1998; 106(11):745–90. https://doi.org/10.1289/ehp.98106745 PMID: 9799191; PubMed Central PMCID: PMC1533471.
- Wheeler W, Brown MJ. Blood lead levels in children aged 1–5 years—United States, 1999–2010. Morbidity and Mortality Weekly Report. 2013; 62(13):245–8. PMID: 23552225; PubMed Central PMCID: PMC4605011.
- 112. Tsoi M-F, Cheung C-L, Cheung TT, Cheung BMY. Continual decreases in blood lead level in Americans: United States National Health Nutrition and Examination Survey 1999–2014. The American Journal of Medicine. 2016; 129:1213–8. https://doi.org/10.1016/j.amjmed.2016.05.042 PMID: 27341956.
- 113. Lively DE. The diminishing relevance of rights: Racial disparities in the distribution of lead exposure risks. Boston College Environmental Affairs Law Review. 1994; 21(2):309–34.
- 114. Lanphear BP, Weitzman M, Eberly S. Racial differences in urban children's environmental exposures to lead. American Journal of Public Health. 1996; 86(10):1460–3. https://doi.org/10.2105/ajph.86.10. 1460 PMID: 8876521; PubMed Central PMCID: PMC1380663.
- 115. Aelion CM, Davis HT, Lawson AB, Cai B, McDermott S. Associations between soil lead concentrations and populations by race/ethnicity and income-to-poverty ratio in urban and rural areas. Environmental Geochemistry and Health. 2013; 35(1):1–12. https://doi.org/10.1007/s10653-012-9472-0 PMID: 22752852; PubMed Central PMCID: PMC4655433.
- 116. Sargent JD, Brown MJ, Freeman JL, Bailey A, Goodman D, Freeman DH. Childhood lead poisoning in Massachusetts communities: Its association with sociodemographic and housing characteristics. American Journal of Public Health. 1995; 85:528–34. https://doi.org/10.2105/ajph.85.4.528 PMID: 7702117
- 117. Whitehead LS, Buchanan SD. Childhood lead poisoning: A perpetual environmental justice issue? Journal of Public Health Management & Practice. 2019; 25:S115–S20. https://doi.org/10.1097/PHH.0000000000000891 PMID: 30507780.
- 118. Fortner SK, Suffoletta MK, Vogt LK, Brown A, Diaz M. An iterative course-based soil lead research and partnerning model to address systemic racism and the enduring legacy of redlining. Environmental Justice. 2021. https://doi.org/10.1089/env.2021.0013
- Arora M, Austin C. Teeth as a biomarker of past chemical exposure. Current Opinion in Pediatrics. 2013; 25(2):261–7. https://doi.org/10.1097/MOP.0b013e32835e9084 PMID: 23429707.
- 120. Arora M, Kennedy BJ, Elhlou S, Pearson NJ, Walker DM, Bayl P, et al. Spatial distribution of lead in human primary teeth as a biomarker of pre- and neonatal lead exposure. Science of the Total Environment. 2006; 371:55–62. https://doi.org/10.1016/j.scitotenv.2006.07.035 PMID: 16950500.
- 121. Horton MK, Hsu L, Claus Henn B, Margolis A, Austin C, Svensson K, et al. Dentine biomarkers of prenatal and early childhood exposure to manganese, zinc and lead and childhood behavior. Environment International. 2018; 121:148–58. https://doi.org/10.1016/j.envint.2018.08.045 PMID: 30205321.