Altered Gray Matter Volume and School Age Anxiety in Children Born Late Preterm

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Objectives To determine if late preterm (LP) children differ from full term (FT) children in volumes of the cortex, hippocampus, corpus callosum, or amygdala and whether these differences are associated with anxiety symptoms at school-age.

Study design LP children born between 34 and 36 weeks gestation and FT children born between 39 and 41 weeks gestation from a larger longitudinal cohort had magnetic resonance imaging scans at school-age. Brain volumes, cortical surface area, and thickness measures were obtained. Anxiety symptoms were assessed using a structured diagnostic interview annually beginning at preschool-age and following the magnetic resonance imaging.

Results LP children (n = 21) had a smaller percentage of total, right parietal, and right temporal lobe gray matter volume than FT children (n = 87). There were no differences in hippocampal, callosal, or amygdala volumes or cortical thickness. LP children also had a relative decrease in right parietal lobe cortical surface area. LP children had greater anxiety symptoms over all assessments. The relationship between late prematurity and school-age anxiety symptoms was mediated by the relative decrease in right temporal lobe volume.

Conclusions LP children, comprising 70% of preterm children, are also at increased risk for altered brain development particularly in the right temporal and parietal cortices. Alterations in the right temporal lobe cortical volume may underlie the increased rate of anxiety symptoms among these LP children. These findings suggest that LP delivery may disrupt temporal and parietal cortical development that persists until school-age with the right temporal lobe conferring risk for elevated anxiety symptoms. (J Pediatr 2014;165:928-35).

Preterm birth is a major public health problem with well-established high risk for adverse medical and developmental outcomes in survivors.1 These poor outcomes may be mediated by a greater risk for altered brain development associated with prematurity including decreased volumes of gray matter, white matter,2 and in particular regions like the hippocampus3 and the corpus callosum.4 These regional alterations have been found during the neonatal period with evidence of persistence into school-age,5,6 adolescence,7,8 and adulthood9,10 especially in those born prior to 34 weeks gestation. Global and regional reductions in gray and white matter volumes have even been found in school-age preterm children that were considered at low risk for neurodevelopmental deficits based on their gestational age (GA) and lack of significant medical complications.11

Studies of volumetric brain changes in children born preterm, however, have been limited to a focus on very preterm infants, even though late preterm (LP) infants, born between 34 and 36 weeks gestation, comprise approximately 70% of preterm births.12 LP infants may also be at elevated risk for disrupted brain development. Underscoring the significant brain development that occurs between 34 and 40 weeks, the 34-week brain weighs only 65% of the full term (FT) brain, and the cortical volume is only about one-half that of what it will be at 40 weeks.13 One study that evaluated neonatal brain volumes of LP infants at term-equivalent age found that they had significantly smaller gray matter volumes and percentage of gray matter volume than term-born infants.14 Conversely, a study focused on corpus callosum volume, did not find differences in callosal volumes at adolescence between 11 children born LP and 53 FT...
controls. Given the high prevalence of LP birth, the suggestion of structural differences in LP infants and the steep trajectory of brain development during these final gestational weeks, further investigation of structural brain outcomes in LP children is clearly indicated.

Prematurity is also associated with increased rates of psychiatric disorders including anxiety disorders. Indeed, increased rates of anxiety symptoms among those born very preterm (prior to 30-32 weeks gestation) have been reported at both childhood and adulthood, although findings have been mixed. Although there is less evidence in the LP population, available evidence supports an increased rate of psychiatric symptoms in LP children as well, including “emotional,” and anxiety symptoms. We have also previously noted increased rates of psychiatric disorders, including a 4-fold risk for anxiety disorders, at preschool age among LP children in the cohort to be examined in the analyses presented here. Elevated rates of generalized anxiety disorder (GAD) and separation anxiety disorders were found in this group. Although we found these early childhood anxiety disorders in LP children were mediated by maternal depression, underlying neurodevelopmental differences could confer additional risk.

There has been a burgeoning literature investigating whether altered brain development associated with preterm birth is associated with the increased risk of psychiatric symptoms, with links noted between both global and regional brain differences and childhood psychiatric symptoms including anxiety symptoms. Given the primary focus of the prior research on children born at earlier GAs, it remains unclear whether abnormalities in brain development are also found at an increased rate among LP children. It is also unknown if these brain alterations could underlie an increased risk of anxiety disorders. To address these research gaps, the current study aimed to assess whether children born LP differed from FT children in regions previously associated with preterm birth (hippocampus, corpus callosum) or anxiety (amygdala), in volume of cerebral gray matter or white matter, or in cortical surface area and thickness, and to determine whether any structural differences were related to differences in anxiety symptoms at school age between LP and FT children.

### Methods

Data for this analysis was obtained from 108 children born between 34 and 36 weeks GA (LP) or between 40 and 41 weeks (FT) from a larger sample enrolled in a 10-year longitudinal study investigating preschool depression (n = 306). The larger sample was recruited from day cares and preschools around metropolitan St. Louis using a screening checklist to oversample preschoolers with depressive and disruptive symptoms and to include healthy controls. Preschoolers with chronic medical or neurologic problems, intellectual disability, or autistic spectrum disorders were also excluded from the larger longitudinal study. As previously described, children and their caregivers participated in 3-6 comprehensive annual diagnostic and developmental assessments prior to their first neuroimaging session. Participants were screened for standard imaging contraindications. In addition, children with a history of head injury, ischemic insults or stroke or other brain injuries, seizure disorders, history of mechanical ventilation, or focal neurologic deficits on neurologic examination were also excluded. Of the 210 LP (n = 40) and FT (n = 170) children eligible for the imaging study, 122 had a magnetic resonance imaging (MRI) scan at school-age (6-12 years). Forty-eight children or families refused, 12 cancelled or did not keep appointments repeatedly, 20 had MRI contraindications, 2 were deceased, and 6 were lost to follow-up/lived out of state. Children also returned for 1 to 3 annual diagnostic assessments after the MRI scan. All study procedures were reviewed and approved by the institutional review board at the Washington University School of Medicine in St. Louis. Written informed consent was obtained from parents, and assent was obtained from children. Children with poor image quality were also excluded (n = 14) leaving a final sample of 108 children for inclusion in the analyses that follow.

### Preterm Birth

GA at birth (completed weeks) was reported by the child’s primary caregiver. GA groups were categorized as follows: LP (34-36 weeks), and FT (40-41 weeks). The focus of this analysis was on the LP population because of our prior findings of increased risk of psychopathology in this group compared with the FT children in this sample.

### Anxiety Symptoms

Trained staff conducted annual behavioral/developmental assessments of children and their parents/guardians. Prior to age 8, the Preschool-Age Psychiatric Assessment (PAPA) was administered to assess psychopathology. The PAPA is an interviewer-based diagnostic assessment with empirically established test re-test reliability that covers a broad range of psychiatric symptoms and impairment. The Childhood and Adolescent Psychiatric Assessment (CAPA) was used after age 8, and it also includes a child-report interview. All interviews were audi-taped for quality control and group calibration. In addition to diagnoses, both the PAPA and CAPA provide dimensional counts of symptoms. For this analysis, an anxiety symptom domain score was created for each annual assessment by summing the symptom counts for GAD, post-traumatic stress disorder (PTSD), and separation anxiety disorder. There are 6 possible GAD symptoms, 17 possible PTSD symptoms, and 8 possible separation anxiety disorder symptoms, for a maximum possible anxiety dimensional score of 31. Only symptoms from these anxiety disorders were included as they are assessed both on the PAPA and the CAPA.

### MRI Acquisition

Two 3-dimensional T1-weighted magnetization-prepared rapid gradient echo scans were acquired on a Siemens 3.0-T Tim Trio scanner (Siemens, Maryland Heights, Missouri).
without sedation (sagittal acquisition; repetition time = 2300 milliseconds; echo time = 3.16 milliseconds; inversion time = 1200 milliseconds; flip angle = 8°; 160 slices; 256 × 256 matrix; field of view = 256 mm; 1.0 mm isotropic voxels; time = 6.3 min per scan). The scan deemed to be of highest quality was used for subsequent processing.

**Image Analyses**

Total gray and white matter volumes were obtained using FreeSurfer v 5.1 (http://surfer.nmr.mgh.harvard.edu/), as previously detailed. Briefly, the white and pial surfaces were visually inspected and were regenerated with manual intervention when needed. Total gray matter volumes (cortical and subcortical), white matter volumes, and whole brain volume (total gray matter volume + white matter volume) were obtained. Freesurfer also utilizes an automated labeling system to parcellate the cortex into 34 gyral-based regions of interest. This parcellation is applied to the cortical surface using a registration procedure that aligns cortical folding patterns and probabilistically assigns a neuroanatomical label to every point on the cortical surface. Regions of interest were then summed to provide lobar cortical gray matter volumes, surface areas (defined as the area of the gray matter-white matter junction). Lobar cortical thickness was obtained by weighting the regional thicknesses by their corresponding surface areas. Freesurfer also provided volumes of the amygdala, hippocampus, cerebellum, and corpus callosum.

**Pubertal Status**

The Tanner staging questionnaire was used to measure children’s pubertal status at the time of the scan.

**Socioeconomic Status**

Income to need was used for socioeconomic status and was defined by total family income divided by the federal poverty level for a family of that size.

**Data Analyses**

All data analysis was conducted utilizing SPSS (v 21) statistical software (SPSS Inc, Chicago, Illinois). Differences in sociodemographic variables between the LP and FT groups were compared via t tests and χ² analyses. Differences in tissue and regional volumes were tested via general linear models, adjusted for age, sex, and pubertal status. Whole brain volume was also entered as a covariate to analyze whether any regional differences were attributable to differences in overall brain volume. To test for mediation, the PROCESS tool for SPSS was used, which is a regression-based approach to test the indirect effect of an independent variable on a dependent variable via a mediator. The significance of the indirect effect was determined via PROCESS by using 10,000 bootstrap resamples to generate 95% CIs (significant when not crossing zero).

**Results**

Characteristics for the participants are noted in Table I. LP children were significantly more likely to be white, have a lifetime history of an anxiety disorder, and have mothers with histories of maternal depression. LP children included in the analysis (n = 21) did not differ significantly from LP children that were not scanned or that had unusable MRI data (n = 19) in birthweight, GA, IQ, adjusted mean anxiety symptoms throughout the assessment periods, preschool-age anxiety symptoms, proportion with a lifetime anxiety diagnosis, income to need, sex, ethnicity, rate of neonatal intensive care unit hospitalization, or rates of maternal major depressive disorder. FT children included in the analysis (n = 87) also did not differ significantly from FT children that were not scanned or that did not have usable MRI data (N = 82) in these same key variables.

**Late Prematurity and Hippocampal, Amygdala, and Corpus Callosal Volumes**

There were no differences as a function of prematurity in either left (P = .78) or right (P = .39) hippocampal volume or left (P = .14) or right (P = .22) amygdala volume or total volume of the corpus callosum (P = .10) after adjusting for age, sex, pubertal status, and whole brain volume.

**Late Prematurity and Cerebral Gray and White Matter Volumes**

LP and FT children did not differ in whole brain volume (P = .45), or total gray (P = .96) or white matter volume (P = .10) after adjusting for age, sex, and pubertal status. After adjusting for whole brain volume in addition to age, sex, and pubertal status, LP children had significantly less relative

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**Table I. Cohort characteristics**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>LP (GA 34-36 wk) (N = 21)</th>
<th>FT (GA 40-41 wk) (N = 87)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male, %</td>
<td>61.9</td>
<td>51.7</td>
</tr>
<tr>
<td>Ethnicity, %</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>71.4</td>
<td>48.3*</td>
</tr>
<tr>
<td>Black</td>
<td>28.6</td>
<td>37.9</td>
</tr>
<tr>
<td>Other</td>
<td>15.4</td>
<td>13.8</td>
</tr>
<tr>
<td>GA at birth, wk, M (SD)</td>
<td>35.24 (0.8)</td>
<td>35.16 (0.4)</td>
</tr>
<tr>
<td>NICU hospitalization, %</td>
<td>23.8</td>
<td>9.2%</td>
</tr>
<tr>
<td>Days in NICU, M (SD)</td>
<td>13.6 (8.8)</td>
<td>5.6 (5.9)</td>
</tr>
<tr>
<td>Age at MRI scan, y, M (SD)</td>
<td>9.71 (1.27)</td>
<td>9.99 (1.28)</td>
</tr>
<tr>
<td>Range</td>
<td>6-11</td>
<td>7-12</td>
</tr>
<tr>
<td>Pubertal status, %</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prepubertal</td>
<td>57.1</td>
<td>48.8</td>
</tr>
<tr>
<td>Early pubertal</td>
<td>19.0</td>
<td>20.9</td>
</tr>
<tr>
<td>Midpubertal</td>
<td>23.8</td>
<td>24.4</td>
</tr>
<tr>
<td>Late pubertal</td>
<td>0.0</td>
<td>5.8</td>
</tr>
<tr>
<td>Income to needs ratio M (SD)</td>
<td>1.98 (1.3)</td>
<td>2.19 (1.32)</td>
</tr>
<tr>
<td>IQ, M (SD)</td>
<td>104.5 (12.7)</td>
<td>105.6 (16.1)</td>
</tr>
<tr>
<td>Anxiety disorder</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lifetime, %</td>
<td>76.2</td>
<td>42.5†</td>
</tr>
<tr>
<td>Current (after MRI), %</td>
<td>23.5</td>
<td>8.4</td>
</tr>
<tr>
<td>Number of annual diagnostic assessments, M (SD)</td>
<td>6.1 (1.2)</td>
<td>5.8 (1.8)</td>
</tr>
<tr>
<td>Time between MRI and last diagnostic assessment after MRI, mo (SD)</td>
<td>22.1 (11.2)</td>
<td>20.7 (10.0)</td>
</tr>
<tr>
<td>History of maternal MDD, %</td>
<td>71.4</td>
<td>34.5†</td>
</tr>
<tr>
<td>History of maternal anxiety, %</td>
<td>14.3</td>
<td>9.2</td>
</tr>
</tbody>
</table>

*P < .05. †P < .01.
total gray matter volume (standardized beta = −0.137; 95% CI −0.237 to −0.037; P = .008). After adjustment for the above mentioned covariates, LP children had on average 1% less percent gray matter volume (gray matter volume divided by whole brain volume) than the FT children (LP 62.3% [61.5%-63.0%]; FT 63.3% [62.8%-63.8%]; P = .007; bracketed values here and henceforth give 95% CIs). This decrease in percentage of relative total gray matter (which includes subcortical gray matter) was mainly due to a reduction in the relative percentage of cortical gray matter volume (LP 46.6%, [46.0%-47.3%]; FT 47.5%, [47.1%-48.0%]; P = .007). Given our prior report of a relationship between income to need and smaller gray matter volume in the larger study population, we added income to need to the model even though there was not a significant difference in socioeconomic status between the LP and FT children. Late prematurity remained a significant predictor of relative gray matter volume (standardized beta = −0.189 [−0.317 to −0.062]; P = .004).

**Exploratory Analyses of Regional Cortical Volume Differences**

Exploratory analyses were conducted to evaluate whether differences in relative gray matter volumes were diffuse or attributable to particular regional volume differences. Free-surface parcellated regions were combined based on the parcellations of Desikan et al.31 to calculate frontal, temporal, parietal, and occipital lobe gray matter volumes and cingulate gray matter volumes per hemisphere. These volumes were analyzed in MANCOVAs adjusting for age, sex, pubertal status, and whole brain volume for each hemisphere. For the right hemisphere there was an overall main effect of late prematurity (P = .002). Figure 1 shows follow-up analyses correcting for multiple comparisons (0.05/5 regions = 0.001) indicated that preterm children had significantly smaller right temporal lobes (adjusted mean 63 305 mm³ [61 972-64 638 mm³] vs adjusted mean 64 966 mm³ [61 158-65 775 mm³], respectively; P = .01) and right parietal lobes (adjusted mean 73 061 mm³ [71 348-74 774 mm³], vs adjusted mean 76 589 mm³ [75 550-77 628 mm³], respectively; P < .001). Further exploration revealed that the parcellated regions in the temporal lobe with significant differences were the right superior temporal gyrus (P = .036) and the right fusiform gyrus (P = .046), though these relationships did not persist after Bonferroni adjustment for multiple comparisons (alpha = 0.05/9 parcellated regions = 0.006). In the right parietal lobe, the right supramarginal gyrus was significantly smaller in the LP children (P < .001), which remained significant after adjustment for multiple comparisons (alpha = 0.05/5 = 0.01). There was no main effect for late prematurity in left hemisphere regional volumes.

**Cortical Thickness, Cortical Surface Area, and LP Birth**

There was no difference in total cortical surface area (P = .63) nor overall (mean) cortical thickness (P = .19) between FT and LP children after adjustment for age, sex, and pubertal status nor for cortical surface area after additionally adjusting for whole brain volume (P = .57). Given the above noted volume findings, we explored whether regions with volume differences between preterm and term children also differed in cortical surface area or cortical thickness. Regional surface area and thickness in the right parietal and right temporal lobes were analyzed in MANCOVAs adjusting for age, sex, pubertal status, and total surface area and mean thickness respectively. Right parietal surface area was significantly smaller in LP children compared with term-born children (adjusted mean 17 463 mm² [17 190-17 736 mm²], vs 17 759 mm² [17 625-17 893 mm²], P = .004). Further analyses demonstrated that the only parcellated region that was significantly decreased in LP children compared with FT children was the right supramarginal gyrus surface area (adjusted mean 3814 mm² [3592-4035 mm²], vs 4160 mm² [4025-4295 mm²], P = .002, alpha = 0.01 after Bonferroni correction). Right temporal lobe area trended toward a difference between groups (P = .057), but thickness did not (P = .19). There was also no difference in right parietal lobe thickness (P = .34) between groups.

**Late Prematurity and School-Age Anxiety Symptoms**

LP children who returned for at least 2 annual assessments (n = 38) had higher mean anxiety scores averaged over all assessments than FT children (n = 150) after adjusting for age and gender (mean 3.94, SD 12.04 vs mean 3.01, SD 1.66; beta 0.568, P = .009). This relationship remained significant when restricted to those who had an available MRI scan for analyses (LP n = 21, FT n = 87) even after adjusting for age at scan and sex (mean 4.19, SD 1.97 vs mean 2.90, SD 1.98; beta 1.31, P = .009).

**Total and Lobar Gray Matter Volume and Anxiety Symptoms**

We investigated whether the differences in total or lobar gray matter volumes (temporal and parietal, since these differed in LP children) between LP and FT children were related to increased anxiety symptoms among LP children. For those children with available assessment data in subsequent study waves following their school-age MRI scan, LP children (n = 17) still had higher anxiety scores than FT children (n = 83) after adjusting for age at scan and sex (mean 2.43, SD 1.84 vs mean 1.43, SD 1.44; beta 0.977, P = .017). Both percent total gray matter and percent right temporal lobe volume were significantly related to their later anxiety symptoms as noted in Table II, whereas percent right parietal lobe volume was not (P = .59). When both percent total gray matter and right temporal lobe volume were entered into the same model, only percent right temporal lobe volume remained significant (Table II). This finding raised the question of whether the alteration in the right temporal lobe mediated the relationship between LP birth and subsequent anxiety symptoms assessed
1.5 years later. We formally tested whether percent right temporal lobe volume mediated the relationship between LP birth and later school age anxiety symptoms. As noted in Figure 2, results yielded a significant indirect effect of LP birth on later anxiety symptoms through percent right temporal lobe volume (beta = 0.171, [-0.357 to 0.048]). The direct effect of preterm group on post MRI anxiety symptoms was no longer significant indicating that the percent right temporal lobe volume mediated the relationship between late prematurity and anxiety symptoms assessed after the MRI scan. We had previously found maternal major depressive disorders (MDD) to mediate preschool age anxiety symptoms. In the current study, LP children at school-age again had higher rates of maternal MDD. Thus, we assessed whether maternal MDD had a similar impact on post-scan anxiety symptoms. Maternal MDD was not significantly related to post scan anxiety symptoms (P = .46) after adjusting for age, sex, and LP birth. Percent right temporal lobe volume remained significant (P = .001) when adjusted for maternal MDD.

We also investigated whether the anxiety symptoms assessed at study waves before the MRI scan were related to the percent right temporal lobe volume. After adjusting for age, sex, LP birth, and puberty, the average of anxiety symptoms across the annual waves until the time of the scan was significantly related to the percent right temporal lobe volume (P = .02). Given this relationship, we added the mean anxiety symptom score measured until the time of the scan in the model predicting postscan anxiety symptoms. Percent right temporal lobe volume remained a significant predictor of postscan anxiety symptoms (beta = 150.48, [-32.94], P = .01). Given the above noted differences in specific regions of the right temporal gyrus (ie, superior) and right parietal cortex (ie, supramarginal gyrus) in FT and LP children, we also similarly assessed if these regional volumes related to anxiety symptoms. The right superior temporal gyrus volume was not related to pre-scan or post-scan anxiety symptoms in the entire cohort or after adjusting for LP birth.

The right supramarginal gyrus was associated with pre-scan anxiety symptoms (P = .04), but this relationship was not significant (P = .25) once LP birth was entered in the model.

### Discussion

This study comparing school-age children born LP and FT found that although there were no differences in total brain volume, LP children had a smaller percentage of their brain volumes as gray matter, particularly in the right temporal and parietal lobes. LP children also had a significantly smaller cortical surface area in the right parietal lobe (and nearly so in the right temporal lobe). In addition, LP children had more
Consistent with our volumetric results, among a sample of school-age children born between 28 and 41 weeks gestation, LP and FT children in this study—the right temporal and right parietal lobes. The temporal lobe has been highlighted as a region very susceptible to the deleterious effects of preterm birth and perhaps the region most vulnerable for LP infants as it is one of the latest maturing regions.36 This study extends those findings by highlighting 2 broad regions that accounted for the difference in gray matter volume between LP and FT children in this study—the right temporal and right parietal lobes. The temporal lobe has been highlighted as a region very susceptible to the deleterious effects of preterm birth and perhaps the region most vulnerable for LP infants as it is one of the latest maturing regions.36

Consistent with our volumetric results, among a sample of school-age children born between 28 and 41 weeks gestation, longer gestation was associated with greater gray matter density particularly in temporal and parietal regions.37 LP and FT children also differed in cortical surface area in the temporal lobe and even more prominently in the supramarginal gyrus in the parietal lobe, which has also been noted in other studies of children and adults born at earlier GAs.38,39 We did not, however, find differences in cortical thickness, as we did with cortical surface area, in the regions where we found volume differences. Prior work has found differences in preterm adolescents cortical thickness have noted this difference to be mostly attributable to children born at earlier GAs.40,41 Others who have investigated cortical thick-
many possible explanations for the volume changes reported in this study.

This study did not detect differences in hippocampus or corpus callosum volumes between LP and FT children. Previous studies of hippocampal volumes in preterm children have been mixed, with differences generally reported only in investigations of children born younger than 34 weeks gestation. As mentioned above, a previous study comparing LP and FT children also did not detect differences in corpus callosal volumes. Despite increased rates of anxiety disorders in the LP children, we also found no evidence of group differences in amygdala volumes. These negative findings, however, must be considered in the context that volumetric changes in these regions are inconsistently demonstrated in child and adolescent populations.

Although this study benefits from the use of longitudinal assessments utilizing semi-structured age-appropriate diagnostic interviews, there are some limitations. Oversampling of children with depressive symptoms, known to be comorbid with anxiety, prohibits inferences about the rates of anxiety among LP in the community. We previously noted, however, increased rates of anxiety disorders in this cohort in LP children without comorbid depression diagnoses. In addition, this study excluded children with known neurologic or major medical disorders. Thus, volumetric differences between more disabled LP children may differ or be more pronounced. Another potential limitation of the study is maternal report of GA, although maternal report has demonstrated high sensitivity and specificity for preterm birth suggesting sufficient accuracy of maternal report. Another potential limitation of the study design is that the interval between MRI and psychiatric evaluation was somewhat variable for each child. There was, however, no significant difference in this interval between the LP and FT children. Also, while percent right temporal lobe volume mediated the relationship between LP birth and later school age anxiety symptoms, mediation analysis does not necessarily prove a causal relationship. Finally, future work would benefit from cohorts with larger samples of LP children.

The findings in this study indicate that altered brain development particularly in the right temporal lobe may relate to the increased rate of anxiety symptoms found in LP children at school-age. Such a risk trajectory is plausible given the known function of this region in anxiety related emotion processing. It remains unclear if these altered brain volumes were present at birth and represent a biomarker of risk for the preschool age anxiety symptoms previously reported in this cohort. The finding of a significant relationship between anxiety symptoms prior to the scan and percent temporal lobe volume is suggestive. Future work that evaluates brain volumes at birth and assesses subsequent development of behavioral inhibition or later preschool anxiety symptoms will be needed to identify these children at the earliest possible time point for intervention. Future work will also need to determine the functional correlates of these structural alterations, which may help guide future treatment development in this population particularly vulnerable to anxiety disorders.

In summary, school-age children born LP had significantly smaller adjusted right temporal lobe and right parietal lobe gray matter than children born FT. The increased rate of anxiety symptoms in LP children persisted into school-age and was mediated by this alteration in right temporal lobe gray matter.

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