Diffusion Tractography and Neuromotor Outcome in Very Preterm Children with White Matter Abnormalities

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Abstract

Background—Moderate-severe white matter abnormality (WMA) in the newborn has been shown to produce persistent disruptions in cerebral connectivity, but does not universally result in neurodevelopmental disability in very preterm (VPT) children. The aims of this hypothesis driven study were to apply diffusion imaging to: 1) examine whether bilateral WMA detected in VPT children in the newborn period can predict microstructural organization at age 7; 2) compare corticospinal tract (CST) and corpus callosum (CC) measures in VPT children at age 7 with neonatal WMA with normal versus impaired motor functioning.

Methods—Diffusion parameters of the CST and CC were compared between VPT 7-year-olds with (n=20) and without (n=42) bilateral WMA detected in the newborn period. For those with WMA, diffusion parameters were further examined.

Results—Microstructural organization of CST and CC tracts at age 7 years were altered in VPT children with moderate-severe WMA detected at term equivalent age compared to those without injury. Furthermore, diffusion parameters differed in the CC for children with WMA categorized by motor outcome (N=8).

Conclusions—WMA on conventional MRI at term equivalent age is associated with altered microstructural organization of the CST and CC at 7 years of age.
Introduction

Early brain development can be disrupted by white matter abnormality (WMA), which is common in infants born preterm (1). Studies employing conventional (T1- and T2-weighted) MRI have demonstrated that up to 50% of very preterm (VPT) infants have WMA accompanied by neuronal or axonal disease (2). In these infants, moderate-severe WMA identified on qualitative MRI assessment, including volume loss, cysts, ventriculomegaly, thinning of the corpus callosum and delayed myelination, are associated with increased risk for adverse motor outcomes in early childhood (3-6). Thus, MRI scans performed at term equivalent postmenstrual age (PMA) in VPT infants may provide valuable prognostic information for clinicians (7, 8). However, conventional MRI studies may not fully elucidate the nature and extent of WMA.

Diffusion tensor imaging (DTI) also provides information on cerebral architecture, enabling quantification of white matter microstructural properties through the measurement of water displacement in the brain. For example, diffusivity values decrease significantly during brain maturation, reflecting changes in overall brain water content (9-11). Anisotropy measures provide information on white matter fiber bundle orientation and integrity, both of which are potentially disrupted in WMA. DTI measurements also enable calculation of quantitative diffusion tensor tractography (DTT), which has been used to detect subtle anatomic connectivity differences in white matter fiber bundles in preterm children (12). Across these measures, alterations in DTI parameters may reflect disruption in axonal and myelin microstructure (13).

It is important to note that while DTI provides information that differs from and is complementary to that obtained using conventional MRI, it is not entirely independent. The severity of white matter signal abnormalities on conventional MRI has been shown to correlate with regional DTI measures in VPT infants (14, 15), and changes in diffusion measures in the corticospinal tract (CST) and corpus callosum (CC) also correlate with qualitative WMA (16, 17). Additionally, DTT studies have revealed alterations in the development of the CST (12, 18) and CC (19, 20) in preterm adolescents and young adults with WMA.

The current study was aimed to determine whether bilateral moderate-severe WMA, detected on conventional MRI at term equivalent age in VPT children, can predict DTI measures at 7 years of age in two major tracts associated with motor function, the CST and CC. Because considerable variability exists in developmental outcomes for individuals with similar qualitative MRI lesions (21-24), we then investigated whether DTI measures at age 7 years differed between VPT children with and without motor impairment in subjects with bilateral moderate-severe WMA identified at term equivalent age.

Materials and Methods

Participants

Two hundred and twenty six VPT subjects (born <30 weeks' gestational age or <1250 g) with no genetic or congenital abnormalities were recruited for the Victorian Infant Brain
Study (VIBeS), a prospective, longitudinal cohort study conducted between 2001 and 2003 at the Royal Women's Hospital, Victoria, Australia. For this investigation, all subjects with either bilateral qualitative moderate-severe WMA or with no WMA on neonatal MRI were eligible.

Parental informed consent was obtained for each subject prior to participation in the study. The study was approved by the Royal Women's Hospital and Royal Children's Hospital Human Research Ethics Committees.

**Neonatal Magnetic Resonance Imaging**

For all participants, brain MRI was performed at term equivalent (37-42 weeks' PMA) using a 1.5T General Electric Signa LX Echospeed System MR scanner (Milwaukee, WI). Acquired sequences included 3D T1 spoiled gradient recalled (SPGR) images (0.8-1.6 mm coronal slices, flip angle 45°, TR 35 ms, TE 9 ms, FOV 210×157 mm², matrix 256×192) and T2 dual echo (interleaved acquisition) fast recovery fast spin echo images (1.7-3.0 mm coronal slices, TR 4000 ms, TE 60/160 ms, FOV 220×165 mm², matrix 256×192 interpolated to 512×512). Scanning was performed without sedation (25).

T1- and T2-weighted images were assessed for WMA according to a standardized scoring system previously validated in this population (3, 8). Severity of WMA was categorized according to total WMA score on a scale of 1-4 (1–none, 2–mild, 3–moderate, 4–severe) based on variables including presence and severity of white matter signal abnormality, white matter volume reduction, cystic white matter injury, ventriculomegaly, myelination of the posterior limb of the internal capsule (PLIC), and thinning of the CC. All scans were scored by a pediatric neurologist and a neonatologist blinded to clinical status. Inter-rater agreement was 96% (8). Figure 1 provides representative examples of subjects with moderate-severe WMA.

**7-Year-Old Magnetic Resonance Imaging**

At 7 years of age, all participants underwent repeat neuroimaging using a 3T Siemens scanner at the Royal Children's Hospital, Melbourne. Data acquisition included sagittal 3D turbo spin-echo T2-weighted images (T2W) (0.9 mm³ isotropic voxels, TR 3200 ms, TE 447 ms, FOV 240×215 mm², matrix 256×230), and a spin-echo echo-planar sequence with 25 non-collinear diffusion gradient directions (1.7 mm³ isotropic voxels, TR 12000 ms, TE 96 ms, FOV 250×250 mm², matrix 144×144). The b values for these 25 directions were chosen to linearly range from 0 to 1200 s/mm². This choice of b values differs from that used in most DTI experiments in which one or more b = 0 values are used and a constant high b value is used for all remaining directions. This choice is helpful in calculating more complicated and non-linear diffusion tensor models.

**Diffusion Tensor Tractography Analysis**

DTI and DTT computations followed the methods detailed in (26). The unsensitized (b = 0 s/mm²) DTI volume was registered to the T2W image with 12-parameter affine transform to compensate for EPI distortion. Internal head motion correction of the DTI data was performed using 9-parameter affine transform among groups of images with similar
sensitization. The tensor was calculated using log-linear regression (27). Measured diffusion parameters included fractional anisotropy (FA), mean diffusivity (MD), axial diffusivity (AD) and radial diffusivity (RD). Deterministic tracking algorithms and tract reconstructions were based on the streamline algorithm, i.e. by propagating along the local diffusion tensor principal eigenvector (26). The propagation increment was 0.5 mm, and the track termination criteria include FA <0.13 and a radius of curvature <1 mm. Full brain DTT on a 1 cubic mm grid was filtered by regions of interest (ROIs) manually placed perpendicular to the length of the tract based on anatomical landmarks in native space. Using a multi-ROI approach (28), the CST was identified using ROIs in the centrum semiovale (CSO) and the level of the ventral pontine crossing fibers (representative example provided in Figure 2A).

The medial portion of the full CC was tracked by placing a single ROI on a midline sagittal slice and ROIs on the sagittal view that were displaced 1 cm to the left and to the right from the midline slice. To ensure acceptable ROI placement, the callosal radiations were visualized in each subject. The filtered tracts were then clipped outside of the limiting ROIs and designated tracking thresholds were implemented. This approach identified the central fibers of each reconstructed tract (representative example provided in Figure 2B). Diffusion parameters were averaged over the reconstructed tract area, with FA, MD, AD and RD measured along each tract.

In order to confirm our DTT findings, whole brain analysis of the DTI data was also performed using the method of Tract Based Spatial Statistics (TBSS) (29). Using each individual's motion-corrected, aligned and averaged DTI data set, FA maps were used to create a target template for registration. FA images in the target group were aligned to each other and to a target image. The target image was the image that required the least amount of warping for all other FA images to align to it. The target image was affine-aligned into MNI152 standard space, and every FA image was then transformed into 1x1x1 mm³ MNI152 space. A mean FA image was calculated and used to produce the mean FA skeleton to represent the center of white matter tracts. FA images were projected onto the mean FA skeleton and thresholded at FA>0.20 for voxel-wise analyses. The registrations for all subjects were visually inspected for accuracy, and 3 of the subjects from the moderate to severe WMA group were removed due to poor registration. FA, MD, AD and RD images were also calculated and all were analyzed with Randomise, a permutation-based multiple comparisons corrected statistical approach, to evaluate for differences between subjects with moderate-severe WMA and those without WMA.

7-Year-Old Developmental Testing

At 7 years of age, motor functioning was assessed with the Movement Assessment Battery for Children, 2nd Edition (MABC2) (30). This tool provides a standardized assessment of both gross and fine motor development. Children were classified as having impaired motor performance if their total motor score was at or below the 15th-centile based on the test norms. The hemisphere contralateral to the hand the child used to write during the 7-year follow-up assessment was designated as dominant.
**Statistical Analyses**

Linear mixed models with subject ID nested within white matter injury (or motor impairment) as a random effect were used to assess the differences in diffusion parameters across groups. Study subject ID was also included as a random variable to account for the repeated measure design. Additional multivariable analyses controlling for gender, PMA at birth, age at scan, and handedness in the CST were conducted. Due to the number of diffusion measures that were simultaneously analyzed in the four regions, Bonferroni multiple comparisons correction was applied with a level of significance determined at $p \leq 0.01$.

**Results**

A total of 62 7-year-old VPT subjects were included in this study, including 42 subjects without WMA and 20 subjects with bilateral moderate-severe WMA on neonatal MRI scan. The remaining 164 subjects were ineligible for the following reasons: death (n=2), incomplete data at age 7 years (n=48), or mild WMA or unilateral injury (n=114). There were no demographic or clinical differences between included and excluded subjects (data not shown). The median PMA at birth for included subjects was 27.3 weeks (range 22-32 weeks) and median birthweight was 937 grams (range 414-1425 grams). Left-handedness was reported in 16.7% of the subjects without WMA and 10% of the WMA group. For the infants with bilateral moderate-severe WMA, four subjects had cystic periventricular leukomalacia (PVL). A total of five subjects had lesions in the area of the CST. No infants had evidence of injury in the region of the PLIC and a single subject had asymmetric myelination of the PLIC. Additional group demographic information is provided in Table 1.

At 7 years of age, VPT children with moderate-severe WMA on qualitative scoring of term equivalent MRI demonstrated lower FA and higher MD and RD in the corpus callosum compared to VPT children who had no WMA on neonatal MRI. The right CST also displayed higher MD and AD in children with neonatal WMA. Notably, there were no differences between groups in the left CST (Table 2).

In relation to motor outcomes, of the 20 VPT infants who were categorized with moderate-severe WMA at term equivalent, there were 12 VPT children with no impairment on the MABC2 and 8 VPT children with impairment at 7 years of age. The only value that differed between groups, when including Bonferroni correction for multiple comparisons, was lower FA in the corpus callosum in VPT children with impaired motor ability at age 7 years. There was no difference in the diffusion measures within the CST (Table 3).

Comparison of diffusion parameters between VPT children with and without motor impairment at 7 years of age (irrespective of WMA status during infancy) demonstrated no differences between groups (data not shown).

In the TBSS analysis (Figure 3), the CC demonstrated lower FA values in the moderate-severe WMA group (n=17, after removal of 3 subjects with poor registration) as compared to subjects without WMA (n=42) (areas colored yellow and red on the FA skeleton in Figure 3A). A similar pattern was identified with RD, with greater values in the WMA group.
No differences in MD or AD were identified between groups. In addition, no group differences were identified for any of the CST parameters.

Discussion

There are two key findings utilizing DTT in this study. The first is that, as previously reported, VPT born children with neonatal qualitative moderate-severe white matter abnormalities displayed alterations in diffusion measures within the CST and CC tracts at 7 years of age compared to those without qualitative neonatal WMA. The second key finding, complementary to prior studies using WMA alone (8), is that 40% of VPT born children with qualitative neonatal moderate-severe WMA (8/20) experience significant motor impairment at 7 years of age, and such impairment is associated with abnormal diffusion measures in the CC compared with non-impaired children with similar patterns of qualitative neonatal bilateral white matter injury. These findings highlight that the perturbations in cerebral microstructure of the CST and CC tracts that occur secondary to neonatal moderate-severe WMA persist and may differentiate VPT children with motor impairments in childhood.

Diffusion Measures affected by WMA

Corticospinal Tract—This study demonstrated that VPT children with moderate-severe WMA demonstrated increased diffusivity measures in the right CST compared to those without, indicative of abnormal microstructural organization. This pattern of injury has been hypothesized to reflect disrupted fiber tracts, less myelin and/or lower axonal density. This premise is consistent with animal models indicating a trophic role for oligodendrocytes in axonal development in rabbit (31) and rat embryonic cortical neurons (32) and disruption in pre-oligodendrocyte maturation following WMA in a neonatal rat model of hypoxic-ischemic injury (33). This pattern of injury has also been reported in a mouse model of multiple sclerosis (34).

The CST is one of the earliest areas of the brain to undergo myelination, and therefore may be particularly vulnerable to injury. Prior investigation has shown the CST to be susceptible to the effects of WMA, with abnormal anisotropy and diffusivity measures reported in infants with PVL (35) and extensive white matter signal abnormalities (14). Furthermore, it is located within the regions dorsal and lateral to the lateral ventricles, areas known to be susceptible to white matter injury. The current investigation demonstrated that these effects persist beyond infancy and into childhood (although this was noted on the right side only as discussed below).

Corpus Callosum—Commissural fibers such as the CC have also been shown to be susceptible to the effects of WMA in VPT born children (36). The current study supports these findings, as the FA was reduced and diffusivity values were increased in children with moderate-severe WMA. These findings in the CC were also confirmed by the TBSS analysis (Figure 3). In comparison to the corticospinal pathways, the corpus callosum matures relatively late, continuing to show growth secondary to myelination of interhemispheric pathways through adolescence. The susceptibility of this tract to complications associated with prematurity has been well established. Yeatman (37) demonstrated alterations in
callosal diffusion correlated with early WMA. In addition, Fan (35) demonstrated children with PVL had reduced FA in the corpus callosum. In this investigation, children with early bilateral moderate-severe WMA had decreased FA in the CC in comparison to children without WMA, a pattern previously reported in delayed or disrupted myelination (38, 39). We have previously shown that altered microstructural organization of the CC in VPT infants is associated with WMA at term equivalent (40), with the current study indicating this association persists into childhood.

**Motor Impairment**

The study demonstrated lower FA in the CC of subjects with moderate-severe WMA and motor impairment. These findings are consistent with studies investigating the relationship between diffusion measures of the CC and motor outcome in PT populations. Counsell and Skranes reported FA within the posterior body and isthmus of the CC correlated with neurodevelopmental ability in preterm children and young adults (38, 39). Rose reported reduced FA in the splenium of the CC in VPT infants with abnormal neurodevelopment (41). In addition, we previously found that higher MD and RD within the splenium of VPT infant was related to worse motor functioning at 2 years of age (40). These findings support that the CC plays an important function in differential motor ability in VPT born children with moderate-severe WMA and supports further investigative focus in this area.

**Hemispheric Asymmetry**

A prior study by Hasan (42) demonstrated DTT asymmetry of the uncinate fasciculus across the lifespan, with elevation of FA in the left CST compared to the right in a group of right handed subjects. A study by Lebel and Beaulieu (43) advanced this premise by demonstrating asymmetry in the arcuate fasciculus and correlating it with cognitive language skills in typically developing children. Both studies imply that diffusion asymmetry is related to remodeling due to asymmetric activity in right-handed subjects.

The findings in the CST of neonates with WMA were confined to the right (non-dominant) side of the brain. This potentially reflects asymmetric recovery due to differences in activity on the dominant side of the brain. It is also plausible that functional adaptation for motor performance is dependent on adequate compensation involving the non-dominant hemisphere. Subsequently, if this area is also affected, the potential for positive adaptation is limited. The current investigation was unable to test this hypothesis in the neonatal period due to limitations in the DTI acquisition, but it would be worthy of further study in preterm infants at term in relation to motor development.

**Strengths and Limitations**

This study benefits from its longitudinal design, enabling exploration of the relationship between MRI results from the neonatal period with neuroimaging and neurodevelopmental outcome measures at 7 years of age. However, the study focused on a specific subset of subjects from a larger cohort, which limited subject numbers available for subgroup analyses. Further studies of children with unilateral injury and those with less severe WMA are needed. In addition, subjects were studied during a period where heterogeneity may exist due to differing rates of individual development, introducing an additional confound into the
interpretation of the results. Further follow-up of these and other cohorts during later developmental periods is warranted.

There are also technical limitations inherent to DTT that should be noted. When applying this technique, there is no measure indicative of the confidence in the reconstructed diffusion trajectories. Nonetheless, for regions where there is a single dominant fiber pathway (such as the CST and CC), uncertainty in the principal diffusion direction is small.

Conclusions

WMA on qualitative conventional MRI at term equivalent is correlated with persistent microstructural effects at 7 years of age and associated with impaired motor outcome. This implies that white matter pathways play a critical role in the pathophysiology of motor disability. The study results also suggest developmental adaptation to injury, with microstructural differences identified in subjects with normal outcomes despite significant neonatal WMA. This highlights the plasticity inherent in this population, and represents a possible marker for future therapeutic interventions.

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Figure 1. WMA in preterm infants scanned at term equivalent PMA
Axial and coronal T2-weighted MR images illustrating representative example of subjects without WMA (A), with moderate WMA (B), and with severe WMA (C) (arrows denote representative areas of WMA).
Figure 2. ROI-based tractography for VPT children scanned at 7 years of age
Images illustrating diffusion tensor tractography results for the (A) corticospinal tract and (B) corpus callosum for a 7-year-old subject without WMA. Results overlaid on diffusion weighted image.
Figure 3. TBSS results comparing VPT children with and without WMA
TBSS results denoting areas of the CC that differed significantly between VPT children scanned at 7 years of age with and without WMA on neonatal MRI. Results demonstrate (A) higher FA and lower RD (B) in the CC in subjects without WMA. Yellow color denotes areas that differ between groups (p ≤0.05). Green color indicates areas with no difference. Results overlaid on averaged FA map.
# Table 1

**Group Characteristics for 7-year-old VPT children with and without moderate-severe WMA**

| DEMOGRAPHICS                      | Mod-Severe WMA | No WMA          | Total N=40 |
|-----------------------------------|----------------|-----------------|------------|
|                                   | Total N=20     | Impaired MABC2 N=8 | Normal MABC2 N=12 | |
| Males:Females                     | 11:9           | 3:5             | 8:4        | 23:19       |
| Median PMA at birth (weeks)       | 26.6 (22-29.6) | 26.5 (23-29.4)  | 26.6 (22-29.6) | 27.9 (24-32) |
| Median Birthweight (grams)        | 892 (414-1395) | 848 (480-1354)  | 899 (414-1395) | 975 (560-1425) |
| IUGR                              | 1 (5%)         | 0 (0%)          | 1 (8%)     | 3 (7%)      |
| Multiple Gestation                | 8 (40%)        | 3 (38%)         | 5 (42%)    | 20 (48%)    |
| NEONATAL COURSE                   |                |                 |            |            |
| Mean Length of IPPV (days)        | 19.4 ± 27.1    | 20.2 ± 22.9     | 18.9 ± 30.5 | 7.2 ± 12.3 |
| Postnatal Corticosteroids         | 2 (10%)        | 0 (0%)          | 2 (17%)    | 2 (5%)     |
| Necrotizing Enterocolitis         | 3 (15%)        | 0 (0%)          | 3 (25%)    | 4 (10%)    |
| Sepsis                            | 7 (35%)        | 4 (50%)         | 3 (25%)    | 6 (14%)    |
| NEONATAL INJURY                   |                |                 |            |            |
| Mean IVH grade (cranial ultrasound)| 0.7 ± 1.5      | 1.4 ± 1.9       | 0.25 ± 0.9 | 0.1 ± 0.5  |
| Mean WMA score (MRI)              | 3.0 ± 0.0      | 3.0 ± 0.0       | 3.0 ± 0.0  | 1.0 ± 0.0  |
| 7 YEAR OLD OUTCOME                |                |                 |            |            |
| Handedness (left:right:mixed)     | 2:18:0         | 2:6:0           | 0:12:0     | 7:33:2     |
| MABC2 Total Standard Score        | 6.8 ± 2.9      | 3.8 ± 1.0       | 8.8 ± 1.7  | 9.4 ± 3.3  |
| MABC2 Percentile                  | 22.6 ± 23.2%   | 2.3% ± 1.8%     | 36.1 ± 20.1% | 44.1 ± 29.2% |
Table 2
Diffusion parameters for 7-year-old VPT children with and without moderate-severe WMA

| Parameter | Tract               | Mod-Severe WMA n=20 | No WMA n=42 | p-value<sup>a</sup>* |
|-----------|---------------------|---------------------|--------------|-----------------------|
| FA        | Left CST            | 0.64 ± 0.03         | 0.64 ± 0.03  | 0.95                  |
|           | Right CST           | 0.57 ± 0.03         | 0.58 ± 0.03  | 0.54                  |
|           | Corpus Callosum     | 0.65 ± 0.03         | 0.68 ± 0.02  | 0.0002                |
| MD        | Left CST            | 0.76 ± 0.02         | 0.76 ± 0.02  | 0.44                  |
|           | Right CST           | 0.90 ± 0.07         | 0.85 ± 0.06  | 0.008                 |
|           | Corpus Callosum     | 1.07 ± 0.08         | 1.02 ± 0.07  | 0.008                 |
| AD        | Left CST            | 1.43 ± 0.04         | 1.42 ± 0.04  | 0.43                  |
|           | Right CST           | 1.53 ± 0.10         | 1.46 ± 0.07  | 0.009                 |
|           | Corpus Callosum     | 1.90 ± 0.10         | 1.88 ± 0.07  | 0.50                  |
| RD        | Left CST            | 0.43 ± 0.03         | 0.43 ± 0.03  | 0.74                  |
|           | Right CST           | 0.59 ± 0.07         | 0.54 ± 0.06  | 0.02                  |
|           | Corpus Callosum     | 0.66 ± 0.08         | 0.59 ± 0.07  | 0.0007                |

Abbreviations: Fractional Anisotropy (FA), Mean Diffusivity (MD), Axial Diffusivity (AD), Radial Diffusivity (RD).

FA is dimensionless. Units for MD, AD and RD are 10⁻³ mm²/s.

<sup>a</sup> Adjusted for gender, gestational age at birth and age at scan for all tracts. Additional adjustment for handedness was done only in the CST tracts.

<sup>*</sup> Bold text denotes significant between group differences following multiple comparisons correction with a level of significance determined at p ≤ 0.01.

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### Table 3
Diffusion parameters for 7-year-old VPT children with moderate-severe WMA categorized by motor outcome

| Parameter | Tract          | Impaired MABC2 n=8 | Normal MABC2 n=12 | p-value<sup>a</sup> |
|-----------|----------------|--------------------|-------------------|---------------------|
| FA        | Left CST       | 0.65 ± 0.03        | 0.64 ± 0.02       | 0.71                |
|           | Right CST      | 0.59 ± 0.02        | 0.56 ± 0.03       | 0.16                |
|           | Corpus Callosum| 0.63 ± 0.03        | 0.66 ± 0.02       | 0.01                |
| MD        | Left CST       | 0.76 ± 0.01        | 0.76 ± 0.03       | 0.89                |
|           | Right CST      | 0.91 ± 0.05        | 0.90 ± 0.08       | 0.63                |
|           | Corpus Callosum| 1.09 ± 0.10        | 1.06 ± 0.06       | 0.75                |
| AD        | Left CST       | 1.44 ± 0.05        | 1.42 ± 0.04       | 0.68                |
|           | Right CST      | 1.57 ± 0.08        | 1.51 ± 0.10       | 0.80                |
|           | Corpus Callosum| 1.89 ± 0.11        | 1.91 ± 0.09       | 0.27                |
| RD        | Left CST       | 0.43 ± 0.02        | 0.43 ± 0.03       | 0.67                |
|           | Right CST      | 0.58 ± 0.04        | 0.59 ± 0.08       | 0.37                |
|           | Corpus Callosum| 0.69 ± 0.10        | 0.64 ± 0.05       | 0.17                |

Abbreviations: Fractional Anisotropy (FA), Mean Diffusivity (MD), Axial Diffusivity (AD), Radial Diffusivity (RD).

FA is dimensionless. Units for MD, AD and RD are $10^{-3}$ mm$^2$/s.

<sup>a</sup> Adjusted for gender, gestational age at birth and age at scan for all tracts. Additional adjustment for handedness was done only in the CST tracts.

<sup>*</sup> Bold text denotes significant between group differences following multiple comparisons correction with a level of significance determined at $p \leq 0.01$. 