ABSTRACT
BACKGROUND: Early low socioeconomic status (SES) is associated with poor outcomes in childhood, many of which endure into adulthood. It is critical to determine how early low SES relates to trajectories of brain development and whether these mediate relationships to poor outcomes. We use data from a unique 17-year longitudinal study with five waves of structural brain imaging to prospectively examine relationships between preschool SES and cognitive, social, academic, and psychiatric outcomes in early adulthood.

METHODS: Children (n = 216, 50% female, 47.2% non-White) were recruited from a study of early onset depression and followed approximately annually. Family income-to-needs ratios (SES) were assessed when children were ages 3 to 5 years. Volumes of cortical gray and white matter and subcortical gray matter collected across five scan waves were processed using the FreeSurfer Longitudinal pipeline. When youth were ages 16+ years, cognitive function was assessed using the NIH Toolbox, and psychiatric diagnoses, high-risk behaviors, educational function, and social function were assessed using clinician administered and parent/youth report measures.

RESULTS: Lower preschool SES related to worse cognitive, high-risk, educational, and social outcomes (|standardized B| = 0.20–0.31, p values < .003). Lower SES was associated with overall lower cortical (standardized B = 0.12, p < .0001) and subcortical gray matter (standardized B = 0.17, p < .0001) volumes, as well as a shallower slope of subcortical gray matter growth over time (standardized B = 0.04, p = .012). Subcortical gray matter mediated the relationship of preschool SES to cognition and high-risk behaviors.

CONCLUSIONS: These novel longitudinal data underscore the key role of brain development in understanding the long-lasting relations of early low SES to outcomes in children.

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not address whether preschool SES relates to the trajectories of brain development within an individual, a question with high relevance for evaluating the timing and duration of SES relationships. However, Hanson et al. found that children from poor families had slower trajectories of growth in whole-brain volume and in regional frontal and parietal lobe volumes, with these effects related to the emergence of externalizing symptoms. Importantly, the patterns of growth suggested few differences as a function of poverty at the very earliest ages (e.g., 5–9 months old) but increasing differences as children grew older (26). Such findings are consistent with the idea that continued exposure to impoverished environments slows brain development. However, not all studies have found similar results (27–30). This literature is constrained by few studies having multiple waves of brain imaging, with only the Hanson study (26) to our knowledge having more than three time points. There is also some evidence for such mediation in the domains of mental health (31–34) and cognition (1,35–37). This has led to call for more studies that assess the same in- dividuals longitudinally (1) to address such critical questions about whether brain structure mediates the relationship of preschool SES to a broader array of adult outcomes. Such investigations have important developmental implications in understanding how early exposures may set trajectories for long-term outcomes.

To address this major gap in our understanding of the pathways from early SES to poor adaptive outcomes as youth transition to adulthood, we analyzed data from the Preschool Depression Study, a 17-year longitudinal study that followed children from preschool through the transition to adulthood, with up to five waves of imaging in each youth. We asked 1) if income-to-needs in preschool related to cognitive function, high-risk behaviors, social function, educational function, and/or psychiatric outcomes when youth were between the ages of 16 and 21; 2) if income-to-needs in preschool related to trajectories of growth in cortical and subcortical gray matter, white matter, or specific cortical and subcortical regions; and 3) whether variation in structural brain development mediated the relationship between preschool SES and outcomes at the transition to adulthood.

METHODS AND MATERIALS

Participants

The Preschool Depression Study is a 17-year longitudinal study that includes five waves of brain scans across school age to early adulthood (Figure S1). At the time of their first interview (T1), 306 children aged 3 to 5 years and their primary caregivers were recruited from the St. Louis area, using a checklist to oversample preschoolers with elevated symptoms of depression (38). At school age (–7–12 years), healthy children and those with a history of depression and/or anxiety were invited to participate in brain imaging; an additional 42 healthy children were also recruited (n = 210 completed the first wave of imaging). See the Supplement for exclusions. All methods were approved by the Institutional Review Board at Washington University (IRB #201502094). Written informed consent and assent was obtained from all participants.

Preschool SES

SES was operationalized as the income-to-needs ratio, defined as the total family income at T1 (Figure S1) divided by the federal poverty level based on family size (39).

Preschool Psychopathology and Life Events

Trained staff from the Early Emotional Development Program conducted up to 10 in-person assessment sessions with participants and their primary caregivers over the course of the study (Figure S1). The children were between the ages of 3 years and 5 years 11 months at T1 and between the ages of 15 years 3 months and 21 years 6 months at the most recent assessment wave (T10/MRI 5). The Preschool-Age Psychiatric Assessment (PAPA) (40,41) was the diagnostic assessment when children were age 3 years to 7 years 11 months. The PAPA is designed for diagnostic use with the caregivers of children ages 2 to 6 years (but has been used up to age 8 years) and has acceptable reliability (41). It consists of questions about developmentally appropriate symptom manifestations of DSM-IV criteria for all Axis I disorders, including major depression disorder, attention-deficit/hyperactivity disorder, and anxiety disorders. We created the following dimensional T1 psychopathology scores by computing the number of core items from the PAPA endorsed by the parents (Figure S1) to estimate early psychiatric challenges in these children: 1) depression (major depressive disorder); 2) anxiety (separation anxiety disorder, generalized anxiety disorder, and post-traumatic stress disorder); and 3) externalizing (oppositional defiant disorder, attention-deficit/hyperactivity disorder, and conduct disorder). Each of these was significantly correlated with T1 income-to-needs (r values = –0.20 to –0.33, p values = .011–.0001), and all were included as covariates in follow-up analyses to determine whether any relationships of SES to later outcomes were over and above early psychopa-thology. Youth’s life events were assessed at each annual assessment wave starting at T1 as part of the PAPA and then the Childhood-Age Psychiatric Assessment (see the Supplement for description of all possible events) (42). Life events were measured as the total number of events through the first scan wave (Figure S1).

Maternal Mental Health

Maternal history of mental illness (number of diagnoses) was assessed using the Family Interview for Genetic Studies (43). This measure was used as a covariate in analyses presented below to determine the degree to which SES associated with adaptive outcomes and brain measures over and above maternal mental health. We focused on maternal mental health because the vast majority of our participating parents were mothers, and we felt most confident in maternal mental health reports. See the Supplement for details.

Late Adolescence/Early Adulthood Adaptive Outcomes

We modeled our outcome metrics after those used in the Great Smoky Mountains Study (44–46) but did not include health, because it was not well assessed in our study, and added cognitive function. We used data from T9/MRI 4 and T10/MRI 5 (Figure S1), but only when the participant was age 16 years or
older. We chose 16+ years as a good balance of capturing late adolescence/early adulthood without having too narrow of an age range for outcomes. We used T9 and T10 because age varied, and some youth were 16+ years at both T9 and T10, while some youth were not 16+ years until T10. The measure of each of these outcomes is described in detail in the Supplement, with a brief overview provided below.

**Cognitive Function.** Participants completed the following NIH Toolbox (HealthMeasures) cognitive measures (47): 1) picture sequence memory, episodic memory; 2) list sorting, working memory; 3) flanker, selective attention; 4) pattern comparison, processing speed; and 5) picture vocabulary, verbal IQ (see the Supplement). We used age-corrected t scores and averaged the five tasks to create a composite.

**High-Risk Behavioral Outcomes.** We coded for the following eight behaviors, each coded “1” for present at either T9 or T10 or “0” for absent at both (see the Supplement): 1) pathological lying; 2) initiating physical fights; 3) breaking and entering; 4) arrested; detained; cited; adjudicated; placed on probation, juvenile detention, or court-ordered treatment; or incarcerated since last assessment; 5) being intoxicated or drunk at least 1 or 2 times per week in the past year; 6) marijuana or other illicit drug use in the past year; 7) consensual sex with someone known for <24 hours; and 8) being pregnant (female) or impregnating someone (male) prior to age 18 years (not the result of rape).

**Social Outcomes.** We coded for the following seven outcomes (Supplement): 1) poor peer relationships; 2) peer acceptance/rejection; 3) being bullied; 4) relational victimization; 5) social withdrawal; 6) social inhibition; and 7) prosocial behavior.

**Education Outcomes.** We coded for two outcomes (Supplement): 1) poor school engagement and 2) poor academic function.

**Psychiatric Outcomes.** We coded for the following seven outcomes (Supplement): 1) anxiety disorder; 2) a unipolar depressive disorder; 3) conduct disorder; 4) alcohol, 5) marijuana, or 6) substance use disorder; and 7) significant borderline personality symptoms. We also conducted secondary analyses with a variable that had only six outcomes, excluding borderline personality symptoms. We also examined each of the six disorder categories individually.

The final outcome score for the high-risk, social, education, and psychiatric outcomes was the mean of the items within each domain.

**Structural Imaging Acquisition**

All five waves of MRI data collection acquired 3D T1-weighted scans using a 3.0T whole-body scanner (Trio or Prisma) (Siemens Healthineers). See the Supplement for details.

**Structural Imaging Processing**

The data from MRI waves 1–3 have been previously processed (48). Processing of structural data used the FreeSurfer Longitudinal processing stream (version 5.3) (http://surfer.nmr.mgh.harvard.edu) (49). See the Supplement for details of processing, quality control, and harmonization across platforms. Volume of cortical gray, cortical white, subcortical gray, hippocampus, amygdala, caudate, putamen, and thalamus were obtained using FreeSurfer’s “aseg.stats” report, as was intracranial volume. Volumes of the dorsolateral prefrontal cortex and dorsal anterior cingulate were obtained from the Destrieux Atlas (50). We did not have hypotheses about asymmetries and thus averaged the left and right. We examined global measures (e.g., cortical gray, cortical white, subcortical gray) and regional measures because of the possibility that SES relations to brain development reflect mechanisms that could have broad effects across the cortex or subcortex or both (e.g., toxins, nutrition, stress).

**Statistical Analysis**

Continuous predictors and outcomes were standardized to facilitate comparison of effect sizes (standardized B; see the Supplement). Covariates for all analyses included sex, and analyses of brain variables included intracranial volume. Significant results were followed up by analyses adding T1 psychopathology (depression, anxiety, and externalizing), life events (see the Supplement), and maternal mental health as covariates (Supplement). Sex did not interact significantly with T1 income-to-needs ratio (T1INR) or any brain variable to relate to any outcome, and thus sex interactions were not included in final models. Because some of the outcome variables were counts, the results of the linear regressions were confirmed using zero-inflated Poisson regression.

Multiple regression was used to examine relationships of early T1INR to each outcome, with bootstrapped confidence intervals (1000) and false discovery rate (FDR) correction (51) across the five outcome measures. For the individual psychiatric disorder outcomes, we conducted logistic regressions. To examine whether T1INR relates to trajectories of brain volume across MRI 1–MRI 5, we used multilevel models that included both random intercept and slope components in addition to fixed effects (see the Supplement for details and code examples). Analyses were FDR corrected across both the main effects of T1INR (intercepts) and interactions with age (slopes) for cortical gray and white matter and subcortical gray matter (i.e., six tests). We conducted follow-up analyses on five subcortical regions (hippocampus, amygdala, putamen, caudate, thalamus) and two cortical regions (dorsolateral prefrontal cortex, dorsal anterior cingulate) previously associated with SES (15) to assess specificity. There were no FDR-significant nonlinear relations of T1INR, so T1INR by age squared interactions were not included in final models (see the Supplement). As an additional exploratory analyses, we examined the relationship of T1INR to the additional 71 regions in the Destrieux Atlas (50).

We generated individual intercepts and slopes of each brain metric across MRI 1–MRI 5 for each youth, using the same type of multilevel models described above. For brain metrics significantly related to T1INR, linear regressions examined whether the brain metric related to any of the outcomes associated with T1INR with FDR correction.
For any brain metric related to both T1INR and an outcome measure also associated with T1INR, we conducted mediation analyses using the PROCESS procedure (model 4) in R (52).

**RESULTS**

**Participant Characteristics**

Demographic characteristics and distributions of adaptive outcomes are shown in Table S1, with zero-order correlations among variables in Table S2. See the Supplement for additional analyses incorporating race.

**Early SES and Outcomes in the Transition to Adulthood**

As shown in Table 1, lower early SES (i.e., lower T1INR) was associated with worse cognitive function, more high-risk behaviors, and worse social and educational outcomes, but not with psychiatric outcomes (Table 1). All of these relationships remained significant when controlling for preschool psychopathology, maternal mental health, and cumulative life events. For psychiatric outcomes, there was still no significant relationship to T1INR if we examined a variable that excluded borderline symptoms (p > .67) and no significant relationships in logistic regressions using T1INR and sex to predict each of the six categories of diagnostic outcomes (all p values > .10). As described above, some of the outcome variables represent counts, for which we took the mean. Thus, we repeated all of the analyses with zero-inflated Poisson regression with the counts themselves, with the same results.

**Early SES and Trajectories of Brain Development**

As can be seen in Table 2, Figure 1, and Figure S2, there were main effects of lower early SES (i.e., lower T1INR) relating to both overall reduced (i.e., lower intercept) cortical gray matter and subcortical gray matter volumes, but no significant effect for white matter. T1INR also related to the slope of subcortical gray matter (i.e., significant interaction with age). As shown in Figure 1, the difference in subcortical gray matter between children with greater versus lesser T1INR increased across the course of development. These effects remained significant when including T1 depression, anxiety, and externalizing severity; life events; and maternal mental health (Table S3).

As shown in Table S4 and Figure S3, T1INR did not significantly relate to the volume of the dorsolateral prefrontal cortex or dorsal anterior cingulate (see Table S5 for exploratory analysis of the other 73 Destrieux regions, a few of which showed significant relations to T1INR after FDR correction). As shown in Table S6 and Figures S4 and S5, there were main effects of lower early SES relating to overall reduced (i.e., lower intercept) hippocampal, caudate, putamen, and thalamus volumes, but not amygdala volume or the slopes of any subcortical brain region. The relationships of T1INR to hippocampal, caudate, putamen, and thalamus volumes remained significant when controlling for T1 psychopathology, life events, and maternal psychopathology (Table S7).

**Trajectories of Brain Development and Early Adult Outcomes**

Early SES related to only the intercept, but not the slope, of cortical gray matter (e.g., main effect of T1INR but no interaction with age). However, early SES related significantly to both the intercept and slope of subcortical gray matter. Cognitive function and high-risk behaviors were significantly associated with intercepts of cortical gray volumes and both intercepts and slopes of subcortical gray volumes (Table 3). Poor social outcomes were related to cortical gray intercepts, and poor educational outcomes were related to subcortical gray slopes. Cognitive function and high-risk behaviors remained significantly related to subcortical intercepts and slopes when covarying for T1 psychopathology, life events, and maternal psychopathology (Table 3). As noted above, some of the outcome variables represent counts. Thus, we repeated all of the analyses with zero-inflated Poisson regression with the same pattern of results, other than education outcomes for subcortical gray slopes (p = .09 for Poisson regression), which did not survive correction for the full set of covariates (Table 3).

To determine whether subcortical intercepts and slopes were accounting for dissociable variance in outcomes, we ran linear models that included both. For cognitive outcomes, subcortical intercepts were no longer significant (standardized B = 0.19, t = 1.16, p = .25), but slopes remained trend level (standardized B = 0.23, t = 1.85, p = .07). For high-risk outcomes, subcortical intercepts were not

<table>
<thead>
<tr>
<th>Table 1. T1 Income-to-Needs Ratio Relating to Outcomes at the Transition to Adulthood</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Outcome Variable</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Cognitive Function Outcome</td>
</tr>
<tr>
<td>High-Risk Behaviors Outcome</td>
</tr>
<tr>
<td>Poor Social Outcomes</td>
</tr>
<tr>
<td>Poor Education Outcomes</td>
</tr>
<tr>
<td>Psychiatric Outcomes</td>
</tr>
</tbody>
</table>

Models controlling for T1 psychopathology were only run if the effect was significant in models not including T1 psychopathology. CI, confidence interval; Std., standardized; T1, time of first interview. *Survives false discovery rate correction across all five outcomes.
Figure 1. Trajectories of cortical and subcortical gray matter as a function of T1 income-to-needs ratio. (A) Trajectory of cortical gray matter, with the estimated fit lines plotted separately for the mean T1 income-to-needs and ±1 standard deviation. (B) Trajectory of subcortical gray matter, with the estimated fit lines plotted separately for the mean T1 income-to-needs and ±1 standard deviation. These graphs were generated using the original values for interpretation, although the analyses were run on standardized variables. Mean values of 0.5 (neutral) for sex and 1547.9 cm³ for intracranial volume were entered into the model equation. T1, time of first interview.

Table 2. Multilevel Models of T1 Income-to-Needs Ratio Relating to Cortical Gray, Cortical White, and Subcortical Gray Volumes

<table>
<thead>
<tr>
<th>Predictors</th>
<th>Std. B</th>
<th>SE</th>
<th>t</th>
<th>p</th>
<th>Semipartial R²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dependent Variable—Cortical Gray Matter Volume</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>0.0039</td>
<td>0.0286</td>
<td>0.14</td>
<td>.89</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>−0.4210</td>
<td>0.0117</td>
<td>−36.09</td>
<td>&lt;.001</td>
<td>0.372</td>
</tr>
<tr>
<td>Age squared</td>
<td>−0.0548</td>
<td>0.0103</td>
<td>−5.31</td>
<td>&lt;.001</td>
<td>0.010</td>
</tr>
<tr>
<td>Female sex</td>
<td>−0.0577</td>
<td>0.0315</td>
<td>−1.83</td>
<td>.07</td>
<td>0.015</td>
</tr>
<tr>
<td>Intracranial volume</td>
<td>0.7647</td>
<td>0.0317</td>
<td>24.12</td>
<td>&lt;.001</td>
<td>0.717</td>
</tr>
<tr>
<td>T1 income-to-needs ratio</td>
<td>0.1229</td>
<td>0.0295</td>
<td>4.16</td>
<td>&lt;.001</td>
<td>0.072</td>
</tr>
<tr>
<td>T1 income-to-needs ratio × age</td>
<td>0.0193</td>
<td>0.0098</td>
<td>1.97</td>
<td>.05</td>
<td>0.002</td>
</tr>
<tr>
<td>Dependent Variable—Cortical White Matter Volume</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>−0.0084</td>
<td>0.0321</td>
<td>−0.26</td>
<td>.79</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>0.2066</td>
<td>0.0063</td>
<td>32.69</td>
<td>&lt;.001</td>
<td>0.114</td>
</tr>
<tr>
<td>Age squared</td>
<td>−0.0695</td>
<td>0.0045</td>
<td>−15.39</td>
<td>&lt;.001</td>
<td>0.014</td>
</tr>
<tr>
<td>Female sex</td>
<td>0.0048</td>
<td>0.0352</td>
<td>0.14</td>
<td>.89</td>
<td>0.000</td>
</tr>
<tr>
<td>Intracranial volume</td>
<td>0.8550</td>
<td>0.0353</td>
<td>24.20</td>
<td>&lt;.001</td>
<td>0.740</td>
</tr>
<tr>
<td>T1 income-to-needs ratio</td>
<td>0.0224</td>
<td>0.0331</td>
<td>0.68</td>
<td>.50</td>
<td>0.002</td>
</tr>
<tr>
<td>T1 income-to-needs ratio × age</td>
<td>0.0078</td>
<td>0.0057</td>
<td>1.3</td>
<td>.17</td>
<td>0.000</td>
</tr>
<tr>
<td>Dependent Variable—Subcortical Gray Matter Volume</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intercept</td>
<td>−0.0009</td>
<td>0.0398</td>
<td>−0.02</td>
<td>.98</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td>0.4176</td>
<td>0.0165</td>
<td>25.34</td>
<td>&lt;.001</td>
<td>0.231</td>
</tr>
<tr>
<td>Age squared</td>
<td>−0.0300</td>
<td>0.0137</td>
<td>−2.19</td>
<td>.03</td>
<td>0.002</td>
</tr>
<tr>
<td>Female sex</td>
<td>−0.0760</td>
<td>0.0403</td>
<td>−1.88</td>
<td>.06</td>
<td>0.013</td>
</tr>
<tr>
<td>Intracranial volume</td>
<td>0.6046</td>
<td>0.0412</td>
<td>14.69</td>
<td>&lt;.001</td>
<td>0.496</td>
</tr>
<tr>
<td>T1 income-to-needs ratio</td>
<td>0.1645</td>
<td>0.0410</td>
<td>4.01</td>
<td>&lt;.001</td>
<td>0.067</td>
</tr>
<tr>
<td>T1 income-to-needs ratio × age</td>
<td>0.0363</td>
<td>0.0142</td>
<td>2.55</td>
<td>.01*</td>
<td>0.003</td>
</tr>
</tbody>
</table>

The multilevel models included both random intercept and random slope components in addition to fixed effects. Std., standardized; T1, time of first interview.

*Survives false discovery rate correction across both main effects and interactions with age (six total tests).
significant (standardized B = −0.15, t = −0.95, p = .34), but slopes were (standardized B = −0.25, t = −2.00, p = .047). Thus, for the mediations below, we focused on slopes and not intercepts.

Neither hippocampal nor caudate intercepts significantly related to any outcomes. Cognitive function related to both putamen and thalamic intercepts (Table S8), although only thalamus remained significant when accounting for additional covariates. High-risk behaviors were significantly related to both putamen and thalamic intercepts, even controlling for additional covariates of preschool psychopathology, maternal mental health, and life events.

Mediation

We only examined mediation for adaptive outcomes related to both T1INR (Table 2) and the brain metric (Table 3) with the full covariate set in the model. Slopes of subcortical gray matter significantly mediated the relationship between T1INR and both cognitive function and high-risk behavior even when controlling for additional covariates (Figure 2; Table S9). The bivariate relationships between T1INR and slopes of subcortical gray matter and between subcortical gray matter and both cognitive function and high-risk behavior are shown in Figure 3. The only significant mediation for individual subcortical volumes was thalamic volumes mediating the relationship between T1INR and cognitive function, but this did not survive inclusion of additional covariates (Table S10).

DISCUSSION

Preschool SES was associated with cognitive function, high-risk behaviors, social function, and educational function over 13 years later, even when controlling for preschool psychopathology, life events, and maternal psychopathology, providing robust evidence for the enduring influence of early childhood SES. Early SES was associated with both cortical and subcortical gray matter development, with the relationship of low preschool SES to differences in subcortical gray matter development increasing with age. Further, subcortical gray matter trajectories mediated the relationship between preschool SES and cognition and high-risk behaviors, even when controlling for preschool psychopathology, life events, and maternal psychopathology. Together, these novel longitudinal data underscore the key role of brain development in understanding the long-lasting relationships of early SES in children to later outcomes and highlight the need to address this pressing public health concern.

An unresolved question is the pathway by which early childhood SES relates to higher risk of poor adult outcomes. One possibility is that low early SES indicates ongoing low SES and thus ongoing exposure to a range of chronic stressors, reduced access to health care, poor nutrition, exposures to

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Table 3. Cortical and Subcortical Brain Volume Trajectories Relating to Late Adolescence/Early Adulthood Outcomes

<table>
<thead>
<tr>
<th>Outcome Variable</th>
<th>Independent Variable With Sex and Intracranial Volume as Covariates</th>
<th>Independent Variable With Sex; Intracranial Volume; T1 Depression, Anxiety, and Externalizing Severity; Cumulative Life Events; and Maternal Mental Health as Covariates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Std. B</td>
<td>Lower 95% CI</td>
</tr>
<tr>
<td>Cortical Gray Matter Intercepts</td>
<td>Cognitive function outcome</td>
<td>0.422</td>
</tr>
<tr>
<td></td>
<td>High-risk behaviors outcome</td>
<td>−0.335</td>
</tr>
<tr>
<td></td>
<td>Poor social outcomes</td>
<td>−0.397</td>
</tr>
<tr>
<td></td>
<td>Poor education outcomes</td>
<td>−0.186</td>
</tr>
<tr>
<td>Subcortical Gray Matter Intercepts</td>
<td>Cognitive function outcome</td>
<td>0.410</td>
</tr>
<tr>
<td></td>
<td>High-risk behaviors outcome</td>
<td>−0.375</td>
</tr>
<tr>
<td></td>
<td>Poor social outcomes</td>
<td>−0.169</td>
</tr>
<tr>
<td></td>
<td>Poor education outcomes</td>
<td>−0.222</td>
</tr>
<tr>
<td>Subcortical Gray Matter Slopes</td>
<td>Cognitive function outcome</td>
<td>0.341</td>
</tr>
<tr>
<td></td>
<td>High-risk behaviors outcome</td>
<td>−0.332</td>
</tr>
<tr>
<td></td>
<td>Poor social outcomes</td>
<td>−0.073</td>
</tr>
<tr>
<td></td>
<td>Poor education outcomes</td>
<td>−0.236</td>
</tr>
</tbody>
</table>

Models controlling for T1 psychopathology, life events, and maternal psychopathology were only run if the effect was significant in models not including these covariates. The multilevel models used to generate the intercepts and slopes including random intercept and random slope components in addition to fixed effects.

Cl, confidence interval; Std., standardized; T1, time of first interview.
<sup>a</sup>Survives false discovery rate correction across all four outcomes.
<sup>b</sup>Significant only when not controlling for T1 psychopathology.
<sup>c</sup>Relationships significant even when controlling for T1 psychopathology.
<sup>d</sup>Survives false discovery rate correction for the number of follow-up models run for that independent variable.
environmental toxins, and so on, that persist into adulthood. If so, it may not be the early low SES per se that relates to poor outcomes, but rather the fact that early low SES is a harbinger of ongoing low SES and chronic stressors that continue to be present through adulthood. The evidence that improving SES during childhood improves cognitive, educational, and mental health outcomes in youth is indirectly consistent with this interpretation (53–55). Alternatively, it may be that early childhood is a particularly sensitive period for poverty, with relationships that last even if a child moves out of poverty at a later point. This interpretation is consistent with the evidence that early childhood poverty has lasting negative relationships into adulthood, even when the individual’s SES improves (56–63). We cannot arbitrate between these possibilities with the current data because there is little social mobility in our sample and therefore we could not examine the relationships

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**Figure 2.** Mediation models for subcortical gray matter volumes. Mediation results were generated using the PROCESS 3.5 macro in R (version 4.03, beta 0.5). The “a” values refer to the relationship between the predictor and the mediator, and the “b” values refer to the relationship between the mediator and the outcome. The “c” values are the direct relationship between the predictor and the outcome with no mediator in the model, and the “c’” values are the relationship between the predictor and the outcome with the mediator in the model.

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**Figure 3.** Graphs illustrating bivariate relationships with subcortical gray matter volumes. (A) Scatterplot depicting the relationship between T1 income-to-needs ratio and the slope of subcortical gray matter volume, with the estimated linear fit and 95% confidence interval overlaid. (B) Scatterplot depicting the relationship between the slope of subcortical gray matter volume and cognitive function, with the estimated linear fit and 95% confidence interval overlaid. (C) Scatterplot depicting the relationship between the slope of subcortical gray matter volume and high-risk behaviors, with the estimated linear fit and 95% confidence interval overlaid. T1, time of first interview.
between variation in SES and brain volume over time within a child, but this is a critical area for future research. It will be important in future research to examine populations where SES varies more over the course of a child’s life or to examine interventions that improve SES. It will also be essential to better disentangle SES as one form of social disadvantage from other factors that may also confer social disadvantage, such as the effects of systemic racism. Further, it is important to acknowledge that there can be many factors that influence SES. We focused on family income-to-needs, but parental education and neighborhood income levels are also relevant.

It is relevant to note that we continued to see relationships of early SES to later adaptive outcomes and brain development even when we controlled for a variety of factors that often co-occur with low early SES and that could, in theory, be mediating mechanisms. This included preschool depression, anxiety, and externalizing symptoms; maternal psychopathology; and cumulative life events experienced by the child. The relationship between cortical gray matter intercepts and later outcomes were no longer significant when controlling for these factors, suggesting that they may play a role in linking cortical brain development to adult cognitive function, high-risk behaviors, and social function. However, the relationships of subcortical intercepts and slopes to both cognitive function and high-risk behaviors remained significant even when controlling for these factors. While efforts to further unpack the differential effects and causal pathways from SES to adaptive outcomes is of course worthwhile, the robust and powerful evidence here and in the literature repeatedly linking low childhood SES to a greater likelihood of poor adaptive outcomes suggests that directly addressing low SES itself may be the most tangible and effective intervention with greatest potential of broad impacts across many different mediating mechanisms. It is also possible that some of the relationship of early SES to brain development and poor adaptive outcomes is accounted for by shared genetics that contribute to poor adaptive function in both children and their parents (resulting in low early SES for their children), although there is evidence for causal effects of poverty and children’s outcomes (53–55).

We did not find a relationship between preschool SES and psychiatric outcomes at ages 16+ years. This result is not consistent with a body of research suggesting that poverty is associated with higher rates of psychopathology (1,20,21) and our own prior research in this sample indicating a relationship between early low SES and depression and externalizing behaviors prior to age 16 years through MRI wave 3 (24,31,64–65). There are two speculative explanations for this finding. One is that because this was a prospective sample followed over time with repeated assessments, youth may have been more likely to receive treatment, given potentially greater parental awareness of psychopathology. A second is that youth with greater mental health problems in childhood were less likely to stay in the study into the transition to adulthood. However, we compared depression, anxiety, and externalizing symptoms for those children who continued in the study at MRI waves 4/5 and those who did not, and there were no significant differences in these variables (Table S11).

We found robust relationships of preschool SES to brain development but with important evidence of specificity. Preschool SES related to cortical and subcortical gray matter but not cortical white matter, relationships that remained when controlling for early child mental health, cumulative life events, and maternal mental health. These results provide evidence that the relationship of early SES to subcortical brain development is not just reflecting greater levels of early psychopathology or some other risk factors that might also relate to brain development. Further, we found that subcortical gray matter volume, and more specifically reduced increases over development (slopes), served as a mediator of the relationship between early SES and cognitive and high-risk outcomes. The finding that early SES relates to slopes of subcortical gray matter has important developmental implications. It suggests that the relationship of SES to brain development is not static (e.g., present from the start with no change across development), but in the case of subcortical volumes, the difference between youth raised in low versus higher SES increases over development. We were somewhat surprised to find that cortical gray matter and white matter did not show similar relationships, but it may be that subcortical regions are uniquely susceptible to early and/or sustained low SES.

We found that early SES related to reductions in hippocampal, caudate, putamen, and thalamic volumes, but not amygdala, dorsolateral prefrontal, or dorsal anterior cingulate volumes, consistent with a number of prior studies (10–12,27,66,67). Reduced hippocampal volume in humans associated with early low SES has been interpreted as consistent with animal literature showing opposite effects of stress and environmental enrichment on hippocampal cell proliferation and dendritic length and branching (68,69). In turn, the hippocampus is important for both cognitive function and stress reactivity (70). Palacios-Barrios and Hanson have argued that impairments in self-regulation may be a final common pathway linking early adversity, alterations in hippocampal development and other brain regions, and poor physical and mental health outcomes (15). The striatum and the thalamus may also be critical components of such a pathway, given the role of the striatum in reward processing (71) and the thalamus as a key pathway of communication with cortical regions (72). Early-life stress in animal models alters the development of striatal regions involved in reward processing (73), as well as disrupting thalamic structure and function (74,75). However, we did not have direct measures of these functions in this work, and therefore these hypotheses about functional relevance are speculative and need prospective confirmation. We may not have found relationships to the amygdala, given the possibility that amygdala structure may be more related to threat-relevant experiences (76).

This study has several limitations. First, we did not start brain imaging until the children were school age and thus do not know how early variation in brain structure related to SES manifests. Second, we experienced a scanner upgrade between scan waves 3 and 4 and cannot make general statements about normative patterns of brain development, because overall shifts in volume across time could be confounded with the upgrade. However, we do not think the scanner change impairs our ability to look at individual differences in such trajectories, because all youth shifted scanner platforms at waves 4 and 5 and we used longitudinal harmonization methods. Third, it is possible that associations in
mediation models may be bidirectional. Fourth, while a number of our findings fit with the extant literature on SES (e.g., relationships to hippocampal volume, gray matter volume, cognitive function), the specialized nature of this sample (overrecruited for early depression) may limit generalizability. Fifth, to create interpretable outcome measures that aggregated multiple indicators, we had to dichotomize measures in ways that may have resulted in some loss of information.

In summary, this study provides highly novel longitudinal prospective data demonstrating that low childhood SES has long-lasting relationships to a range of outcomes at the transition to adulthood that are mediated in part by brain development. These data add to the ongoing need for policy initiatives that address this crisis, because childhood poverty is one of the most critical factors relating to long-term outcomes for far too many youth across the world.

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