Neighborhood deprivation, prefrontal morphology and neurocognition in late childhood to early adolescence

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Abstract

\textbf{Background:} Neighborhood deprivation adversely effects neurodevelopment and cognitive function; however, mechanisms remain unexplored. Neighborhood deprivation could be particularly impactful in late childhood/early adolescence, in neural regions with protracted developmental trajectories, e.g., prefrontal cortex (PFC).

\textbf{Methods:} The Adolescent Brain Cognitive Development (ABCD) study recruited 10,205 youth. Geocoded residential history was used to extract individual neighborhood characteristics. A general cognitive ability index and MRI scans were completed. Associations with neurocognition were examined. The relation of PFC surface area and cortical thickness to neighborhood deprivation was tested. PFC subregions and asymmetry, with putative differential environmental susceptibility during key developmental periods, were explored. Analyses tested PFC area as a possible mediating mechanism.

\textbf{Results:} Neighborhood deprivation predicted neurocognitive performance ($\beta = -0.11$), even after accounting for parental education and household income ($\beta = -0.07$). Higher neighborhood deprivation related to greater overall PFC surface area ($\eta^2_p = 0.003$), and differences in leftward asymmetry were observed for area ($\eta^2_p = 0.001$), and thickness ($\eta^2_p = 0.003$). Subregion analyses highlighted differences among critical areas that are actively developing in late childhood/early adolescence and are essential to modulating high order cognitive function. These included orbitofrontal, superior frontal, rostral middle frontal, and frontal pole regions (Cohen’s $d = 0.03–0.09$). PFC surface area partially mediated the relation between neighborhood deprivation and neurocognition.

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CRediT authorship contribution statement

Teresa Vargas: Conceptualization, Methodology, Writing - original draft, Writing - review & editing. Katherine S.F. Damme: Data curation, Methodology, Writing - original draft, Writing - review & editing. Vijay A. Mittal: Supervision, Methodology, Writing - original draft, Writing - review & editing.

Declaration of competing interest
None.

Appendix A. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.neuroimage.2020.117086.
Discussion: Neighborhood deprivation related to cognitive function (a foundational skill tied to a range of lifetime outcomes) and PFC morphology, with evidence found for partial mediation of PFC on neurocognitive function. Results inform public health conceptualizations of development and environmental vulnerability.

Keywords

Neighborhood deprivation; Prefrontal cortex; Neurodevelopment; Environmental vulnerability; Cognition

1. Introduction

In recent years, seminal investigations have aimed to characterize phenomenologically distinct environmental influences on neurocognition during development (Farah et al., 2006; Laus et al., 2011; McLaughlin et al., 2014a; Noble et al., 2005; Pechtel and Pizzagalli, 2011; Sirin, 2005; Wu et al., 2015). Exposure to deprivation, defined as environments that do not provide needs or resources necessary for healthy development, has been identified as a critical environmental vulnerability factor (Cubbin and Winkleby, 2005; Jessop, 1992; Krishnadas et al., 2013; Matheson et al., 2008; McLaughlin et al., 2014b; Verhaeghe and Tampubolon, 2012). Animal and human studies on exposure to deprivation at the individual-level have yielded compelling evidence to its impact on both cognitive function and brain morphology (Akman et al., 2004; Beckett et al., 2006; Gee et al., 2013; Mackes et al., 2020; McLaughlin et al., 2014b; Mehta et al., 2009; Sheridan et al., 2012; Wiesel and Hubel, 1965). Largely these studies have examined individual cases of deprivation (e.g., parental neglect), but broad societal environments are also potential sources of deprivation (e.g., neighborhood poverty). Fewer studies, have broadened the scope of deprivation to considering structural, or systems level exposure (Wu et al., 2015). Local, neighborhood level, and even country-level structural characteristics (such as neighborhood socioeconomic status and national income inequality) have been consistently shown to impact individual health and development (Forsberg et al., 2018; Jaffe et al., 2005; Matheson et al., 2008; Tuliani et al., 2017). Yet, few investigations have sought to clarify whether brain morphology neural gray matter features can be impacted as a result of systems-level exposure to environmental vulnerability factors such as deprivation. Likewise, associations with general and specific functioning, such as cognitive performance, have yet to be integrated into these conceptualizations for key neurodevelopmental stages. The present investigation utilized a large, geographically diverse, nationally representative sample of youth to clarify possible relations between structural factors, cognitive and neural features. Identifying mechanisms through which neighborhood deprivation could be impactful is a necessary step toward identifying targets for epidemiological and public health models of healthy development.

Deprivation environments theoretically comprise settings in environmental resources are lacking (McLaughlin et al., 2014a); these can include scarcity of educational, cognitive, economic or health resources, which could lead to negative effects on neurodevelopment (Akman et al., 2004; Diamond et al., 1972; Globus et al., 1973). Animal and human research on deprivation environments has to date largely focused on individual-level exposure to
deprivation (i.e. individual socioeconomic status in human studies, experimental manipulations of enriched versus deprived environments in animal studies) (Akman et al., 2004; Diamond et al., 1972; Farah et al., 2006; Globus et al., 1973; Hackman et al., 2010; Lawson et al., 2017; Mackes et al., 2020; Mackey et al., 2015; Noble et al., 2005). Although these investigations have yielded valuable insights as to the possible effect of deprivation on neural function, deprivation can exist at a number of levels of a social structure. Exposure can range from individual-level deprivation (e.g. deprivation inside the immediate home), to deprivation of the broader environmental context (e.g. neighborhood features).

Research on the broader environmental and social context (i.e. structural characteristics) has long highlighted the necessity of taking into account systems-level factors potentially impacting healthy development (Bronfenbrenner, 1994; Glass and McAtee, 2006). This structural approach complements the valuable literature on individual-level factors (e.g., childhood institutionalization), generalizing the scope of what we consider to be influential to relatively distal factors (e.g., neighborhood resource scarcity). High deprivation environments occur at the systems level when various socioeconomic needs or resources are lacking, which may possibly hinder species-normative development. Existing literature exploring neighborhood deprivation shows evidence that it could be associated with a host of adverse health outcomes, including increased mortality rates (Forsberg et al., 2018; Jaffe et al., 2005; Matheson et al., 2008; Tuliani et al., 2017). Exposure during childhood could have particularly pervasive effects lasting throughout the lifetime (Mensah and Hobcraft, 2008). Neighborhood deprivation could be impactful on functional outcomes (such as neurocognitive function) and neural features, though these questions remain unexplored.

Lower neurocognitive function constitutes a foundational marker strongly related to a host of life outcomes (Aichele et al., 2016), and has a detrimental impact on health that is observable as early as adolescence (Sörberg et al., 2014). Research looking into individual-level deprivation exposure supports the notion that deprivation relates to lower cognitive function in the case of childhood institutionalization (Beckett et al., 2006) and socioeconomic status (Noble et al., 2005). Although evidence exists that neighborhood deprivation impacts cognition in older adults (Lang et al., 2008; McCann et al., 2018; Sheffield and Peek, 2009; Wu et al., 2015; Zeki Al Hazzouri et al., 2011), it has not been examined in earlier development, where individual-level deprivation has a demonstrated impact.

Exposure to deprived environments could affect neural regions with a protracted neurodevelopmental trajectory, such as the prefrontal cortex (PFC) (Mackes et al., 2020; McLaughlin et al., 2011, 2014a; Pechtel and Pizzagalli, 2011). PFC development undergoes a critical developmental milestone preceding puberty and adolescence, during which synaptic pruning, trophic glial and vascular changes, cell shrinkage and neuronal specialization begins to take place at a large scale (Gogtay et al., 2004; Tamnes et al., 2017). Prefrontal surface area and thickness, which follow unique developmental trajectories and thus may reflect differences in timing of impact, would be particularly informative (Lyall et al., 2015). Despite the fact, studies tying together neural mechanisms and neurocognition in the context of neighborhood deprivation are, to our knowledge, lacking in the literature. An association of neighborhood deprivation to PFC development and neurocognitive function...
could pinpoint larger community enrichment as a valuable strategy alongside targeted individual approaches to care at the population level.

The current study comprised a nationally representative sample of children aged 9–11 years old, an age range that has been established as a critical period of healthy PFC development (Gogtay et al., 2004; Tamnes et al., 2017). First, the study related neighborhood deprivation to neurocognition over and above proximal individual-level factors (such as parental education and household income). Then, the study examined neighborhood deprivation relations to PFC surface area and cortical thickness. Given their unique neurodevelopmental trajectories, surface area and cortical thickness were honed in on in order to gauge possibly distinct influences of neighborhood deprivation on PFC neurodevelopment (Raznahan et al., 2011; Shaw et al., 2008; Wierenga et al., 2014). The final, third aim explored whether PFC area would mediate an existing relation between neighborhood deprivation and neurocognitive function as a potential, contributing mechanism.

2. Materials and methods

The multisite Adolescent Brain Cognitive Development (ABCD) study aims to better understand adolescent development through a multimodal perspective (Volkow et al., 2018). The ABCD study utilized a school-based recruitment strategy, collecting cognitive and neuroimaging data from 9 to 11 year old children (Garavan et al., 2018). Written informed consent was obtained from participants, and data collection was approved by respective institutional review boards. Recruitment was conducted in a way that ensured the sample was representative of the U.S. population. The current study extracted data from the ABCD Release 2.0 (March 2019; DOI: 10.1515/506121). Data is available as part of the Adolescent Brain Cognitive Development (ABCD) study. Permission to access the data can be applied for at nda.gov.

2.1. Neighborhood deprivation

Residential history was established by collecting information on addresses where the participants had lived. Addresses were used to establish the Census tracts corresponding to each address. Publicly available Census data was then used to calculate the area deprivation index (ADI). The ADI was calculated based on the American Community Survey 2015 5-year summary, which has been successfully adapted in numerous investigations to assess deprivation at the neighborhood level (Kind et al., 2014). The ADI metric is compiled for each individual’s tract, representing Census-delineated neighborhoods of each participant. The area deprivation index has 17 subscores, including percentage of population aged ≥25 years with <9 years of education, percentage of population aged ≥25 years with at least a high school diploma, percentage of employed persons aged ≥16 years in white collar occupations, median family income, income disparity (log of 100 x ratio of the number of households with 50000 annual income), median home value, median gross rent, median monthly mortgage, percentage of home owners, percentage of occupied housing units with >1 person per room (crowding), percentage of civilian labor force population aged ≥16 years unemployed (unemployment rate), percentage of families below the poverty level, percentage of population below 138% of the poverty threshold, percentage of single parent
households with children aged <18 years, percentage of occupied housing units without a motor vehicle, percentage of occupied housing units without a telephone, and percentage of occupied housing units without complete plumbing. A composite ADI was computed for each participant based on the average of each address in which they had lived. Quintiles were then created based on ADI scores. ADI quintiles are predominantly used in the literature, their utility having been demonstrated across relevant socioeconomic domains (Knighton et al., 2016). Quintiles were numbered such that the first quintile has the lowest degree of deprivation, and the fifth quintile has the highest degree of deprivation (see Table 1).

2.2. Cognitive function

The NIH Toolbox Cognition domain comprises seven measures measuring the constructs of executive function, episodic memory, language, processing speed, working memory, and attention (Weintraub et al., 2013). NIH Toolbox instruments were validated in a sample of 476 participants ranging in age from 3 to 85 years, with representative sex, racial/ethnic categories, and education levels, and have been found to have appropriate test-retest reliability, as well as convergent and discriminant construct validity (Gershon et al., 2010, 2013; Weintraub et al., 2013). The current analyses utilized the NIH Toolbox composite score for Cognition. Scores were converted to T-scores derived from the NIH Toolbox nationally representative normative sample. Demographic variables were adjusted for in the corrected scores, including age, gender, race/ethnicity and educational attainment.

2.3. Structural imaging

Participants completed a high-resolution T1-weighted structural MRI scan (1-mm isotropic voxels) using scanners from GE Healthcare (Waukesha, Wisconsin), Philips Healthcare (Andover, Massachusetts), or Siemens Healthcare (Erlangen, Germany) (Casey et al., 2018). Structural MRI data was processed using FreeSurfer version 5.3.0 (http://surfer.nmr.mgh.harvard.edu/) (Dale et al., 1999; Fischl et al., 1999) according to standard processing pipelines (Casey et al., 2018). Processing included removal of nonbrain tissue, segmentation of gray and white matter structures (Fischl et al., 2002) and cortical parcellation (Fischl et al., 2004). All scan sessions underwent radiological review whereby scans with incidental findings were identified. Quality control for the structural images comprised visual inspection of T1 images and FreeSurfer outputs for quality (Hagler et al., 2019). Quality review was conducted by the ABCD team. Subjects whose scans failed inspection (due to severe artifacts or irregularities) were excluded. The Desikan-Killiany Atlas was used for cortical parcellation (Hagler et al., 2019). Regions of interest included caudal middle frontal, lateral orbitofrontal, medial orbitofrontal, rostral middle frontal, superior frontal, and frontal pole.

2.4. Parental education and household income

Data was collected on how many years of education the participant’s parent(s) had completed. In the case that the participant lived in a one-guardian household, the number of years for that parent was used. In the case that the participant lived in a two-guardian household, the number of years of education for each was averaged for both guardians. For household income, information was collected according to household income in the past 12
months (see Table 1). Of the original sample, 9375 subjects had data on both parental education and household income.

2.5. Data analysis

SPSS 25 was used for analyses. Chi-squares/One-way ANOVAS were used as appropriate to test for demographic differences among quintiles. The first aim was to determine whether neighborhood deprivation relates to cognitive function (using the composite NIH toolbox cognitive score). The second aim, in turn, was to determine whether neighbourhood deprivation would relate to PFC area and thickness, which exhibit distinct developmental trajectories and structural features (Wierenga et al., 2014). The third aim sought to determine whether the relationship with neurocognition, if present, would be mediated by PFC. For the first aim, a general linear model examined if composite cognitive scores would be predicted by composite ADI quintiles beyond the variance related to individual SES and parental education. To address the second aim, two repeated-measures ANCOVAs were run separately for deprivation quintiles (between-groups) predicting area and thickness (within groups, including subregions), respectively accounting for the variance related to age, sex, weight, parental education, household income, and a whole brain correction. Examining overall PFC with subregions as a repeated measure allows for detection of general, global PFC differences between groups (main effect of quintile) as well as for subregion-specific differences between groups (quintile by subregion interactions). For area analyses, variance related to total surface area was accounted for in the model. For thickness analyses, variance related to mean total thickness was accounted for in the model. Within-subject factors included hemisphere (left and right) as well as PFC regions (caudal middle frontal, lateral orbitofrontal, medial orbitofrontal, rostral middle frontal, superior frontal, and frontal pole). The model included tests for main effects, as well as interactions of quintiles by hemisphere, quintiles by region, and quintiles by hemisphere by region. In the case of interactions by quintile, Bonferroni-corrected, post-hoc tests were run to better understand relations between quintiles.

A mediation analysis was conducted using PROCESS v3.4 Model 4 (Hayes, 2017). In the model, age, sex, total surface area, parental education and household income were controlled for. Given the observed main effect from aim 1, PFC surface area was tested as a mediator. All analyses were re-run having identified outliers (Schwertman et al., 2004), these values were then Winsorized to the next non-outlier value. Given that this did not significantly alter observed findings, results are presented using the original data.

3. Results

3.1. Demographic characteristics and neurocognition

The current study comprised 10,205 participants. Quintiles did not differ by sex, $\chi^2$(4, 10204) = 5.33, $p = 0.26$. Significant differences in months of age $F(4, 10200) = 5.27$, $p < 0.001$ and weight $F(4, 10200) = 50.25$, $p < 0.001$ were detected between quintiles. As expected, significant differences were detected between quintiles with regards to parental education, $F(4,10043) = 436.04$, $p < 0.001$ and household income, $\chi^2(36, 9375) = 3008.68$, $p < 0.001$. The association of neighborhood deprivation and neurocognitive function was
significant, $\beta = -0.11$, $t = 10.62$, $p < 0.001$ (and persisted when accounting for parental education and household income, $\beta = -0.07$, $t = 5.60$, $p < 0.001$). Greater neighborhood deprivation related to lower neurocognitive performance (Fig. 1).

4. PFC surface area

Within a nested repeated-measures general linear model, within-subject factors were defined as frontal regions (6) nested within hemisphere (right and left), and between-subject factors were defined as quintiles, with the model accounting for variance related to total area, age, sex, weight, parental education, household income and site. As for between-subject effects, there was a main effect of quintiles on PFC surface area, $R(4,9262) = 6.29$, $p < 0.001$, $\eta^2_p = 0.003$. Quintile 5 showed greater total PFC area than Quintile 1 (mean difference = 22.52, SE = 4.72, $p < 0.001$, 95% CI 9.26–35.78) and Quintile 2 (mean difference = 16.18, SE = 4.56, $p = 0.004$, 95% CI 3.38–28.98). Quintile 3 also showed greater total PFC area than Quintile 1, mean difference = 13.48, SE = 4.19, $p = 0.01$, 95% CI 1.72–25.24. For within subjects effects, there was no main effect of hemisphere, Greenhouse-Geisser $R(1, 9262) = 2.64$, $p = 0.10$, though there was an effect of hemisphere by quintiles, Greenhouse-Geisser $R(4, 9262) = 2.47$, $p = 0.04$, $\eta^2_p = 0.001$. Across quintiles the left hemisphere exhibited greater surface area (Table 2). Quintile 1 (exhibiting the lowest levels of deprivation in the current samples) had greatest leftward asymmetry with confidence intervals that did not overlap with quintiles 2–5. As expected, there was a main effect of area, Greenhouse-Geisser $R(2.99, 27714.73) = 109.33$, $p > 0.001$, $\eta^2_p = 0.012$, and an area by quintiles interaction, Greenhouse-Geisser $R(11.97, 27714.73) = 3.70$, $p > 0.001$, $\eta^2_p = 0.002$. An interaction of hemisphere by area by quintiles was also observed, Greenhouse-Geisser $R(11.93, 27616.31) = 2.03$, $p = 0.02$, $\eta^2_p = 0.001$. For the frontal pole, greater neighborhood deprivation related to lower surface area. For right rostral middle frontal and superior frontal regions, greater neighborhood deprivation related to greater surface area (Figs. 2 and 3; Table 3).

4.1. PFC thickness

Within a nested repeated-measures general linear model, within-subject factors were defined as frontal regions nested within hemispheres, and between-subject factors were defined as quintiles, with the model accounting for variance related to total average thickness, age, sex, weight, parental education, household income and site. For between-subject effects, there was no main effect of quintiles on PFC thickness, $R(4,9262) = 1.07$, $p = 0.37$. For within-subject effects, there was a significant main effect of hemisphere exhibiting leftward asymmetry, Greenhouse-Geisser $R(1, 9262) = 4.53$, $p = 0.03$, $\eta^2_p = 0.001$. In addition, there was an interaction of hemisphere by quintiles, Greenhouse-Geisser $R(4, 9262) = 6.58$, $p < 0.001$, $\eta^2_p = 0.003$. Quintiles 3, 4 and 5 (exhibiting greater levels of deprivation in the current samples) had greater leftward asymmetry with confidence intervals that did not overlap with quintile 1, the quintile with the lowest deprivation levels. As expected, there was a main effect of thickness (Greenhouse-Geisser $R(2.66, 24672.84) = 334.06$, $p < 0.001$, $\eta^2_p = 0.035$) and an interaction of thickness by quintiles (Greenhouse-Geisser $R(10.66, 24672.84) = 2.40$, $p = 0.01$, $\eta^2_p = 0.001$). Finally, an interaction of hemisphere by region by quintiles was observed, Greenhouse-Geisser $R(8.69, 20114.09) = 3.71$, $p < 0.001$, $\eta^2_p = 0.002$. For the left lateral orbitofrontal, right medial orbitofrontal, right superior frontal and
right rostral middle frontal regions, greater neighborhood deprivation related to lower thickness (Table 3).

4.2. Mediation of PFC area

In line with the main effect observed in aim 1, the regression measuring the association of neighborhood deprivation and PFC area was significant, $\beta = 0.02$, $t = 4.84$, $p < 0.001$. Greater deprivation related to greater PFC area. While controlling for neighborhood deprivation, the relationship of the mediator (PFC area) with neurocognitive function was significant, $\beta = 0.09$, $t = 2.81$, $p = 0.005$. Greater PFC area predicted higher neurocognitive performance. Further, analyses showed that while controlling for the mediator (PFC area), neighborhood deprivation remained a significant predictor of neurocognitive function, $\beta = -0.06$, $t = -5.50$, $p < 0.001$. The Sobel test for indirect effects had a 95% confidence interval of $-0.0031$ to $-0.0004$, calculated using 5000 bootstrap samples. The model suggests partial mediation; 2.6% of the association between neighborhood deprivation and neurocognitive function is accounted for by PFC area ($\beta = 0.002$, $SE = 0.001$) (Fig. 4).

5. Discussion

Exposure to neighborhood level deprivation was explored in relation to cognitive function and protracted brain development (within PFC) in a nationally representative sample of youth. First, greater levels of neighborhood level deprivation predicted lower neurocognitive performance. Second, neighborhood deprivation related to distinct prefrontal gray matter features. Greater PFC area was observed with increased neighborhood deprivation levels. Additionally, neighborhood deprivation impacted the frontal asymmetry of gray matter area and thickness; the normative leftward bias was significantly different between the high and low deprivation groups. Further, the effect of neighborhood deprivation varied by PFC metric. Both surface area and thickness showed regional specificity; neighborhood deprivation particularly impacted critical regions including orbitofrontal, superior, rostral medial frontal, and frontal pole regions. To examine relevance of PFC area as a possible mechanism underlying cognitive deficits, associations between neighborhood deprivation and neurocognitive function were examined, with PFC area tested as a potential mediator. Analyses showed PFC surface area independently predicted neurocognitive performance; strikingly, the relation between neighborhood deprivation and neurocognition was found to be partially mediated by PFC surface area. Taken together, findings suggest that exposure to deprivation at the neighborhood level relates to both functional outcomes (i.e. neurocognition) and neurodevelopment (i.e. PFC). Results suggest that systems level environmental features are relevant and ought to be accounted for in models of environmental effects during critical periods of human neurodevelopment.

As hypothesized, increased levels of deprivation related to lower cognitive function. These results are consistent with, and of a similar effect size to those observed in older adults (50 years and older) (Drukker and van Os, 2003; Lang et al., 2008). The current study extends the existing literature on older adults (McCann et al., 2018; Sheffield and Peek, 2009; Wu et al., 2015; Zeki Al Hazzouri et al., 2011) to youth in late childhood to early adolescence undergoing critical normative neurodevelopmental processes. Results suggest that while the
cognitive developmental literature has largely focused on individual-level factors, more attention to structural factors such as neighborhood characteristics may be warranted. Structural level trends may be present even after accounting for the impact of individual-level features; future investigations will be necessary to determine whether systems level policy initiatives addressing structural characteristics such as neighborhood features (even in the absence of individual-targeted initiatives) could have a protective impact on neurocognitive development at the aggregate level.

The age of our current sample, directly preceding puberty and adolescence, is critical for PFC development, with synaptic pruning, cell shrinkage, neuronal specialization, trophic glial and vascular changes starting to take place at a broad scale (Gogtay et al., 2004; Pechtel and Pizzagalli, 2011; Tamnes et al., 2017). These processes make the PFC particularly susceptible to environmental influence at this stage of neurodevelopment. In the present study, greater levels of deprivation predicted increased PFC surface area. Results support the broader literature on individual-level early life stress and effects on neural development (Gee et al., 2013; McLaughlin et al., 2014a, 2014b; Tottenham et al., 2010; Tottenham and Sheridan, 2010). The fact that an effect was not observed with regards to total PFC cortical thickness could mean the effect at this age is specific to surface area. Surface area and cortical thickness follow distinct developmental trajectories throughout childhood and early adolescence (Alemán-Gómez et al., 2013; Raznahan et al., 2011; Shaw et al., 2008; Tamnes et al., 2017). For example, while both cortical thickness and surface area show normative decreases throughout late childhood and adolescence, cortical thinning happens at a much greater degree compared to surface area (Tamnes et al., 2017). The observed difference relating to deprivation could be due to a delayed trajectory of synaptic pruning and dendritic arborisation (Alemán-Gómez et al., 2013; Bourgeois and Rakic, 1993; Huttenlocher and Dehghani, 1997; Klein et al., 2014; Petanjek et al., 2011; Raznahan et al., 2011; Tamnes et al., 2017; White et al., 2010).

For surface area, decreased leftward asymmetry was observed as deprivation levels increased. Interestingly, for cortical thickness an opposing pattern was observed—increasing leftward asymmetry as deprivation levels increased. Perhaps divergences in leftward asymmetry at this age relate to altered developmental processes due to exposure to adverse environmental factors (Lawson et al., 2013; Luders et al., 2005); indeed, there is evidence that individual differences in cerebral lateralization may be vastly influenced by environmental factors occurring during early neurodevelopment (Bishop, 2013; Raj and van Oudenaarden, 2008). Further, asymmetry has been associated with impactful outcomes including psychopathology risk and cognitive function (Avnit et al., 2019; Damme et al., 2020; Keune et al., 2015). Future studies will be needed to further enrich and corroborate these hypotheses and incidental findings.

Subregion specific analyses yielded compelling results. Superior frontal and rostral middle frontal areas were larger in quintiles with greater neighborhood deprivation. By contrast, more ventral prefrontal areas including orbitofrontal and frontal pole regions showed lower area/thickness in quintiles with greater deprivation. Results are partially consistent with investigations on individual institutionalization conferring childhood deprivation (Mackes et al., 2020). Given these region’s involvement in fundamental cognitive functions (Bahlmann
et al., 2015; Lopatina et al., 2017), and the lack of precedent in the literature, future studies will benefit from further assessing relations to environmental influence. On the contrary, bilateral frontal pole surface area decreased as neighborhood deprivation exposure increased; this could be due to it being among the first prefrontal regions to fully develop (Gogtay et al., 2004). Perhaps the region is more susceptible to early deprivation, with less compensatory mechanisms or developmental flexibility occurring; these processes could be functionally relevant given frontal pole relations to complex cognition (Burke et al., 2013; Gilbert et al., 2006; Liu et al., 2013; Semendeferi et al., 2001).

Decreases in left lateral orbitofrontal, right medial orbitofrontal, right superior frontal, and right rostral middle frontal regions related to greater neighborhood deprivation. Orbitofrontal thickness results are consistent with previous investigations on children exposed to maltreatment (Kelly et al., 2013) and institutionalization (Mackes et al., 2020; Sheridan et al., 2012). Subregion analyses are also partially consistent with an investigation in children exposed to maltreatment (Kelly et al., 2013) and another examining relations between children’s socioeconomic status and prefrontal cortical thickness (Lawson et al., 2013). Future studies will be needed to further flesh out relations between neighborhood deprivation and prefrontal cortical subregions.

Finally, analyses tested PFC area as a relevant mechanism. The model found evidence for PFC area partially mediating the relation between neighborhood deprivation and neurocognitive function. Results suggest that the association between distal exposure to deprivation and neurocognitive function becomes apparent at an early age, and is robust to related individual-level factors (i.e. household income and parental education). The proportion of the effect that PFC area explained was modest and is important to note; nonetheless, given the milieu of environmental, genetic, and neurodevelopmental factors that may each uniquely impact and be impacted by neighborhood deprivation, this is to be reasonably expected. PFC area is one of many candidate neurodevelopmental processes through which neighborhood deprivation could impact neurocognition. The model showed increased PFC area related to greater neurocognitive function in the current sample. Perhaps increased PFC surface area serves as a developed protective mechanism attenuating the established relation between increased deprivation and decreased neurocognitive function. Given these are novel questions, future investigations are needed to further build on these initial findings.

It is critical to highlight that effect sizes were rather modest for neighborhood deprivation predicting PFC brain metrics and neurocognition. These effect sizes are in line with studies of distal neighborhood-level characteristics predicting biological outcomes (Laraia et al., 2012; Lopez, 2007), as well as gray matter MRI pooled samples research (Sacher et al., 2012), which typically observe small effect sizes. Further, these findings were robust to accounting for more proximal characteristics relating to deprivation (parental education and household income), which are typically impactful to a greater magnitude. Although observed effects are small according to convention, it will be paramount for future investigations to ascertain clinical significance and possible relevance to public health initiatives. There is reason to think this pursuit could be fruitful. Indeed, public health and medicine research has successfully based treatment decisions on comparable effect sizes that
would conventionally be viewed as small. For example, low-dose aspirin remains recommended to reduce the risk of heart attacks for those under 70 years old without bleeding risk, based on a 0.03 correlation (Steering Committee of the Physicians’ Health Study Research Group, 1988; Steering Committee of the Physicians’ Health Study Research Group, 1989), and fruit and vegetable consumption is recommended to address weight and abdominal obesity based on an effect size of −0.05 (Schwingshackl et al., 2015). However, it is also noteworthy that while the initial policy impact of the aspirin studies was initially broader, it was eventually scaled back to include a more circumscribed set of circumstances (Arnett, 2019). Thus, while there is good reason to consider the present effects as an important step in clarifying our understanding in this critical area, it should not be forgotten that the effects were small. Future investigations would benefit from clarifying and identifying whether there are practical consequences to the presently observed effects, which will aid with interpreting and understanding their magnitude (Funder; Ozer, 2019). Perhaps targeting these individually small effects at the societal level through policy initiatives for neighborhood features has potential to provide public health improvements at the aggregate level—future study will be critical in determining these possibilities.

Overall, this study’s observations suggest that more attention to environmental characteristics may be warranted in public health and neurodevelopmental models; a general approach to addressing deprivation (including both individual-level and neighborhood/systems-level exposure) could hold widespread benefits to communities. Although the current study provides promising introductory evidence, there are several lines of inquiry that remain unanswered. First, it will be necessary to tease apart exposure during several different developmental periods, in order to confirm critical periods of impact for systems level exposures such as neighborhood deprivation. The timing of the most critical developmental periods may vary within-subjects, and so it will be essential to allow for multiple time points across the lifespan. In addition, gauging more detailed information on different environments that individuals are exposed to beyond their home environment could provide valuable information for developing interventions. Further fleshing out the intricate interactions between proximal exposure to environmental adversity and distal exposure at the systems level is also a much-needed area of inquiry. In all, results suggest that larger community enrichment could be a valuable tool alongside targeted individual approaches to care, and follow up investigations exploring this notion would be valuable for treatment and intervention.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Association between neurocognitive performance and neighborhood deprivation quintiles. Increasing levels of neighborhood deprivation relate to lower neurocognitive performance, $p < 0.001$. Neurocognition values are adjusted for age, sex, parental education, and household income.
Fig. 2.
Regions (for area and thickness) associated with neighborhood deprivation in Bonferroni-corrected analyses ($p_{\text{Bonferroni}} < 0.05$). Analyses adjust for age, sex, a whole brain correction, weight, parental education, and household income.
Fig. 3.
Bilateral frontal pole surface area plotted according to individual Area Deprivation Index (ADI) values used to sort participants into quintiles (color-coded to the right). The highest deprivation quintile (Q5) shows smaller surface area relative to Q2 and Q3. $p_{Bonferroni} < 0.05$. Adjusted area values constitute estimated marginal means partialling out age, sex, a whole brain correction, weight, parental education, and household income.
Fig. 4.
Association between neighborhood deprivation and neurocognitive function, with prefrontal area as a partial mediator. Greater neighborhood deprivation predicts lower neurocognitive performance. Independently, greater prefrontal surface area predicts higher neurocognitive performance. The association between neighborhood deprivation and neurocognitive performance is partially mediated by prefrontal surface area.
Table 1

Demographic characteristics.

| Group Comparison | Q1 n | % | Q2 n | % | Q3 n | % | Q4 n | % | Q5 n | % |
|------------------|------|---|------|---|------|---|------|---|------|---|
| Sex (Female)     | 977  | 47.80 | 935  | 45.90 | 956  | 46.70 | 996  | 49.00 | 990  | 48.50 |
| Household Income |      |      |      |      |      |      |      |      |      |      |
| Less than $5000  | 24   | 1.3 | 32   | 1.7 | 47   | 2.5 | 61   | 3.3 | 170  | 9.5 |
| $5000–11999      | 19   | 1   | 30   | 1.6 | 49   | 2.6 | 83   | 4.4 | 172  | 9.6 |
| $12000–15999     | 20   | 1.1 | 19   | 1   | 30   | 1.6 | 66   | 3.5 | 101  | 5.7 |
| $16000–24999     | 26   | 1.4 | 42   | 2.2 | 67   | 3.5 | 89   | 4.8 | 203  | 11.4 |
| $25000–34999     | 43   | 2.3 | 67   | 3.5 | 78   | 4.1 | 140  | 7.5 | 234  | 13.1 |
| $35000–49999     | 72   | 3.8 | 95   | 4.9 | 143  | 7.5 | 231  | 12.4 | 249  | 13.9 |
| $50000–74999     | 144  | 7.6 | 196  | 10.2 | 310  | 16.3 | 348  | 18.6 | 311  | 17.4 |
| $75000–99999     | 186  | 9.8 | 315  | 16.4 | 354  | 18.6 | 342  | 18.3 | 180  | 10.1 |
| $100000–199999   | 750  | 39.6 | 856  | 44.5 | 690  | 36.2 | 454  | 24.3 | 162  | 9.1 |
| $200000+         | 608  | 32.1 | 271  | 14.1 | 136  | 7.1 | 55   | 2.9 | 5    | 0.3 |

| Group Comparison | Q1 Mean | SD | Q2 Mean | SD | Q3 Mean | SD | Q4 Mean | SD | Q5 Mean | SD |
|------------------|---------|---|---------|---|---------|---|---------|---|---------|---|
| Age (months)     | 119.34  | 7.49 | 119.44  | 7.44 | 118.88  | 7.58 | 119.06  | 7.53 | 118.49  | 7.41 |
| Weight (inches)  | 80.42   | 21.12 | 81.65  | 22.11 | 83.49  | 23.18 | 86.09  | 25.81 | 90.08  | 29.61 |
| Cognition total score\(a\) | 46.76   | 16.86 | 46.19  | 15.88 | 45.08  | 16.36 | 44    | 15.46 | 41.82  | 15.24 |
| Parental education | 17.71  | 2.17 | 17.23  | 2.32 | 16.57  | 2.6 | 16    | 2.54 | 14.79  | 2.52 |
| ADI index score  | 55.22   | 27.05 | 89.52  | 3.5 | 98.4   | 2.15 | 105.83  | 2.31 | 115.63  | 3.72 |

\(\chi^2(4,10204) = 5.33, p = 0.026^*\)
\(\chi^2(36,9375) = 3008.68, p < 0.001^*\)
\(F(4,10200) = 5.27, p < 0.001^*\)
\(F(4,10200) = 50.25, p < 0.001^*\)
\(\beta=0.11, t=10.62, p < 0.001^*\)
\(F(4,10043) = 436.04, p < 0.001^*\)
\(F(4,10200) = 7120.82, p < 0.001^*\)
Table 2

Bonferroni-corrected significant mean differences between quintiles of leftward asymmetry for frontal area and thickness.

| Surface area | Quintile | Cohen’s $d$ | Mean difference | SE  | $P_{Bonferroni}$ | 95% CI     |
|--------------|----------|-------------|-----------------|-----|-----------------|------------|
|              | 5        | 0.069       | 23.123          | 3.504 | <0.001          | 16.254–29.992 |
|              | 4        | 0.075       | 22.795          | 3.153 | <0.001          | 16.614–28.977 |
|              | 3        | 0.073       | 21.834          | 3.106 | <0.001          | 15.746–27.922 |
|              | 2        | 0.088       | 26.584          | 3.138 | <0.001          | 20.433–32.734 |
|              | 1        | 0.110       | 34.515          | 3.250 | <0.001          | 28.145–40.885 |

| Thickness    | Quintile | Mean difference | Mean difference | SE  | $P_{Bonferroni}$ | 95% CI     |
|--------------|----------|-----------------|-----------------|-----|-----------------|------------|
|              | 5        | 0.109           | 0.021           | 0.002 | <0.001         | 0.016–0.025 |
|              | 4        | 0.119           | 0.023           | 0.002 | <0.001         | 0.019–0.027 |
|              | 3        | 0.088           | 0.017           | 0.002 | <0.001         | 0.013–0.021 |
|              | 2        | 0.062           | 0.012           | 0.002 | <0.001         | 0.008–0.016 |
|              | 1        | 0.047           | 0.009           | 0.002 | <0.001         | 0.005–0.013 |
Table 3

Bonferroni-corrected significant associations between quintiles and prefrontal subregion frontal area/thickness.

| Surface area PFC region | Hemisphere | Summary | Cohen’s $d$ | Mean difference | SE | $p_{Bonferroni}$ | 95% CI       |
|-------------------------|------------|---------|-------------|-----------------|----|-----------------|-------------|
| Caudal Middle Frontal   | –          | no difference |             |                 |    |                 |             |
| Lateral Orbitofrontal   | Left       | 5 < 3 = 2 > 1; 5 < 2 | [5 < 3] $-0.031$; [5 < 2] $-0.031$ | [5 < 3] $-22.369$; [5 < 2] $-23.100$ | 22.369; 23.100 | $7.457$; $7.668$ | $0.027$; $0.026$ | $-43.307$ to $-1.432$; $-44.629$ to $-1.571$ |
|                         |            |         |             |                 |    |                 |             |
|                         | Right      | 5 = 4 < 2 = 1 | [5 < 2] $-0.031$; [5 < 2] $-0.031$ | [5 < 2] $-23.773$; [5 < 2] $-23.773$ | 23.773; 23.773 | $8.022$; $8.022$ | $0.031$; $0.031$ | $-46.296$ to $-1.249$; $-46.296$ to $-1.249$ |
| Medial Orbitofrontal    | –          | no difference |             |                 |    |                 |             |
|                         | Left       | 5 < 4 < 3 = 2 = 1 | [5 < 3] $0.040$; [5 < 2] $0.045$ | [5 < 3] $55.909$; [5 < 2] $85.697$ | 55.909; 85.697 | $18.770$; $19.808$ | $0.029$; $0.020$ | $30.082$ to $141.311$; $19.263$ to $30.082$ |
|                         | Right      | 5 < 3 = 2 = 1 | [5 < 2] $-0.031$; [5 < 2] $-0.031$ | [5 < 2] $-22.064$; [5 < 2] $-22.064$ | 22.064; 22.064 | $7.408$; $7.408$ | $0.029$; $0.029$ | $-42.863$ to $-1.265$; $-42.863$ to $-1.265$ |
| Superior Frontal        | Right      | 5 < 3 = 2 > 1; 5 > 2 | [5 < 3] $-0.031$; [5 < 2] $-0.031$ | [5 < 3] $-20.740$; [5 < 2] $-20.740$ | 20.740; 20.740 | $7.668$; $6.960$ | $0.027$; $0.026$ | $-42.100$ to $-1.022$; $-42.100$ to $-1.022$ |
|                         | Right      | 5 < 3 = 2 > 1; 5 > 2 | [5 < 3] $-0.031$; [5 < 2] $-0.031$ | [5 < 3] $-21.470$; [5 < 2] $-21.470$ | 21.470; 21.470 | $6.960$; $6.960$ | $0.026$; $0.020$ | $-41.012$ to $-1.928$; $-41.012$ to $-1.928$ |
| Frontal Pole            | Left       | 5 < 3 = 2 > 1; 5 < 2 | [5 < 3] $-0.037$; [5 < 2] $-0.045$ | [5 < 3] $-1.728$; [5 < 2] $-1.728$ | 1.728; 1.728 | $-2.059$; $-2.059$ | $0.003$; $0.002$ | $-4.857$ to $0.072$; $-4.857$ to $0.072$ |
|                         | Right      | 5 < 3 = 2 > 1; 5 < 2 | [5 < 3] $-0.036$; [5 < 2] $-0.036$ | [5 < 3] $-5.405$; [5 < 2] $-5.405$ | 5.405; 5.405 | $-1.547$; $-1.547$ | $0.005$; $0.005$ | $-9.749$ to $-9.749$; $-9.749$ to $-9.749$ |
| Thickness PFC region    | Hemisphere | Summary | Mean difference | SE | $p_{Bonferroni}$ | 95% CI       |
| Caudal Middle Frontal   | –          | no difference |             |                 |    |                 |             |
| Lateral Orbitofrontal   | Left       | 5 < 3 | [5 < 3] $-0.036$; [5 < 3] $-0.036$ | [5 < 3] $-0.036$; [5 < 3] $-0.036$ | $-0.036$; $-0.036$ | $-0.014$; $-0.014$ | $0.004$; $0.004$ | $-0.026$ to $-0.026$; $-0.026$ to $-0.026$ |
| Medial Orbitofrontal    | Right      | 5 < 3 = 2; 4 < 2 | [5 < 3] $-0.035$; [5 < 2] $-0.050$ | [5 < 3] $-0.035$; [5 < 2] $-0.050$ | $-0.035$; $-0.050$ | $-0.020$; $-0.020$ | $0.006$; $0.006$ | $-0.036$ to $-0.036$; $-0.036$ to $-0.036$ |
|                         | Right      | 5 < 4 < 3 = 2 = 1; 4 < 2 | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | $-0.039$; $-0.039$ | $-0.018$; $-0.018$ | $0.005$; $0.005$ | $-0.034$ to $-0.034$; $-0.034$ to $-0.034$ |
| Rostral Middle Frontal  | Right      | 5 < 4 < 3 = 2 = 1; 4 < 2 | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | $-0.039$; $-0.039$ | $-0.019$; $-0.019$ | $0.005$; $0.005$ | $-0.034$ to $-0.034$; $-0.034$ to $-0.034$ |
|                         | Right      | 5 < 4 < 3 = 2 = 1; 4 < 2 | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | $-0.039$; $-0.039$ | $-0.019$; $-0.019$ | $0.005$; $0.005$ | $-0.034$ to $-0.034$; $-0.034$ to $-0.034$ |
|                         | Right      | 5 < 4 < 3 = 2 = 1; 4 < 2 | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | $-0.039$; $-0.039$ | $-0.019$; $-0.019$ | $0.005$; $0.005$ | $-0.034$ to $-0.034$; $-0.034$ to $-0.034$ |
|                         | Right      | 5 < 4 < 3 = 2 = 1; 4 < 2 | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | [5 < 4] $-0.039$; [5 < 2] $-0.039$ | $-0.039$; $-0.039$ | $-0.019$; $-0.019$ | $0.005$; $0.005$ | $-0.034$ to $-0.034$; $-0.034$ to $-0.034$ |
| Surface area PFC region | Hemisphere | Summary                | Cohen’s $d$     | Mean difference | SE      | $p_{Bonferroni}$ | 95% CI            |
|-------------------------|------------|------------------------|----------------|-----------------|--------|-----------------|------------------|
| **Superior Frontal**    | Right      | 5 < 2 = 1; 4 < 2 = 1  | $[4 < 3] -0.031$; | $[4 < 3] -0.009$; | $[4 < 3] 0.003$; | $[4 < 3] 0.037$; | $[4 < 3] -0.018$ - 0.000; |
|                         |            |                        | $[4 < 2] -0.052$; | $[4 < 2] -0.015$; | $[4 < 2] 0.003$; | $[4 < 2] 0.000$; | $[4 < 2] -0.024$ to −0.006; |
|                         |            | 4 < 1 −0.038;          | $[4 < 1] 0.003$; | $[4 < 1] 0.008$; | $[4 < 1] -0.020$ to −0.002; |
| **Frontal Pole**        | no difference |                        | −                | −               | −      | −               | −                |