Archival Report

Microstructure of the Dorsal Anterior Cingulum Bundle in Very Preterm Neonates Predicts the Preterm Behavioral Phenotype at 5 Years of Age

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ABSTRACT

BACKGROUND: The cingulum bundle (CB), specifically the dorsal anterior portion of the CB, plays an important role in psychiatric illnesses; however, its role during early development is unclear. This study investigated whether neonatal white matter microstructure in the CB and its subregions is associated with subsequent preterm behavioral phenotype symptoms (internalizing, inattention, and social deficits) in very preterm (VPT) children.

METHODS: Diffusion magnetic resonance imaging data were obtained on a 3T scanner in 138 sleeping nonsedated neonates: 55 full-term neonates (gestational age ≥ 36 weeks) and 83 VPT neonates (gestational age < 30 weeks). The CB was tracked using probabilistic tractography and split into anterior and posterior portions. When children were 5 years of age, parents (n = 80) and teachers (n = 63) of VPT children completed questionnaires of preterm behavioral phenotype symptoms. Linear regression models were used to relate measures of neonatal CB microstructure and childhood preterm behavioral phenotype symptoms (n = 56 parent report, n = 45 teacher report).

RESULTS: Mean diffusivity in the anterior and posterior CB was increased in VPT neonates compared with full-term neonates. Increased fractional anisotropy and decreased mean diffusivity in the right anterior CB, but not in the posterior CB, were related to increased preterm behavioral phenotype symptoms in VPT children as reported by parents and teachers.

CONCLUSIONS: Aberrations in the anterior portion of the right CB may underlie the early development of the preterm behavioral phenotype. This finding provides the foundation for future mechanistic and therapeutic investigations into the role of the anterior cingulum in the development of psychopathology in VPT infants.

Keywords: ADHD, Anxiety, Autism, DTI, Neonatal, Preterm

https://doi.org/10.1016/j.biopsych.2020.06.015

Premature birth remains a major public health issue in the United States, with approximately 1 in 10 infants born prior to 37 weeks’ gestation (1). Preterm infants can experience a range of adverse outcomes, including psychiatric impairments such as internalizing disorders, attention-deficit/hyperactivity disorder (ADHD), and autism (2,3). Numerous studies have shown that preterm children demonstrate specific impairments in three key domains of psychopathology—emotional processing, attention, and social communication—with rates of impairment three to four times higher in preterm children compared with full-term (FT) children (2,4). Furthermore, preterm infants born at earlier gestational ages (GA) have greater risks of psychiatric illnesses, with very preterm (VPT) infants (born <32 weeks gestation) demonstrating greater risk than moderate to late preterm infants (5,6). The consistent pattern of elevated rates of internalizing disorders, ADHD, and autism (without concurrently elevated rates of conduct disorder) has been recognized as the preterm behavioral phenotype (2,3,7,8).

Preterm infants have also been shown to demonstrate altered structural and functional brain connectivity compared with their FT peers. Multiple studies using conventional and diffusion magnetic resonance imaging have demonstrated global gray and white matter alterations in VPT infants at term-equivalent age (9–12), with differences persisting into adolescence and adulthood (13,14). Diffusion tensor imaging (DTI) specifically has been used to assess these white matter abnormalities and link aberrant white matter microstructure with later psychiatric impairments. In VPT infants, neonatal white matter abnormalities qualitatively graded based on overall severity have been associated with lower socioemotional competence scores at 2 years of age (15), with additional evidence demonstrating that the increased fractional anisotropy (FA) in the cingulum bundle (CB) may play a key role during early development of socioemotional competence (16).

The CB is a white matter tract that connects the anterior cingulate, dorsolateral prefrontal, medial prefrontal, and orbitofrontal cortices to the insula and amygdala ventrally and to the
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Methods

Sample and Procedure

This study used a prospective longitudinal design that included neonatal neuroimaging and behavioral follow-up at 5 years of age in a cohort of VPT and FT infants. VPT infants (born at <30 weeks GA, range = 23–30 weeks; n = 83) were recruited prospectively from the St. Louis Children’s Hospital level 3 neonatal intensive care unit. Singleton FT infants (born at ≥36 weeks GA, range = 36–41 weeks; n = 55) were recruited from the adjoining maternity ward at Barnes-Jewish Hospital. Exclusion criteria for both groups included chromosomal abnormalities, congenital infection, high-grade intraventricular hemorrhage, cystic periventricular leukomalacia, moderate to severe cerebellar hemorrhage, and deep nuclear gray matter lesions. Additional exclusion criteria for FT infants included a positive maternal urine drug screen or neonatal acidosis on cord blood gas (pH < 7.20).

DTI data were obtained as part of a multimodal acquisition on a Siemens Tim Trio 3T scanner (Erlangen, Germany) using an infant-specific quadrature head coil (Advanced Imaging Research, Cleveland, OH). All infants were imaged without sedation during natural sleep or while resting quietly. FT infants were scanned within the first 48 to 72 hours of life, and VPT infants were scanned at term-equivalent age (35–42 weeks GA). Medical personnel were present in the scanner room throughout the scan acquisition and monitored each infant’s heart rate and oxygen saturation continuously.

When children were 5 years of age (mean = 5.6 years, range = 4.7–6.5), the parents of a subset of the VPT infants with neonatal neuroimaging (n = 80) completed questionnaires to assess children’s psychiatric symptoms. Teachers of the children who participated at 5 years of age were contacted and asked to complete behavioral questionnaires that were analogous to the parent questionnaires. Written informed consent was obtained from primary caregivers prior to participation in the study. All study procedures were approved by Washington University Institutional Review Board.

Measures

Diffusion Tensor Imaging. DTI data were collected using a diffusion-weighted sequence (repetition time = 13,300 ms, echo time = 112 ms, field of view = 128 mm, voxel size = 1.2 × 1.2 × 1.2 mm³, bandwidth = 1266 Hz/pixel, and 48b amplitudes and directions). The diffusion signal attenuation curve was modeled as a monoexponential function plus a constant, and diffusion parameters were established using Bayesian probability theory (56). Maps of FA, MD, RD, and AD were generated. Region-of-interest masks were placed in the front and back of the dorsal CB with a forced waypoint in the middle. The dorsal CB (Figure 1A) and a control tract, the corpus callosum (see Supplement), were tracked in native space using Bayesian probabilistic tractography with FDT in FSL (FMRI Software Library, version 6) (57,58). Owing to technical limitations, the subgenual portion of the CB was not tracked.

The dorsal CB was manually segmented into anterior and posterior sections using the intersection of the anterior portion of the internal capsule and the posterior portion of the internal capsule as an anatomical landmark by a single rater (RGB with oversight from CDS and JSS) (Figure 1B). This landmark was chosen based on the anatomical landmarks corresponding to previous segmentations of the CB in nonhuman primates (23). Measures of FA, MD, RD, and AD were extracted from the CB.
Extreme outlier values (>3 SD) for each DTI metric within each tract segment were excluded from analyses. The demographic characteristics of the birth sample are displayed in Table 1.

**Behavioral Phenotype: Parent Report.** Parents completed the Child Behavior Checklist for ages 1.5 to 5 years (CBCL) (59), the Conners Rating Scale–Revised (Conners) (60), and the Social Responsiveness Scale–Second Edition (SRS-2) (61) in 80 VPT children. Internalizing problems were measured using the internalizing symptoms score on the CBCL, which comprises four subscales: anxiety/depression, withdrawn, somatic complaints, and emotional reactivity. Autism symptoms, including social communication impairments and restrictive and repetitive behaviors, were assessed using the SRS-2 total score (61). ADHD symptoms, including hyperactivity/impulsivity, inattention, and cognitive problems, were measured using the Conners ADHD Index (62). The CBCL, SRS-2, and Conners ADHD Index all have been well validated for use in diverse samples of children, including those from low socioeconomic status backgrounds and racial/ethnic minorities (63–66). The SRS-2 and Conners were chosen to assess autism and ADHD symptoms, respectively, because they assess multiple domains within each diagnostic category and are more specific than the CBCL subscales. On all measures, higher scores indicated greater symptom severity. The clinical cutoff score is 65 for the CBCL and 60 for the SRS-2 and Conners (59–61). In this cohort, previous research has shown that VPT children have increased psychiatric symptoms.

### Table 1. Neonatal White Matter Microstructure Differences in the CB

<table>
<thead>
<tr>
<th></th>
<th>Full-term, n = 55</th>
<th>Very Preterm, n = 69</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational Age at Birth, Weeks</td>
<td>38.9 ± 1.3</td>
<td>26.7 ± 1.8</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>Gestational Age at Scan, Weeks</td>
<td>38.7 ± 1.2</td>
<td>37.5 ± 1.4</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>FA of Right CB</td>
<td>0.184 ± 0.02</td>
<td>0.181 ± 0.02</td>
<td>.715</td>
</tr>
<tr>
<td>FA of right anterior CB</td>
<td>0.168 ± 0.02</td>
<td>0.154 ± 0.02</td>
<td>.060</td>
</tr>
<tr>
<td>FA of right posterior CB</td>
<td>0.219 ± 0.03</td>
<td>0.222 ± 0.02</td>
<td>.479</td>
</tr>
<tr>
<td>FA of Left CB</td>
<td>0.194 ± 0.02</td>
<td>0.187 ± 0.01</td>
<td>.724</td>
</tr>
<tr>
<td>FA of left anterior CB</td>
<td>0.175 ± 0.02</td>
<td>0.162 ± 0.02</td>
<td>.155</td>
</tr>
<tr>
<td>FA of left posterior CB</td>
<td>0.223 ± 0.03</td>
<td>0.230 ± 0.02</td>
<td>.278</td>
</tr>
<tr>
<td>MD of Right CB</td>
<td>1.35 ± 0.06</td>
<td>1.40 ± 0.06</td>
<td>&lt;.002*</td>
</tr>
<tr>
<td>MD of right anterior CB</td>
<td>1.37 ± 0.07</td>
<td>1.42 ± 0.08</td>
<td>&lt;.002*</td>
</tr>
<tr>
<td>MD of right posterior CB</td>
<td>1.32 ± 0.06</td>
<td>1.36 ± 0.06</td>
<td>.005*</td>
</tr>
<tr>
<td>MD of Left CB</td>
<td>1.34 ± 0.05</td>
<td>1.41 ± 0.07</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>MD of left anterior CB</td>
<td>1.36 ± 0.06</td>
<td>1.43 ± 0.08</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>MD of left posterior CB</td>
<td>1.32 ± 0.06</td>
<td>1.37 ± 0.06</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>Male/Female, n</td>
<td>24/31</td>
<td>29/40</td>
<td>.857</td>
</tr>
<tr>
<td>African American/Caucasian, n</td>
<td>40/15</td>
<td>29/34</td>
<td>.003*</td>
</tr>
</tbody>
</table>

Values are mean ± SD except where noted.
CB, cingulum bundle; FA, fractional anisotropy; MD, mean diffusivity.

*Value that passed false discovery rate correction.

**Figure 1.** (A) Dorsal cingulum bundle in a representative single subject. (B) Anterior–posterior divide, which was placed at the intersection between the anterior limb of the internal capsule (ALIC) and the posterior limb of the internal capsule (PLIC) for each subject. (C) Dorsal anterior cingulum bundle and dorsal posterior cingulum bundle after the division. Diffusion measures were extracted from the tracts in (A) and (C).
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Biological

of autism symptoms and ADHD symptoms, respectively. ADHD Index score were used for the teacher report measures parent report data, the SRS-2 total score and the Conners clinical cutoff score for the TRF is 65 (59). Analogous to the (see Supplement).

Because the aim of the study was to examine whether the CB is associated with common shared variance across the three domains of the preterm behavioral phenotype, a principal component analysis (PCA) was performed to assess the degree of variance that was explained by a composite measure of the preterm behavioral phenotype and to identify the amount of common variance in parent and teacher ratings (see Supplement). In the initial PCA, parent and teacher ratings loaded onto separate factors, and because the sample size was reduced when both parent and teacher variables were used in the model, separate PCAs were performed for parent and teacher variables. Both PCAs produced single-factor models explaining 80% of the variance with factor loadings between 0.89 and 0.92 for all variables, which eliminated the need to rotate the solution. These principal component factors were subsequently used in behavioral analyses labeled as the parent-rated preterm behavioral phenotype score and the teacher-rated preterm behavioral phenotype score.

To address the second aim of the study, we first examined the relationship between neonatal white matter microstructure of the dorsal CB and preterm behavioral phenotype symptoms. Then, we investigated whether the dorsal anterior and/or dorsal posterior portions of the CB were responsible for any significant associations between the dorsal CB and psychiatric outcome measures. Linear regression models adjusted for age at scan, sex, and medical risk were performed in VPT children (n = 56 parent rated; n = 45 teacher rated). Medical risk is a composite measure of perinatal clinical factors (see Supplement) that has been previously validated in the literature (3,71). Age at scan, sex, and medical risk were included in the model because they were related to the CB or outcome measures (see Supplement). Social risk was not included in the models because it was not related to any variables of interest for VPT children (see Supplement). Separate linear regression models were run for each diffusion measure (i.e., FA, MD, AD, and RD). Cronbach’s alpha (α) was set at .05. Multiple comparisons were accounted for using a Benjamini–Hochberg procedure for false discovery rate correction across all the statistical tests in the article (72).

RESULTS

White Matter Microstructure of the CB in VPT Versus FT Neonates

Diffusion measures in the whole, right, and left CB differed between VPT infants (n = 69) and FT infants (n = 55). Specifically, VPT infants had increased MD, AD, and RD in the CB.
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compared with FT infants (Table 1 and Table S2), FA of the CB did not differ between VPT and FT neonates (Table 1). In both the anterior and posterior portions of the CB, VPT neonates had greater MD, AD, and RD than FT neonates but had similar FA values (Table 1 and Table S2).

Neonatal CB and Parent-Rated Preterm Behavioral Phenotype

Increased FA of the right CB in the neonatal period was associated with greater parent-rated preterm behavioral phenotype symptoms at 5 years of age in VPT children (Figure 2). In contrast, there was no association with the FA of the left CB (Table 3). Similarly, MD, AD, and RD of the right and left CB were not associated with the parent-rated preterm behavioral phenotype scores (Table 3). To investigate whether the association between FA of the right CB and the parent-rated preterm behavioral phenotype score was regionally specific, we examined subregions of the CB and found that increased FA of the right anterior CB, but not of the right posterior CB, was associated with greater parent-rated preterm behavioral phenotype symptoms (Figure 2). Additional supplemental analyses supported the specificity of these findings given that the FA and MD of the corpus callosum were not related to the parent-rated preterm behavioral phenotype scores and the FA and MD of the CB were not related to Full Scale IQ (see Supplement).

Neonatal CB and Teacher-Rated Preterm Behavioral Phenotype

Increased FA and decreased MD of the right CB in the neonatal period were associated with greater teacher-rated preterm behavioral phenotype symptoms at 5 years of age (Figure 3). Decreased RD of the right CB was related to greater teacher-rated preterm behavioral phenotype symptoms, but decreased AD of the right CB was not (Table 3). The left CB was not related to the teacher-rated preterm behavioral phenotype symptoms on any diffusion measures (Table 3). When evaluating subregions of the CB, decreased MD, AD, and RD of the right anterior CB were associated with greater teacher-rated preterm behavioral phenotype symptoms (Figure 3). The right posterior CB was not related to any teacher-rated preterm behavioral phenotype scores (Figure 3). Additional supplemental analyses supported the specificity of these findings given that the FA and MD of the corpus callosum were not related to the teacher-rated preterm behavioral phenotype scores and the FA and MD of the CB were not related to Full Scale IQ (see Supplement).

DISCUSSION

Summary of Key Findings

The critical finding in this prospective longitudinal study was that neonatal white matter microstructure alterations in the right anterior CB are related to the severity of preterm

Table 3. Regression Models Predicting the Preterm Behavioral Phenotype

<table>
<thead>
<tr>
<th></th>
<th>Parent</th>
<th>Teacher</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>β</td>
<td>p</td>
</tr>
<tr>
<td>Fractional Anisotropy</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Right CB</td>
<td>.329</td>
<td>.017*</td>
</tr>
<tr>
<td>Right anterior CB</td>
<td>.301</td>
<td>.038*</td>
</tr>
<tr>
<td>Right posterior CB</td>
<td>.124</td>
<td>.400</td>
</tr>
<tr>
<td>Left CB</td>
<td>-.022</td>
<td>.873</td>
</tr>
<tr>
<td>Mean Diffusivity</td>
<td></td>
<td></td>
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<tr>
<td>Right CB</td>
<td>-.191</td>
<td>.232</td>
</tr>
<tr>
<td>Right anterior CB</td>
<td>-.078</td>
<td>.506</td>
</tr>
<tr>
<td>Right posterior CB</td>
<td>.007</td>
<td>.940</td>
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<tr>
<td>Left CB</td>
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<td>.583</td>
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<tr>
<td>Axial Diffusivity</td>
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<tr>
<td>Right CB</td>
<td>.001</td>
<td>.932</td>
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<td>Right anterior CB</td>
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<td>.911</td>
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<tr>
<td>Right posterior CB</td>
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<td>.414</td>
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<td>Left CB</td>
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<tr>
<td>Radial Diffusivity</td>
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<td>Right CB</td>
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<tr>
<td>Right anterior CB</td>
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<td>.300</td>
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<tr>
<td>Right posterior CB</td>
<td>-.042</td>
<td>.790</td>
</tr>
<tr>
<td>Left CB</td>
<td>-.093</td>
<td>.527</td>
</tr>
</tbody>
</table>

CB, cingulum bundle.

*p Value that passed false discovery rate correction.
behavioral phenotype symptoms at 5 years of age as rated by both parents and teachers. This result advances the field by showing that the white matter alterations in the anterior CB seen in adults with psychiatric illnesses (24,25,33) may be identifiable in the neonatal period as early as term-equivalent age. Specifically, we found that increased FA of the right anterior CB in the neonatal period was related to greater parent-rated preterm behavior phenotype symptoms at 5 years in VPT children. Decreased MD, RD, and AD of the right anterior CB were related to greater teacher-rated preterm behavioral phenotype symptoms. Importantly, this work suggests that white matter alterations in the anterior CB are more likely to contribute to the emergence of psychiatric symptoms than to be a downstream consequence of psychiatric illnesses, identifying the CB microstructure as a potentially viable therapeutic target.

The Anterior CB’s Role in Internalizing Disorders, Autism, and ADHD

In older children and adults, the anterior CB has been associated with internalizing disorders, autism, and ADHD, which comprise the preterm behavioral phenotype. Many studies show reduced FA in the dorsal and anterior portions of the CB in patients with internalizing disorders (26–30). Similar studies in autistic populations show decreased FA, increased MD, and increased RD in the anterior and dorsal segments of the CB (31–33). In addition, Makris et al. (24) and Konrad et al. (25) demonstrated that there was decreased FA and increased MD in the right anterior CB in adults with ADHD (24,25). Across these investigations, right hemisphere–specific findings were thought to occur because the right prefrontal cortex directs attention (73) and contributes to emotional-attentional modulation (74). Previous studies also supported the specificity of this association given that patients with anterior cingulotomies did not have lasting cognitive deficits (75,76). Our study further demonstrates this specificity by showing that early variance in right anterior CB microstructure is not associated with Full Scale IQ. This may indicate that white matter microstructure disruption in the CB plays a specific role in the early development of psychiatric illnesses.

The white matter of the CB may be particularly important during the early development of psychiatric illnesses because it connects key socioemotional brain regions. The CB connects the anterior cingulate, dorsolateral prefrontal, medial prefrontal, and orbitofrontal cortices to the insula and amygdala ventrally and to the posterior cingulate, parietal, hippocampus, and parahippocampal cortices posteriorly (22). These regions play important roles in social and emotional functioning (22,34), with the prefrontal, orbitofrontal, and anterior cingulate cortices thought to be particularly important. These frontal regions—and their connections to the amygdala—are thought to support emotion processing, emotion regulation, and executive control (34–40) and to be involved in internalizing disorders, ADHD, and autism (16,24,34,40–65). Disruptions in the anterior CB may be associated with psychiatric symptoms because of the multitude of short-range connections between prefrontal areas and the anterior cingulate cortex as well as long-range connections to the amygdala (22).
Microstructural Differences Between Neonates and Older Individuals

The prior literature linking the dorsal anterior CB to psychiatric illnesses aligns with our results except for a developmental shift in directionality. In adults, lower FA and increased MD, which are thought to represent decreased myelination and reduced axonal strength, have been associated with greater psychiatric impairment; however, in neonates, higher FA and lower MD have been associated with increased symptomatology (77–79). This change in directionality is thought to occur because late-developing white matter tracts, such as the CB, are incompletely myelinated at term-equivalent age (17,80). In the newborn brain, increases in FA and decreases in MD may reflect reduced fiber branching, smaller axonal diameters, and/or fewer crossing fibers (61). This is especially true for VPT infants who have reduced myelination compared with FT infants owing to disruptions in late oligodendrocyte progenitor cells and subplate neurons (19,82,83), which are selectively vulnerable to oxidative stress and excitotoxicity during development (20,84–86). Therefore, comparisons between VPT and FT infants may reflect differences in myelina- tion (10,12,87), whereas comparisons within VPT populations may be more sensitive to reduced fiber branching, smaller axonal diameters, and/or fewer crossing fibers.

Unexpected brain–behavior relationships in VPT children may also be attributable to the fact that DTI relies on average measures within a single voxel, making crossing fibers particularly difficult to delineate in the neonatal brain. If the vectors of crossing fibers point in different directions within a voxel, the average of the vectors will have less directionality (low FA) and greater diffusivity (high MD) (56,88). Prior anatomical investigations show that the CB contains many short-range corticocortical association fibers that cross the medial portions of the frontal, parietal, and temporal lobes (23,79,89,90). These beneficial crossing fibers may be absent in VPT infants with greater preterm behavioral phenotype symptoms. A previous analysis conducted in this VPT cohort showed that higher FA in the CB was related to increased autism symptoms at 2 years of age (77). Similar associations regarding higher FA have also been reported in other samples of VPT infants as well as infants at risk for autism spectrum disorder during childhood, suggesting a developmentally specific relationship between neonatal DTI values and later psychiatric illnesses (79,91).

Comparison of Results Across Parents and Teachers

Our findings linking the microstructure of the CB to parent and teacher reports of the preterm behavioral phenotype were largely similar; however, teacher reports were associated with more diffusion measures than parent reports. These differences in brain–behavior relationships were not unexpected given that multiple studies report parent–teacher disagreements, especially in preterm populations (72,92,93). The behavioral demands placed on children at school and in the home may differ in a context-dependent manner. Teachers are able to observe children in relation to a large peer group and may be better positioned to identify social communication difficulties as well as classroom inattention (94). Parents, on the other hand, may be more aware of internalizing problems because they spend more one-on-one time with their children (95). Differences in associations between the CB and the parent-rated and teacher-rated preterm behavioral phenotypes likely stem from atypical behaviors that are more observable in one context than in another. In support of this interpretation, diffusion measures in the right anterior CB were related to teacher, but not parent, ratings of autism symptoms (see Supplement), which would be expected if social communication deficits are more observable at school than at home. Each rater’s biases and expectations may also play a role.

Limitations

The first limitation of this study is that the subgenual portion of the CB was not assessed because of technical limitations. While the dorsal portion of the CB is thought to be more important for psychopathology in older populations, the subgenual portion has many connections within limbic regions. If these subgenual intralimbic connections are important during development, this study may underestimate the importance of the CB in later psychopathology. The second limitation of this study was our inability to differentiate crossing fibers owing to the limited dimensionality of the DTI data. In addition, owing to sample size limitations, we were unable to assess the relationship between the neonatal CB and psychiatric outcomes in FT infants. Even though FT infants do not exhibit the preterm behavioral phenotype, examining FT infants could help determine whether the link between white matter disruptions in the CB and psychiatric impairments is specific to prematurity. Future studies with higher DTI dimensionality, novel computational approaches, and larger sample sizes may allow for assessment of the subgenual CB and crossing fibers in both VPT and FT neonates.

Conclusions

This prospective, longitudinal, dual-informant study provides the first evidence that alterations in the anterior CB in neonates relate to specific psychiatric outcomes 5 years later. The results show that increased FA and decreased MD, AD, and RD of the right anterior CB at term-equivalent age is related to greater internalizing, autism, and ADHD symptoms at 5 years of age in VPT children. In addition, these results show that subregions of the CB have early functional specificity in the context of prematurity. Taken together, these findings highlight the importance of studying the role of the CB and its subregions in the emergence of the preterm behavioral phenotype. Early assessment of the CB in preterm neonates may be useful for the prognostication of later psychiatric disease and may help clinicians better allocate resources and services prior to the onset of symptoms. Crucially, this work provides the foundation for future studies of the CB that both advance our understanding of the pathophysiology underlying psychiatric diseases in the context of prematurity and may lead to new therapeutic interventions for vulnerable premature infants.

ACKNOWLEDGMENTS AND DISCLOSURES

Research reported in this publication was supported by the National Institutes of Health (Grant Nos. T32-EB014855 [to RGB], R01-HD057098 [to CDS, CER], R01-MH113570 [to CDS, CER], K02-NS089852 [to CDS], UL1-TR000448 [to CDS, CER], K23-MH105179 [to CER], and R01-HD061619 TR000448 [to CDS, CER], K23-MH105179 [to CER], and R01-HD061619
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