Longitudinal Changes in the Corpus Callosum Following Pediatric Traumatic Brain Injury as Assessed by Volumetric MRI and Diffusion Tensor Imaging

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Longitudinal Changes In The Corpus Callosum Following Pediatric Traumatic Brain Injury

Trevor C. Wu

A dissertation submitted to the faculty of Brigham Young University in partial fulfillment of the requirements for the degree of Doctor of Philosophy

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ABSTRACT

Longitudinal Changes In The Corpus Callosum Following Pediatric Traumatic Brain Injury

Trevor C. Wu

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Doctor of Philosophy

Atrophy of the corpus callosum (CC) is a documented consequence of moderate-to-severe traumatic brain injury (TBI), which has been expressed as volume loss using quantitative magnetic resonance imaging (MRI). Other advanced imaging modalities such as diffusion tensor imaging (DTI) have also detected white matter microstructural alteration following TBI in the CC. The manner and degree to which macrostructural changes such as volume and microstructural changes develop over time following pediatric TBI and their relation to a measure of processing speed is the focus of this longitudinal investigation. As such, DTI and volumetric changes of the CC in participants with TBI and a comparison group at approximately three and 18 months post injury and their relation to processing speed were determined.

Keywords: corpus callosum, pediatrics traumatic brain injury, diffusion tensor imaging
ACKNOWLEDGMENTS

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Longitudinal Changes In The Corpus Callosum Following Pediatric Traumatic Brain Injury

Traumatic brain injury (TBI) is a major public health concern. According to Langlois and colleagues (Langlois, Rutland-Brown, & Thomas, 2005), approximately 475,000 children are treated for TBI annually in the United States, with the highest rate of hospitalization in the adolescent age range (129/100,000). Closed head trauma arising from falls and motor vehicle crashes is the most frequent cause of unintentional pediatric TBI, while penetrating brain injury accounts for less than five percent of TBI in the pediatric population. Children and adolescents are particularly vulnerable to closed head trauma (Thurman, Coronado, & Selassie, 2007), and TBI sustained at this critical developmental stage can have a profound negative impact on learning, behavior and psychological outcome (Yuan, et al., 2007).

Neuropathology associated with TBI

The neuropathology associated with TBI has been traditionally classified as either focal or diffuse injury. Focal injury, such as cerebral contusions or intracranial hemorrhage, can be readily identified on computed tomography (CT) or magnetic resonance imaging (MRI) (Benson, et al., 2007; Povlishock & Katz, 2005; Smith, Meaney, & Shull, 2003). In contrast, diffuse injury is usually less evident on conventional imaging. Recent studies have demonstrated the sensitivity and utility of diffusion tensor imaging (DTI) in identifying and characterizing microstructural pathologies, including those occurring from TBI (Alexander, Lee, Lazar, & Field, 2007; Arfanakis, et al., 2002; Ashwal, Holshouser, & Tong, 2006; Chu, et al., 2010; Ewing-Cobbs, Hasan, Prasad, Kramer, & Bachevalier, 2006; Mayer, et al., 2010; Nakayama, et al., 2006; Rugg-Gunn, Symms, Barker, Greenwood, & Duncan, 2001;
Salmond, et al., 2006; Wilde, et al., 2006; Wilde, et al., 2008; Wu, et al., 2010; Xu, Rasmussen, Lagopoulos, & Haberg, 2007).

**Diffusion Tensor Imaging**

Diffusion tensor imaging is based on two theoretical assumptions: water molecules in the brain are constantly moving, and the directionality and magnitude of diffusivities of water molecules are indicative of the integrity of underlying neuronal structures (Alexander, et al., 2007; Mori & Zhang, 2006; Mukherjee, Berman, Chung, Hess, & Henry, 2008). Healthy axonal membranes constrain water molecules intracellularly and are closely compacted in tracts minimizing free water in extracellular spaces as well (Yamada, Sakai, Akazawa, Yuen, & T., 2009). The DTI technique capitalizes upon this combination of constraining water diffusivity within and around the axon where aggregate white matter tracts can be inferred by linking similar characteristics of water diffusivity (Yamada, et al., 2009). This also means that when an axon is damaged water diffusion is altered which can be detected by various DTI metrics, including fractional anisotropy (FA) and apparent diffusion coefficient (ADC). FA measures the degree to which diffusion occurs in one particular direction, and is indicated by a coefficient ranging from 0 to 1, where 0 is completely random and 1 is maximally anisotropic. ADC reflects the overall magnitude of diffusivity. A small ADC represents greater constrained diffusivity. Neuronal swelling, cytoskeleton, or alterations of extracellular space secondary to injury and edema, often seen immediately following mild TBI, have been shown to disrupt normal diffusivities, resulting in increased FA and decreased ADC (Bazarian, et al., 2007; Chu, et al., 2010; Mayer, et al., 2010; Wilde, et al., 2008; Wu, et al., 2010). It is not clear whether the initial increase in FA is limited to mild TBI or whether this finding reflects the feasibility of performing DTI during the subacute
phase of mild injury. In contrast, during axonal degeneration following TBI, water diffusivity increases, which reduces FA and increases ADC (Alexander, et al., 2007; Assaf, Beit-Yannai, Shohami, Berman, & Chohen, 1997; Belanger, Vanderploeg, Curtiss, & Warden, 2007; Benson, et al., 2007; Ducreus, et al., 2005; Newcombe, et al., 2007; Povlishock & Katz, 2005; Sidaros, et al., 2008; Wilde, et al., 2008; Xu, et al., 2007), which relates to histologically identified axonal pathology (Mac Donald, Dikranian, Bayly, Holtzman, & Brody, 2007; Mac Donald, Dikranian, Song, et al., 2007; S. Wang, et al., 2009).

**Microstructural Alterations Following TBI**

Following TBI, several axonal pathways have robustly demonstrated altered diffusivities including the corpus callosum (CC), anterior limb of the internal capsule, cingulum bundles, uncinate fasciculi, centrum semiovale, thalamic white matter projections and inferior and superior longitudinal fascicule though results vary across ages, amount of time post-injury and severities of injury (Arfanakis, et al., 2002; Babikian, et al., 2010; Bendlin, et al., 2008; Huisman, et al., 2004; Inglese, et al., 2005; Kumar, et al., 2010; Levin, et al., 2008; Little, et al., 2010; Nakayama, et al., 2006; Niogi, et al., 2008; Ptak, et al., 2003; Rugg-Gunn, et al., 2001; Sidaros, et al., 2008; J. Wang, Bakhadirov, Devous, Adbdi, & McColl, 2008; Wilde, et al., 2006; Wilde, et al., 2008; Xu, et al., 2007). Among these pathways, the CC is selectively vulnerable where callosal damage is common in children and adolescents with TBI (Beauchamp, et al., 2009; Ewing-Cobbs, et al., 2008). The long coursing nature and the midline location adjacent to the falx cerebri make CC particularly susceptible to the traumatic axonal injury (TAI) resulting from shear-strain forces. Decreased FA and increased ADC (Bendlin, et al., 2008; Ewing-Cobbs, et al., 2008; Gorrie, Duflou, Brown, Gibson, & Waite, 2001; Kraus, et al., 2007; Levin, et al., 2008; Wilde, et al.,
and volumetric reduction (Anderson & Bigler, 1994; Benavidez, et al., 1999; Kim, et al., 2008; Levin, et al., 1990; Tomaiuolo, et al., 2004; Wilde, et al., 2006) in the CC following TBI are consistently documented in children and adolescents during the subacute to chronic stage. However, to date, no study has systematically examined how volumetric changes in the CC relate to changes in DTI metrics in a longitudinal design.

Given the widespread interhemispheric connections that course through the CC and its primary function in mediating interhemispheric communication (Pannek, et al., 2010), callosal pathway disruptions can have a profound impact on cognitive functioning, including the speed of information processing (Aukema, et al., 2009; Biegon, Eberling, & Richardson, 1994; Chepuri, et al., 2002; Jokinen, Ryberg, & Halska, 2007; Turken, et al., 2008). Patients with TBI often demonstrate slowed information processing on neuropsychological testing (Battistone, Woltz, & Clark, 2008; Johansson, Berglund, & Ronnback, 2009; Madigan, DeLuca, Diamond, Tramontano, & Averiall, 2000). In a previous cross-sectional study, we have found a significant relation between speed of processing and CC integrity as measured by DTI FA in children and adolescents with TBI at three months post-injury (H. S. Levin, et al., 2008).

**Hypothesis**

In the present study, we examined longitudinal changes in the CC volume and microstructural integrity from three to 18 months post injury in a group of children and adolescents with complicated mild and moderate to severe TBI and a comparison group of children and adolescents with orthopedic injury (OI), and the relation between these imaging findings and speed of information processing. We hypothesized that the OI group would have higher FA and lower ADC in the total CC and its sub-regions (genu/anterior,
body/central and splenium/posterior) at three months post injury and these differences would become more pronounced at 18 months as a result of either the disruption of a normal developmental trajectory or continued degeneration. We anticipated that the TBI group would demonstrate slower cognitive processing speed than the OI group.

Method

Participants

This study was approved by the Institutional Review Board of all institutions from which data were collected. Informed consent was obtained from each participant or his/her legal guardian as appropriate. All participants were recruited from consecutive admissions to emergency rooms at Level-1 trauma centers in Houston, Dallas, and Miami. All participants were fluent in English and were between the ages of seven and 17 years at the time of study enrollment. As part of the study design, magnetic resonance imaging (MRI) and cognitive assessments were planned for three and 18 months post-injury.

The TBI group was comprised of 23 participants (15 males and 8 females) between the ages of 7.8 and 17.2 years ($M = 12.9 \pm 3.2$) who sustained either complicated mild or moderate-to-severe TBI. The severity of the injury was determined by the postresuscitation Glasgow Coma Scale (GCS) scores (Teasdale & Jennett, 1974). Severe TBI was defined as a GCS score of 3 – 8, and moderate TBI as 9 – 12. A complicated mild TBI was defined as a GCS score of 13 – 15 with acute (within 24 hours post-injury) trauma-related CT findings (e.g., contusions and hematomas). Participants with complicated mild TBI were recruited because the presence of cerebral lesions is predictive of poor cognitive outcome at 12 month post-injury (Levin, et al., 2008). Given these classifications, there were 16 severe, 4 moderate and three complicated mild cases. Other eligibility criteria included Abbreviated
Injury Scale (AIS) (Committee on Injury Scaling, 1998) score of < 4 for body parts other than the head and an Injury Severity Scale ISS score of < 12. The modal mechanisms of injury were motor-vehicle or motorcycle crashes. All but one TBI participants were right-hand dominant.

The OI group included 25 participants (18 males and 7 females), ages ranging 7.1 – 16.3 years (\(M \ 11.8 \pm 2.7\)) who were hospitalized overnight, generally for upper or lower extremity bone fractures. The OI comparison group controls for risk factors that predispose individuals to injury, including pre-existing behavioral problems, subtle learning disabilities, and family variables. The absence of significant previous head trauma in the OI group was confirmed through a detailed developmental questionnaire administered to the parent or legal guardian, and the absence of concurrent head injury was confirmed through medical records and/or physician report of relevant history and physical examination findings and, when available, clinical imaging results (i.e., negative CT). OI in this cohort primarily resulted from sports and play-related injuries and falls. All but four controls were right-hand dominant.

Potential participants in both groups were excluded if there was evidence of previous head injury, child abuse, pre-existing neurologic disorders (e.g., cerebral palsy, mental retardation, and epilepsy) or diagnosed learning disabilities, pre-existing severe psychiatric disorders (e.g., schizophrenia, or pervasive developmental disorders), prematurity or low birth weight (birth weight < 2500 grams and gestation < 37 weeks), penetrating gunshot wounds to the brain, contraindication to undergoing MRI, hypoxia (PO\(_2\) < 96mmHg for age 6 – 16 years) or hypotension (systolic blood pressure 2 standard deviations below the mean for the age group).
Imaging

All participants underwent MRI without sedation on Philips 1.5 Tesla Intera scanners (Netsch, 2001) at Texas Children’s Hospital (Houston), the Rogers MRI Center, University of Texas Southwestern Medical Center (Dallas) or the Miami Children’s Hospital (Miami) using comparable platforms and software. Regular quality assurance testing was performed on all three scanners including American College of Radiology phantom and Weisskoff testing for echo planar imaging (EPI) sequences, and all scanners were consistently noted to be within an acceptable range throughout the study.

DTI Acquisition

Transverse multislice spin echo, single-shot, EPI sequences (10,150.5 ms repetition time (TR), 90 ms echo time (TE), 2.7 mm thick slices with 0 mm gap) were applied. A 256 mm FOV (RFOV = 100%) was used with a measured voxel size of 2.69 × 2.69 × 2.7 mm. Diffusivities will be evaluated along 15 directions (number of b-value = 2, low b-value = 0, and high-b value = 860 s/mm²). Two acquisitions of high-b images were obtained and averaged to ensure better signal-to-noise ratio. A total number of 55 slices were acquired, and each acquisition took approximately 6 minutes.

Volumetric Acquisition

T1-weighted 3D sagittal acquisition series were used for volumetric analysis. Parameters included 15ms TR, 4.6 ms TE, 1.0 mm thick slices with 0 mm gap). A 256 mm FOV (RFOV = 100%) was used with a voxel size of 1.0 × 1.0 × 1.0 mm.

DTI Analysis

Prior to computing FA maps with the Philips fiber tracking 4.1V3 Beta 2 software, shear and eddy current distortion and head motion artifact were corrected by using the Philips
PRIDE-registration tool (Netsch, 2001). ROIs of the CC were manually traced on the mid-sagittal plane following a previously published protocol (Wilde, et al., 2006) and included the genu/anterior, body/central and splenium/posterior sub-regions as well as the total CC. The automated Philips three-dimensional fiber tracking program was utilized to examine fiber tracks passing through the selected regions in the CC. The mean of FA and ADC, automatically calculated by the software, was used to quantitate DTI variables. The algorithm for fiber tracking is based upon the Fiber Assignment by Continuous Tracking (FACT) method (Mori, Crain, Chacko, & Vanzijl, 1999). If the FA in the voxels was less than 0.2 or if the angle between adjacent voxels was greater than 7 degrees, fiber tracking terminated.

**Intra-rater and Inter-rater Reliability**

DTI analysis was performed by two experienced raters following the specified protocol. To ensure intra-rater reliability, each ROI was independently analyzed by each rater twice. To examine inter-rater agreement, two trained raters performed analysis on each sub-region in a sample of six participants randomly selected (three TBI patients and three controls). Shrout-Fleiss intra-class correlation coefficients were performed and revealed satisfactory intra- (range: 0.937-1.000) and inter-rater reliability (range: 0.913-0.976)

**Volumetric Analysis**

Volumetric segmentation was performed with the FreeSurfer image analysis suite version 4.5.0 (Athinoula A. Martinos Center for Biomedical Imaging, 2005; [http://surfer.nmr.mgh.harvard.edu/](http://surfer.nmr.mgh.harvard.edu/)). In order to reduce bias with respect to a particular time point, the longitudinal processing scheme was initialized with a template based on the cross-sectionally processed scans at both three and 18 months. Details of the cross-sectional
(Fischl, et al., 2004) and longitudinal (Morey, et al., 2010) procedures have been described previously. Briefly, the cross-sectional processing involves intensity normalization (Sled, Zijdenbos, & Evans, 1998), automated Talairach transformation, the removal of non-brain tissue using a hybrid watershed/surface deformation procedure, and segmentation and labeling of the subcortical white matter and deep gray matter volumetric structures (Fischl, et al., 2002; Fischl, et al., 2004). Each scan was aligned to the unbiased template space, and the segmentation of the template was used as an initial approximation for the segmentation of each time point in the longitudinal processing. The use of the unbiased template for initializing the processing of cases in a longitudinal series is intended to reduce the random variation in the longitudinal procedure and improve the robustness and sensitivity of the overall longitudinal analysis, as stated by the Freesurfer developers. Results for each subject were visually inspected to ensure accuracy of registration, skull stripping, and segmentation. No editing was necessary in the automated segmentation of subcortical structures. The five standard subregions of the corpus callosum generated by FreeSurfer were used in analysis (anterior, mid-anterior, central, mid-posterior, and posterior) as well as a composite for the total corpus callosum, which was the sum of the volume of each of these sub-regions. The corpus callosum volumes derived in FreeSurfer are considered true volume estimates that compose the entire structure, with partial volume correction in order to account for the fraction of the structure in each border voxel. The anterior, central and posterior sub-regions used in volumetric analysis corresponded to genu/anterior, body/central and splenium/posterior sub-regions (respectively) used in DTI analysis.

**Cognitive Measures**

The baseline condition of the Flanker Arrow Test (Bunge, Dudukovic, Thomason,
Vaidya, & Gabrieli, 2002; Eriksen & Eriksen, 1974) was selected as a measure of information processing speed because we found that this measure of visual choice reaction time (without distraction) was related to corpus callosum integrity as measured by DTI variables (Levin, et al., 2008; Wilde, et al., 2006). One advantage of using this particular cognitive measure is that it is a language-free and multi-trial task with a high reliability (split-half $r = 0.96$) in the TBI population. Missing data were due to technical problems with this computerized task, the child’s refusal to complete the task, or time constraints of the parent/guardian which precluded administration. There did not appear to be any source of systematic bias for the subjects with missing data such as greater injury severity (i.e., GCS) or cognitive impairment, and data was missing from all three sites. In the baseline condition of this task, a horizontal central arrow pointing to the left or to the right appeared on each trial. The child was asked to quickly press the button on the right or the left consistent with the direction that the arrow was pointing. There were 28 trials of the baseline task condition. Reaction time (RT) in each condition was the performance measure.

**Data Analysis**

Demographic statistics between groups were compared using t tests for continuous variables, namely, 1) age at injury, 2) Socioeconomic Composite Index (SCI) $z$-score, a measure of socioeconomic status (Yeates, et al., 1997), 3) maternal education, and 4) time post-injury for both the three and 18-month assessment time points. Categorical variables were analyzed using $\chi^2$ (for gender) or Fisher’s exact test (for handedness, race/ethnicity distribution, and mechanism of injury). We utilized $t$-tests to examine the group differences in FA and ADC of the CC sub-regions and total CC at both three and 18 months. Group differences in CC sub-region and total CC volumes at three and 18 months were also
analyzed with general linear models, including total intracranial volume (TICV) as a
covariate to account for differences in head size. To examine the within-group changes over
time in FA, ADC, and volumes of the CC sub-regions and total CC, paired t-tests were used
for each group and each sub-region on the difference score, as calculated by three month
minus 18 month values. A two-sample t-test was used to test the group difference on the
change (two-way interaction of group by occasion). To examine the possibility that
complicated-mild injury may differ in degree from more severe injuries on DTI, we repeated
analyses excluding the participants with complicated-mild TBI and found that this did not
change the results of the analyses. Within-group differences on the Flanker baseline reaction
time (RT) between three and 18 months were tested using a paired t-test of the difference
score between RT at three and 18 months. Pearson’s partial correlation coefficients with age
at injury and SCI controlled were used to examine the relation of DTI and volumetric
variables to performance on Flanker baseline RT at both three and 18 months. The
significance level was adjusted by using the family-wise Bonferroni method for the sub-
regions (three sub-regions for DTI, alpha = 0.017, and five sub-regions for volumetrics, alpha
= 0.01).

Results

Demographic and Injury Characteristics

The TBI group did not significantly differ from the OI group in age, gender,
handedness, SCI, maternal education, or the duration of the post-injury interval at either the
three- or 18-month assessment interval. As expected, groups differed significantly in the
mechanism of injury, with more participants in the TBI group sustaining injury via motor
vehicle and motorcycle crashes, and more participants in the OI group sustaining injury via
sports- and play-related injuries and falls. T-values (or $\chi^2$ or Fisher exact test values) and $p$-values are presented in Table 1.

**Group Differences at Three and 18 Months Post-Injury for DTI**

At three months post-injury, the FA of the genu/anterior, body/central, splenium/posterior and total CC differed significantly between groups, with the TBI group demonstrating decreased FA. Similarly, ADC for the genu/anterior, splenium/posterior and total CC differed significantly between groups, with the TBI group demonstrating increased ADC. The ADC for the CC body/central sub-region exhibited a positive trend, but this was not considered significant after correction for multiple comparisons.

At the 18 month post-injury assessment, the FA for the genu/anterior and splenium/posterior sub-regions and the ADC for the genu/anterior and the body/central sub-regions and total CC differed significantly between groups, with lower FA and higher ADC in the TBI group. FA for the body/central sub-region and total CC as well as ADC for the splenium/posterior sub-region demonstrated trends after Bonferroni correction. Tables 2-3 detail means, standard deviations, t-values and $p$-values for all CC regions at both time points.
Table 1

*Demographic and Injury Characteristics of TBI and OI Groups*

<table>
<thead>
<tr>
<th>Demographics</th>
<th>TBI Group (n=23), Mean ± SD</th>
<th>OI Group (n=25), Mean ± SD</th>
<th>t-value / χ² value</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at Injury (years)</td>
<td>12.9 ± 3.2, range = 7.8 – 17.2</td>
<td>11.8 ± 2.7, range = 7.1 – 16.3</td>
<td>-1.20</td>
<td>0.235</td>
</tr>
<tr>
<td>Gender distribution</td>
<td>8 female, 15 male</td>
<td>7 female, 18 male</td>
<td>0.26*</td>
<td>0.613</td>
</tr>
<tr>
<td>Race/Ethnicity distribution</td>
<td>8 Caucasian, 1 AA, 1 American Indian, 13 Hispanic</td>
<td>10 Caucasian, 7 AA, 6 Hispanic, 1 Asian, 1 Biracial</td>
<td>Fisher’s</td>
<td>0.026</td>
</tr>
<tr>
<td>Maternal Education (yrs)</td>
<td>12.5 ± 3.4, range = 5.0 – 16.0</td>
<td>14.0 ± 2.7, range = 8.0 – 20.0</td>
<td>1.71</td>
<td>0.094</td>
</tr>
<tr>
<td>Handedness</td>
<td>22 right, 1 left</td>
<td>21 right, 4 left</td>
<td>Fisher’s</td>
<td>0.350</td>
</tr>
<tr>
<td>Mechanism of Injury (accident type)</td>
<td>12 MVC, 4 Motorcycle, 2 RV, 1 Bicycle, 2 Fall, 2 Hit by motor vehicle</td>
<td>1 Motorcycle, 1 RV, 7 Fall, 1 Hit by objects, 14 Sports, 1 Other</td>
<td>Fisher’s</td>
<td>0.0001</td>
</tr>
<tr>
<td>Glasgow Coma Score</td>
<td>7.5 ± 4.1; range = 3.0 – 15.0</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Time post-injury: 3M (months)</td>
<td>4.0 ± 0.9, range = 2.5 – 5.3</td>
<td>4.2 ± 1.0, range = 2.7 – 7.1</td>
<td>0.49</td>
<td>0.629</td>
</tr>
<tr>
<td>Time post-injury: 18M (months)</td>
<td>18.9 ± 1.5, range = 16.7 – 22.6</td>
<td>18.8 ± 1.3, range = 16.6 – 20.9</td>
<td>-0.26</td>
<td>0.795</td>
</tr>
</tbody>
</table>

*Note.* TBI = traumatic brain injury; OI = orthopedically injured; MVC = Motor vehicle crash, †RV = Recreation vehicle crash. M=mean. SD=Standard Deviation. AA = African American. *=performed with Chi-square test.
Table 2

*Group Difference of the CC for FA at three and 18 months*

<table>
<thead>
<tr>
<th>Regions</th>
<th>Interval</th>
<th>TBI (Mean ± SD)</th>
<th>OI (Mean ± SD)</th>
<th>F value</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total CC</td>
<td>3M</td>
<td>0.42 ± 0.03</td>
<td>0.47 ± 0.02</td>
<td>5.03</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.43 ± 0.03</td>
<td>0.47 ± 0.02</td>
<td>2.36</td>
<td>0.043</td>
</tr>
<tr>
<td>Genu</td>
<td>3M</td>
<td>0.42 ± 0.03</td>
<td>0.47 ± 0.03</td>
<td>2.37</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.43 ± 0.03</td>
<td>0.48 ± 0.02</td>
<td>6.97</td>
<td>0.0001</td>
</tr>
<tr>
<td>Body</td>
<td>3M</td>
<td>0.41 ± 0.04</td>
<td>0.45 ± 0.03</td>
<td>4.16</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.42 ± 0.03</td>
<td>0.46 ± 0.02</td>
<td>2.36</td>
<td>0.043</td>
</tr>
<tr>
<td>Splenium</td>
<td>3M</td>
<td>0.46 ± 0.04</td>
<td>0.49 ± 0.02</td>
<td>4.70</td>
<td>0.0004</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.47 ± 0.04</td>
<td>0.51 ± 0.02</td>
<td>3.64</td>
<td>0.003</td>
</tr>
</tbody>
</table>

*Note.* CC = corpus callosum. SD = standard deviation. FA = fractional anisotropy. Bold typeface indicates significance after Bonferroni correction for multiple comparisons.
Table 3

*Group Difference of the CC for ADC at three and 18 months*

<table>
<thead>
<tr>
<th>Regions</th>
<th>Interval</th>
<th>TBI (Mean ± SD)</th>
<th>OI (Mean ± SD)</th>
<th>F value</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total CC</td>
<td>3M</td>
<td>0.98 ± 0.07</td>
<td>0.93 ± 0.05</td>
<td>-3.03</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.99 ± 0.07</td>
<td>0.91 ± 0.06</td>
<td>-4.30</td>
<td>0.0001</td>
</tr>
<tr>
<td>Genu</td>
<td>3M</td>
<td>0.91 ± 0.03</td>
<td>0.86 ± 0.04</td>
<td>-3.80</td>
<td>0.004</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.93 ± 0.07</td>
<td>0.85 ± 0.05</td>
<td>-4.58</td>
<td>0.0001</td>
</tr>
<tr>
<td>Body</td>
<td>3M</td>
<td>0.97 ± 0.10</td>
<td>0.92 ± 0.07</td>
<td>-2.06</td>
<td>0.046</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.97 ± 0.08</td>
<td>0.90 ± 0.06</td>
<td>-3.43</td>
<td>0.001</td>
</tr>
<tr>
<td>Splenium</td>
<td>3M</td>
<td>1.02 ± 0.09</td>
<td>0.95 ± 0.06</td>
<td>-3.18</td>
<td>0.003</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>1.04 ± 0.10</td>
<td>0.95 ± 0.07</td>
<td>32.30</td>
<td>0.049</td>
</tr>
</tbody>
</table>

*Note.* CC = corpus callosum. SD = standard deviation, ADC = apparent diffusion coefficient. ADC values are represented in $10^{-3}$ mm$^2$/sec. Bold typeface indicates significance after Bonferroni correction for multiple comparisons.
Longitudinal DTI Analyses

Both groups demonstrated increased FA in total CC and all CC sub-regions at 18 months as compared to three months post-injury. However, only the increase of FA in the splenium/posterior sub-region in both the TBI ($t(22) = -3.36, p = 0.003$) and OI ($t(24) = -2.76, p = 0.011$) groups reached significance after correction for multiple comparisons. The TBI group showed a marginally significant increase in the FA for the total CC ($t(22) = -2.21, p = 0.038$) over time. The OI group showed a marginally significant increase of FA in the total CC ($t(24) = -2.04, p = 0.053$) and marginally significant decreases in ADC for the body/central subregion ($t(24) = 2.10, p = 0.047$) and total CC ($t(24) = 1.76, p = 0.092$). No significant between-group differences for longitudinal changes were noted in any region. Figure 1 illustrates the change between three and 18 months for FA and ADC of both the splenium/posterior sub-region of the CC and the total CC in both groups.
Figure 1. Bar graphs with error bars illustrating the longitudinal change between three and 18 months post-injury in FA (a) and ADC (b) as measured by quantitative DTI tractography, and volume (c) as measured by volumetric analysis in the splenium/posterior sub-region of the CC in both the TBI and OI groups. Note that the FA remains significantly lower in the TBI group as compared to the OI group, but increases comparably to the OI group. The ADC decreases over time in the OI group, as would be expected given normal developmental change, but ADC in the TBI group continues to increase at a modest rate, which is unexpected given the typical relation between FA and ADC. The volume of the splenium increases in the OI group, as expected given normal developmental change that is occurring in this age range, but evidences continued degenerative change in the TBI group over time. Changes in the total CC are reflected in FA (d), ADC (e) as measured by quantitative DTI tractography, and volume (f) as measured by volumetric analysis.

Relation of DTI to Cognitive Functioning

Cognitively, the OI group demonstrated a significant improvement in speed of information processing over time ($t(18) = 2.55, p = 0.02$). The speed of information processing did not differentiate the TBI and OI groups at either three or 18 month occasions when all participants were included. However, when the participants with complicated-mild injuries were excluded, results were unchanged at three months, but there was a trend for group difference at 18 months ($t(28) = -1.90, p = .086, d = 1.25$) such that the participants with TBI had slower RTs.

At three months post-injury in the TBI group, FA in the splenium ($\rho = -0.58, p = 0.008$) and total CC ($\rho = -0.55, p = 0.012$) were significantly correlated with baseline
condition processing speed. Higher FA was related to faster reaction time. These correlations became stronger at the 18 month post-injury occasion (for the splenium/posterior sub-region: \( \rho = -0.84, p = 0.001 \); for total CC: \( \rho = -0.71, p = 0.014 \)). Figure 2 reflects the relation between FA in the splenium/posterior sub-region of the CC and reaction time on the Flanker task in each group.

\[ \text{Figure 2. Scatterplots with regression lines demonstrating the relation between Flanker baseline condition RT and FA of the splenium/posterior sub-region (a) and the total CC (b) at 18 months post-injury in the TBI and OI groups. Note the significant correlation for the TBI group.} \]

**Group Differences at Three and 18 Months Post-Injury for Volumetrics**

At three months post-injury, the mid-anterior \( (F(1,37) = 6.25, p = 0.017) \) and body/central \( (F(1,37) = 5.36, p = 0.026) \) sub-regions and total CC \( (F(1,37) = 4.27, p = 0.046) \) marginally differed between the groups, though the genu/anterior and
splenium/posterior regions did not significantly differ (see Table 4). In each case, the TBI group exhibited decreased volume in the sub-region relative to the OI comparison group. At 18 months post-injury, the genu/anterior \(F(1,37) = 7.76, p = 0.008\), mid-anterior \(F(1,37) = 15.13, p = 0.0004\), body/central \(F(1,37) = 10.20, p = 0.003\), and mid-posterior \(F(1,37) = 8.96, p = 0.005\) sub-regions of the CC as well as the total CC \(F(1,37) = 11.83, p = 0.002\) significantly differed between groups (see Table 4 for least squares means), again with the TBI group demonstrating significantly decreased volume relative to the OI group.

**Longitudinal Volumetric Analyses**

In terms of within group differences, in the OI group, the genu/anterior \(p = 0.010\), mid-posterior \(p = 0.002\), and splenium/posterior \(p = 0.0008\) sub-regions of the CC as well as the total CC \(p = 0.001\) demonstrated significant volume increase between three and 18 months post-injury. Mid-anterior and body/central sub-regions marginally differed, again demonstrating an increase in volume over time, but this change was not significant. In contrast, the TBI group demonstrated decreases in volume between three and 18 months post-injury in the genu/anterior \(p = 0.009\) and mid-posterior \(p = 0.007\) sub-regions and total CC \(p = 0.012\). Change in CC volume between three and 18 months revealed significant between-group differences in the genu/anterior (Satterthwaite \(t(17.8) = -3.70, p = .002\)), mid-anterior (Satterthwaite \(t(20) = -2.95, p = 0.008\)), mid-posterior (Satterthwaite \(t(20) = -4.19, p = 0.0004\)) and splenium/posterior (Satterthwaite \(t(16.5) = -3.20, p = 0.005\)) regions and the total CC (Satterthwaite \(t(17.2) = -3.73, p = 0.002\)). A marginally significant difference was noted for the body/central sub-region \((t(38) = -1.82, p = 0.077)\). Figures 1c and 2c illustrate the pattern of change between three and 18 months in the TBI and OI groups for the splenium/posterior sub-region and the total CC, respectively.
Table 4

*Volumetric Differences in Corpus Callosum Sub-regions at three and 18 months*

<table>
<thead>
<tr>
<th>Regions</th>
<th>Interval</th>
<th>TBI (LSM)</th>
<th>OI (LSM)</th>
<th>F value</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total CC</td>
<td>3M</td>
<td>2.72</td>
<td>2.92</td>
<td>4.27</td>
<td>0.46</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>2.52</td>
<td>2.99</td>
<td>11.83</td>
<td><strong>0.002</strong></td>
</tr>
<tr>
<td>Anterior</td>
<td>3M</td>
<td>0.72</td>
<td>0.79</td>
<td>3.19</td>
<td>0.082</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.67</td>
<td>0.80</td>
<td>7.76</td>
<td><strong>0.008</strong></td>
</tr>
<tr>
<td>Mid-anterior</td>
<td>3M</td>
<td>0.41</td>
<td>0.46</td>
<td>6.25</td>
<td><strong>0.017</strong></td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.37</td>
<td>0.47</td>
<td>15.13</td>
<td><strong>0.0004</strong></td>
</tr>
<tr>
<td>Central</td>
<td>3M</td>
<td>0.40</td>
<td>0.47</td>
<td>5.36</td>
<td>0.026</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.38</td>
<td>0.48</td>
<td>10.20</td>
<td><strong>0.003</strong></td>
</tr>
<tr>
<td>Mid-posterior</td>
<td>3M</td>
<td>0.40</td>
<td>0.42</td>
<td>1.57</td>
<td>0.218</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.36</td>
<td>0.44</td>
<td>8.96</td>
<td><strong>0.005</strong></td>
</tr>
<tr>
<td>Posterior</td>
<td>3M</td>
<td>0.79</td>
<td>0.79</td>
<td>0.11</td>
<td>0.74</td>
</tr>
<tr>
<td></td>
<td>18M</td>
<td>0.74</td>
<td>0.81</td>
<td>2.14</td>
<td>0.152</td>
</tr>
</tbody>
</table>

*Note.* LSM = least squares means. Volumetric measures are represented in cubic centimeters. Bold typeface indicates significance after Bonferroni correction for multiple comparisons.
Relation of Volume to Cognitive Function

There were no significant correlations between processing speed and volumes of the sub-regions of the corpus callosum in either group at three and 18 month post-injury interval assessments. Total corpus callosum volume was marginally related to processing speed at 18 months in the OI group only ($\rho = -0.433$, $p = 0.06$), with greater volume associated with faster processing speed.

Discussion

This study demonstrated continued late degenerative change as reflected in the gross morphometry of nearly all regions of the corpus callosum as measured by volumetric analysis between three and 18 months post-injury in the TBI group. These degenerative changes were most prominent in the genu/anterior, mid-posterior and splenium/posterior sub-regions of the corpus callosum. In contrast, the genu/anterior, mid-posterior, and splenium/posterior sub-regions of the corpus callosum demonstrated significant volume increases in the comparison group of children with orthopedic injury, consistent with known developmental changes in this age range (Lobel, et al., 2009; Zhang, et al., 2005). Previous developmental studies have demonstrated that the thinly myelinated and densely packed fibers more characteristic of the genu mature later than those with thicker myelination (Kochunov, et al., 2010), and hence the developmental changes may be greater in these regions during the age range of the children whom we studied. It is also possible that these fibers that may sustain a differential vulnerability to aging (Kochunov, et al., 2010) or insult, including TBI-induced axonal injury in particular (Reeves, Phillips, & Povlishock, 2005). Traumatic-induced atrophy of the corpus callosum likely occurs as a consequence of complex neuropathological features that involve both direct trauma to the CC (Mac Donald,
Dikranian, Song, et al., 2007; Parizel, et al., 1998) as well as secondary degeneration of a Wallerian type occurring because of diffuse damage elsewhere in the brain disrupts the integrity of white matter tracts (Di Paola, et al., 2010).

Typical CC development in childhood follows a predictable course of myelination, which at least in animal models also relates to environmental complexity (Markham, Herting, Luszpak, Juraska, & Greenough, 2009). In the typically-developing brain, this likely also leads to complex brain-environment-developmental interactions resulting in increased CC volume during the age ranges of the children in this study and as confirmed in the results of the OI participants. However, brain injury may induce deficits in cognitive processing which likely diminish the child’s ability to process complex material (Taylor, 2004). Hampered by slowed processing speed and diminished integration of inter-hemispheric information due to the CC damage, theoretically this could interact negatively with development and minimize CC growth trajectories. As reflected in further reduced volume over time, the TBI children did not exhibit maturation-related volume increases in the CC. Likewise, it is not unexpected that processing speed would improve in typically-developing children, as was observed in the children with OI, but not TBI participants. These findings support the notion that brain injury disrupts normal white matter maturation and development, disconnecting the typical link with maturational changes that improve processing speed.

However, the current study shows a most interesting apparent adaption in CC functioning in that children with TBI could adequately perform the Flanker Task and at a level that did not differ from OI controls at the three month baseline or 18 month follow-up. This suggests that within the initial three month post-injury time frame, adaptations in CC functioning were sufficient for these patients to adequately perform the task despite the
volume and DTI changes that had already occurred at that stage. In case of surgical treatment of the CC, typically performed as a procedure to treat chronic epilepsy, adaptive rerouting of CC pathways has long been documented (Bloom & Hynd, 2005; Tanriverdi, Olivier, Poulin, Andermann, & Dubeau, 2009). This is likely part of the explanation within the TBI group that part of the recovery from TBI is some form of programmed adaptive mechanisms that engage intact tracts to adapt. It is also possible that new inter- and intra-hemispheric communications could potentially be formulated via alternative routes, such as the anterior commissure (Patel, Toussaint, Charles-Edwards, Lin, & Batchelor, 2010). Even though the TBI group was not significantly slower than the OI group on the Flanker task, DTI findings did relate to processing speed in the TBI group in the splenium/posterior sub-region, which became stronger in this group over time.

The continued degenerative changes over this time interval on volumetrics in the TBI group were not reflected in longitudinal changes in DTI metrics such as FA. Despite loss of volume in the TBI group in some sub-regions, likely reflective of some degree of neuronal death and axonal degeneration, the FA generally increased in both the TBI and the OI groups in all regions. In animal studies, histological findings have related tissue volume loss to cellular loss (Grady, Charleston, Maris, Witgen, & Lifshitz, 2003) and specifically, CC volume loss to neuronal loss (Bramlett & Dietrich, 2002). On the other hand, DTI is likely showing how injured neurons, while damaged, recover over time (Kumar, et al., 2010), at least to some degree. In the current study, these two methods likely reflect quite different neuropathological end-points over time where reduced volume reflects overall neuronal loss whereas DTI findings more specifically reflect axonal integrity, restorative and adaptive processes that develop over time after injury. Figure 3 illustrates the qualitative differences
in DTI tractography over time that compliment the quantitative findings we describe in this study in one participant. However, it should be noted that although FA in the TBI group did increase, ADC in the TBI group did not decrease as would have been anticipated given the traditional inverse relation between FA and ADC, and in fact, ADC also increased over time. This suggests that the expected developmental trajectory reflected in FA increase and ADC increase observed in the comparison group may, in fact, be altered somewhat by TBI despite the apparent increase in FA. As both FA and ADC may be impacted by multiple factors related to the microstructural characteristics of the tissue, including fiber density, axonal diameter and the ratio of intracellular/extracellular space, further investigation into the more specific nature of these changes is warranted.

*Figure 3.* T1-weighted sagittal images of a 16 year old male adolescent with orthopedic injury at three months (a) and 18 months (b) post-injury. T1-weighted images of a 15-year old male adolescent with moderate TBI (Glasgow Coma Scale score of 9) at three months (c)
and 18 months (d) post injury is also presented. Note the subtle atrophy of the CC in the TBI child. DTI tractography of the CC is overlaid on the T1-weighted image for the adolescent with OI at three months (e) and 18 months (f) post-injury, which reflects subtle qualitative increase in the fiber density. Similarly, the DTI tractography of the CC overlaid on the T1-weighted imaging from the adolescent with TBI at three months (g) and 18 months (h) post-injury reveals qualitative increase in both the fiber density and intensity of the CC over time, though the density of fibers is still significantly less than that of the comparison OI participant.

Although the trajectory of change in FA for both groups followed a similar slope post-injury, the difference between the TBI and OI groups remained significant, with the TBI group exhibiting a significantly lower mean FA at both time points in several regions. This also reflects that DTI changes such as FA may be more specific to adaptive mechanism of recovery than are traditional volumetric measures (van der Zijden, van der Toorn, van der Marel, & Dijkhuizen, 2008). This difference further indicates that neuropathologically DTI is assessing something quite different than just neuronal loss or axonal degradation. Because DTI did relate to processing speed, it may be the more sensitive biomarker of actual CC integrity than measuring atrophic changes through volume loss. Because DTI is such a new technology, studies are just now being done to ascertain whether microstructural changes as found with DTI may occur independently of volumetric parenchymal changes (Canu, et al., 2010). Clearly, multimodality imaging studies may be useful in further examination of TBI-related consequences to brain parenchyma.
Strengths, Limitations and Future Directions

Few studies have specifically examined longitudinal changes following TBI using advanced imaging modalities such as volumetric analysis and DTI (Bendlin, et al., 2008; Kumar, et al., 2009; Kumar, et al., 2010; Sidaros, et al., 2008; Sidaros, et al., 2009), and to our knowledge, no study has examined these changes on DTI in a pediatric TBI population (Levin, et al., 2000). Strengths of this study include its prospective, longitudinal design and its use of an orthopedically-injured comparison group.

We limited this investigation to changes in the corpus callosum, but additional brain regions should be examined in future studies. As we have shown elsewhere, diffuse changes in the brain result from moderate to severe TBI (Bigler, et al., 2010) including thinning of the cerebral cortex (Merkley, et al., 2008) but how these diffuse changes relate specifically to the atrophic changes of the CC and cognitive outcome is not known at this time. Additionally, other methods of DTI analysis such as voxel based analysis and region of interest analysis will be applied in future studies to examine these differences.

We emphasize that these analyses represent group trends, and we acknowledge the possibility that different patterns of change over time may emerge in some individuals with TBI based upon factors related to the heterogeneity of injury type and location, age at injury, mechanism of injury, and pre- and post-injury characteristics that may render some individuals more resilient to the effects of injury than others. Future studies may examine some of these factors.

Finally, missing data may have limited the power to detect differences in the behavioral and cognitive measures. For instance, the effect size for group differences on the Flanker task RT at three months was 0.37, which is small to moderate, but the power to
detect a significant difference was only of 33.7%. Similarly at 18 months, the effect size was 0.62, which is moderate to large, but the power to detect a significant difference was only 43.4%.

Conclusions

The current findings indicate that at the macro- and micro-structural level, two seemingly opposite trajectories were noted following moderate to severe TBI in children and adolescents. Volumetric analyses demonstrated degeneration in the CC over time, whereas DTI findings showed some degree of increase in the FA over time, indicative of maturation or recovery. FA was also related to speed of information processing at both time points, though even more strongly at the 18 month post-injury interval. White matter changes as measured by multimodality imaging studies may be useful in better understanding changes indicative of both degeneration and residual loss of tissue and cognitive difficulty as well as plasticity and recovery, particularly in an age range that is also undergoing dynamic developmental change.
References


Gorrie, C., Duflou, J., Brown, J., Gibson, T., & Waite, P. M. (2001). Extent and
distribution of vascular brain injury in pediatric road fatalities. *Journal of
Neurotrauma, 18*, 849-860.

glial cell number in the hippocampus after experimental traumatic brain injury:

Huisman, T. A. G. M., Schwamm, L. H., Schaefer, P. W., Koroshetz, W. J., Shetty-Alva, N.,
Ozsunar, Y., et al. (2004). Diffusion tensor imaging as potential biomarker of white
376.

of Neurosurgery, 103*, 298-303.

information processing after mild and moderate traumatic brain injury. *Brain Injury,
23*, 1027-1040.

mental slowing and executive deficits in subjects with age-related white matter
hyperintensities: the LADIS study. *Journal of Neurology, Neurosurgery, Psychiatry*,
78, 491-496.

consequences of diffuse traumatic brain injury: a large deformation tensor-based


Neuroradiology, 28, 1919-1925.