Tractography of developing white matter of the internal capsule and corpus callosum in very preterm infants

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Abstract

Objectives To investigate in preterm infants associations between Diffusion Tensor Imaging (DTI) parameters of the posterior limb of the internal capsule (PLIC) and corpus callosum (CC) and age, white matter (WM) injury and clinical factors.

Methods In 84 preterm infants DTI was performed between 40–62 weeks postmenstrual age on 3 T MR. Fractional anisotropy (FA), apparent diffusion coefficient (ADC) values and fibre lengths through the PLIC and the genu and splenium were determined. WM injury was categorised as normal/mildly, moderately and severely abnormal. Associations between DTI parameters and age, WM injury and clinical factors were analysed.

Results A positive association existed between FA and age at imaging for fibres through the PLIC (\( r = 0.48 \), \( p < 0.001 \)) and splenium (\( r = 0.24 \), \( p < 0.01 \)). A negative association existed between ADC and age at imaging for fibres through the PLIC (\( r = -0.65 \), \( p < 0.001 \)), splenium (\( r = -0.35 \), \( p < 0.001 \)) and genu (\( r = -0.53 \), \( p < 0.001 \)). No association was found between DTI parameters and gestational age, degree of WM injury or categorical clinical factors.

Conclusions These results indicate that in our cohort of very preterm infants, at this young age, the development of the PLIC and CC is ongoing and independent of the degree of prematurity or WM injury.

Keywords Diffusion tensor imaging · Fibre tractography · Very preterm infants · Term equivalent age · White matter injury

Abbreviations

ADC apparent diffusion coefficient
BPD bronchopulmonary dysplasia
CC corpus callosum
DEHSI diffuse and excessive high signal intensity
DWI diffusion-weighted imaging
DTI diffusion tensor imaging
FA fractional anisotropy
GA gestational age
HC head circumference
IUGR intrauterine growth restriction
PDA persistent ductus arteriosus
PLIC posterior limb of internal capsule
PMA postmenstrual age
PWML punctuate white matter lesion
RDS respiratory distress syndrome
TEA term equivalent age
VPTI very preterm infants
WM white matter

Introduction

Infants born at a gestational age (GA) below 32 weeks are prone to diffuse white matter (WM) injury [1, 2]. This may eventually result in damage, underdevelopment and atrophy of the internal capsule and the corpus callosum (CC) [3–7]. Conventional MRI techniques detect WM injury and depict features of brain maturation [2, 8–15]. Diffusion Tensor Imaging (DTI) has been proposed as an additional tool in the assessment of WM injury and may provide more
adequate diagnostic and prognostic information in relation to neurological outcome than conventional MR imaging [16–21]. DTI enables quantitative assessment of maturation, tract organisation and injury by calculating Fractional Anisotropy (FA) and Apparent Diffusion Coefficient (ADC) values. Fibre tractography offers insight into developing WM by visualisation of the WM tracts [17–19, 22–25]. The non-invasive nature of DTI permits longitudinal and diagnostic clinical studies [26–33].

Until now, the clinical use of DTI fibre tractography in very preterm infants (VPTI) has been restricted by the lack of normal reference values for FA and ADC [16]. Only a few studies have used DTI in VPTI with a GA below 32 weeks [16, 18, 23–25]. In most of these studies MRI was performed before term equivalent age (TEA). Little is known about the changes in FA or ADC in the developing brain around TEA [18, 34–37]. Data on the influence of diffuse WM injury on tract maturation and organisation around TEA are scarce [35, 38, 39].

We studied DTI parameters (FA and ADC values and lengths of fibres) of WM tracts passing through the posterior limb of the internal capsule (PLIC) and the CC in VPTI undergoing MRI at or within the first months of TEA. The aims of this exploratory study were to establish the association between DTI parameters and:

- GA and postmenstrual age (PMA) at the time of imaging
- The degree of WM injury
- Categorical clinical factors that may influence brain maturation

Materials and methods

Preterm infants

As part of an ongoing prospective study of VPTI with a GA ≤32 weeks, admitted to the neonatal intensive care unit of our institution, 113 infants underwent MRI (3 Tesla) between May 2006 and October 2007.

MRI was preferably performed at TEA (PMA 40–44 weeks). For infants who were unstable and/or ventilator dependent around that age, MRI was postponed (PMA range 40–62 weeks). Ethical approval for the study was given by the institutional review board and informed parental consent was obtained for each infant. In 102 infants DTI was performed. Three children with congenital brain abnormalities were excluded. In 15 cases, fibre tractography was not possible due to motion artefacts. In the remaining 84 infants a complete DTI dataset was acquired. Clinical parameters were retrospectively collected from the patient’s medical records (Table 1).

Image and data acquisition

All MRI examinations were performed on a 3 Tesla MR system (Philips Medical Systems, Best, the Netherlands) according to a standard protocol [40]. The infants were sedated using chloral hydrate (55 mg/kg), laid supine and snugly swaddled during the imaging procedure. Ear protection consisted of neonatal earmuffs (Natus Mini Muffs; Natus Medical Inc., San Carlos, CA, USA) covered by a headphone. All MRI examinations included a 3D T1-TFE sequence (TR 9.7 ms, TE 4.6 ms, flip angle 8, TFE factor 128, slice thickness 1 mm) and a T2-TSE sequence (TR 6269 ms, TE 120 ms, turbo factor 18, slice thickness 2 mm). In addition a DTI sequence (SE-EPI, TR 7,456 ms, TE 54 ms, slice thickness 2 mm, voxel size 1.4×1.4×2 mm with diffusion acquisitions in 32 directions and a b-value of 1,000 s/mm2, EPI factor 56) with an imaging time of 5 min and 34 s was acquired.

Fibre tractography of the PLIC and CC

DTI datasets were analysed on an off-line workstation using commercially available processing software as provided by the manufacturer (FiberTrak, by Philips Medical Systems, Best, the Netherlands). DTI colour-coded maps were automatically computed, red representing a right-left, green an anterior-posterior and blue a superior-inferior orientation. These colour-coded DTI maps were used to place a single seed to perform fibre tracking using an automated 3D seeded algorithm. First, for the PLIC in the axial plane at the level of the lateral ventricles and subsequently for the genu and splenium of the CC in the mid-sagittal plane regions of interest were defined. The seed was placed as is shown in Fig. 1. Position changes of the seed-point in the structure of interest resulted in identical fibre tracts, indicating the robustness of the method used. Two investigators (FTdB and AvS, blinded to subject’s age and degree of WM injury) identified the regions of interest and manually placed the individual seeds in consensus for each region of interest. Subsequently, fibre tracts were generated through the PLIC and the callosal genu and splenium resulting in visualisation of fibres (Fig. 2). The quality of all tracts was visually assessed in order to minimise erroneous pathways, which were erased and not used in further analyses. In all analyses, default settings were used consisting of a minimum FA of 0.15, a maximum angle change of 27.0 degrees and a minimum fibre length of 10.0 mm. We used the manufacturer default settings, since (small) changes of these settings, to optimize the performance of the fibre tracking, did not have any influence on the fibres picked by the tracking routine. Finally FA and ADC values of these fibre bundles were obtained and the length of the fibres was calculated.
WM injury

All T1-W and T2-W sequences were analysed by two investigators (FTdB, paediatric neuroradiologist with more than 15 years of experience, LML, with more than 4 years’ experience) by consensus to assess WM injury. We categorised the WM injury into [41]:

• Normal/mildly abnormal WM: normal appearing WM or homogeneous diffuse and excessive high signal intensity (DEHSI) as seen on T2-weighted images, or few (≤6) punctate white matter lesions (PWML)
• Moderately abnormal WM: multiple (> 6) PWML and/or small localised cystic lesions and/or heterogeneous DEHSI
• Severely abnormal WM: extensive or diffuse heterogeneous signal intensity changes and/or haemorrhagic or cystic lesions involving the periventricular and/or subcortical WM

Accordingly, 18 infants were classified as having normal/mildly, 51 infants moderately and 15 infants severely abnormal WM (Table 1)

Statistical analysis

Data were analysed using SPSS 16.0.2 for Windows. Frequency counts and percentages were used to summarise categorical variables. For continuous variables, the mean, standard deviation of the mean and range are reported (Table 1).

Differences between DTI values of left and right PLIC and between the genu and splenium of the CC were determined by using Wilcoxon signed rank tests for related samples.

To establish the association between DTI parameters and age, weight and head circumference (HC) Spearman Rank correlation coefficients were calculated.

Stepwise linear regression analysis was used to assess the relative importance of the various study population characteristics that were associated with the DTI parameters through the PLIC and CC. Applying this model we corrected all other parameters for PMA at MRI and HC. Variables, for which the p value by univariate analyses tests was ≤0.1, were included in the subsequent linear stepwise regression analysis.

To determine the association between DTI parameters and WM injury and categorical clinical factors, Mann-Whitney U analyses or Kruskal-Wallis tests were performed where appropriate. One-way ANOVA with post-hoc Scheffe tests were used to compare DTI parameters between groups categorised as normal/mildly, moderately or severely abnormal WM.

A p value ≤0.05 was considered statistically significant for all analyses.

Results

General characteristics of the study population

Clinical characteristics for all 84 infants are shown in Table 1.

Quantitative DTI values

The FA and ADC values and length of fibres (mean, SD and range) through the PLIC and genu and splenium of the CC are shown in Table 2. As we found no statistical differences or any laterality between left and right DTI values in the PLIC, these DTI values were averaged. In the genu lower FA (p<0.001), lower ADC values (p<0.001) and shorter fibres (p<0.001) were found compared with the splenium of the CC.

Association between DTI parameters and continuous clinical variables

Table 3 shows the associations between DTI parameters for the PLIC and genu and splenium and continuous clinical

### Table 1 Distribution of categorical clinical factors, white matter injury and continuous clinical parameters in the study population (n=84)

| Categorical clinical factors | Number (N) | Percentage (%) |
|-----------------------------|------------|----------------|
| Male                        | 55         | 65.5           |
| Female                      | 29         | 34.5           |
| Plurality                   | 23         | 27.4           |
| IUGR                        | 10         | 11.9           |
| Antenatal corticosteroids   | 43         | 51.2           |
| Perinatal infection         | 30         | 35.7           |
| RDS                         | 44         | 52.4           |
| PDA                         | 23         | 27.4           |
| Hypotension                 | 29         | 34.5           |
| Use of dexamethasone for BPD| 41         | 48.8           |
| Degree of WM injury         |            |                |
| Normal/mildly abnormal WM   | 18         | 21.4           |
| Moderately abnormal WM      | 51         | 60.7           |
| Severely abnormal WM        | 15         | 17.8           |

| Continuous clinical parameters | Mean (SD) | Range         |
|--------------------------------|------------|---------------|
| GA (weeks)                     | 29.0 (2.0) | 25.6-31.9     |
| Birth weight (g)               | 1199 (365) | 585-1960      |
| Head circumference at birth (cm)| 25.8 (4.0)| 22.0-31.2     |
| PMA at MRI (weeks)             | 45.0 (4.2) | 40.0-62.1     |
| Weight at MRI (gram)           | 4002 (867) | 2010-7005     |
| Head circumference at MRI (cm)  | 37.7 (1.8) | 32.7-41.5     |

IUGR intrauterine growth restriction, RDS respiratory distress syndrome, PDA patent ductus arteriosus, BPD bronchopulmonary dysplasia, WM white matter, GA gestational age, PMA postmenstrual age
variables. DTI parameters in the PLIC ($p<0.001$) and CC ($p<0.001$ and $p<0.01$) correlated strongly with PMA at MRI, except for the FA in the genu of the CC. The FA ($p<0.01$), ADC ($p<0.001$) and fibre length ($p<0.01$) through the PLIC and fibre length through the genu and splenium of the CC ($p<0.001$, $p<0.01$ and $p<0.001$, $p<0.001$ respectively) correlated with weight and HC at MRI. No or only weak associations were found between FA, ADC and fibre length through the PLIC or CC and GA or birth weight.

Because of the wide age range, we re-analysed data for a subgroup of infants (69/84) who underwent imaging between 40–46 weeks PMA. For this subgroup, we still found a significant association between FA ($r=0.252$, $p<0.05$), ADC ($r=-0.470$, $p<0.01$) and fibre length ($r=0.410$, $p<0.01$) through the PLIC and PMA at MRI. The association between FA, ADC values or fibre length in the PLIC and weight and HC at MRI no longer existed. For the genu, no association between FA and any continuous clinical variable was found, while the association between ADC ($r=-0.268$, $p<0.05$) or fibre length ($r=0.402$, $p<0.01$) and PMA at MRI remained. For the splenium we found an association between fibre length and weight and HC at MRI (respectively $r=0.337$, $p<0.01$ and $r=0.332$, $p<0.05$). The association with PMA at MRI was no longer present, still the association between fibre length and HC at birth remained ($r=0.417$, $p<0.05$).

Figure 3 shows the individual data depicting the association between the FA and ADC of fibres through the PLIC and GA as well as PMA at MRI. This figure illustrates the association between FA and ADC values with PMA at imaging, whereas no such association exists with GA.

Results of the stepwise regression analyses are shown in Table 4, delineating the strongest contributing factor of clinical parameters. In all but three cases, PMA at MRI was the strongest contributing factor to most of the DTI parameters (FA and ADC of the fibres through the PLIC, ADC and fibre length of the fibres through the genu and splenium, and FA of the splenium). For two of the remaining three parameters (FA in the fibres through the genu and fibre length in the fibres through the splenium) HC at birth contributed most. For the length of fibres through the PLIC, the HC at MRI contributed most.

Association between DTI parameters and WM injury and categorical clinical factors

We found no association between DTI parameters for the PLIC or CC and degree of WM injury. When comparing the groups with different degrees of WM injury, we also found no significant differences in DTI parameters (Table 5). These results remained, even after correction for PMA at MRI.

No associations existed between DTI parameters and categorical clinical factors.

Discussion

In this study, we determined FA and ADC values and the length of WM fibres passing through the PLIC and CC in a cohort of 84 VPTI imaged between 40 and 62 weeks PMA. We assessed the association between DTI parameters and age, diffuse WM injury and clinical factors.

All DTI parameters of the PLIC and most DTI parameters of the CC were strongly associated with PMA at imaging. We found no associations between DTI parameters and GA, the degree of WM injury or any categorical clinical factor.
Fig. 2 a-b: Fibre tracts in an infant undergoing imaging at PMA of 41 weeks a through the PLIC and b through the genu and splenium of the CC using an automated 3D seeded technique according to default settings (FiberTrak, Philips Medical Systems, Best the Netherlands; minimum FA 0.15, maximum angle change 27.0 degrees, minimum fibre length 10.0 mm)
In a previous study by Dudink et al. of very low birth weight infants undergoing imaging within 4 days of birth an association was found between FA of the PLIC and GA. They found no association between ADC values and GA [16]. Although we did not find an association between FA or ADC of the PLIC and GA, we found a positive association between FA and PMA at MRI and a negative association between ADC and PMA at MRI. Increase in FA and decrease in ADC have been described before and reflect the (pre-)myelination and parallel organisation of fibres, such as expected in the development of the PLIC [17, 23, 24]. As we found no association with GA, the maturation of the PLIC seems to progress with age, independent of the degree of prematurity. Similar to our findings in the PLIC, no associations were found between DTI parameters of fibres passing through the CC and GA, while DTI parameters of the CC showed a significant association with PMA at MRI, except for the FA values of the genu. This indicates ongoing maturation of the splenium of the CC around TEA, again independent of GA, while for the genu this seems less clear. These findings are in agreement with the results of Partridge et al. who found no significant maturational trends of the genu of the CC in their serial study of 14 premature neonates with a GA between 25 and 34 weeks, of whom eight underwent a second MRI around TEA [24].

We found no associations between DTI values for the PLIC or CC and the degree of WM injury. These findings are only partly in agreement with the results of Hüppi et al., who at TEA found no difference in ADC values of WM between preterm infants with and without WM lesions. They found a 20% lower regional anisotropy in fibres descending from the internal capsule in children with WM injury compared with those without [18]. Our data do not confirm this latter finding. This may be explained by the fact that we studied a prospective cohort of unselected VPTI, with relatively small numbers of children with normal or severely abnormal WM. Hüppi et al. selected ten infants with WM injury and matched these with ten infants of the same GA and neonatal course but with normal MRI. Our results also differ from those of Counsell et al., who demonstrated in preterm infants who underwent imaging around TEA (38.86–43.86 weeks), that radial diffusivity in the PLIC and the splenium of the CC and axial and radial diffusivity in the WM were significantly elevated in infants with DEHSI compared with those with normal appearing WM and term control.

### Table 2

|                      | Mean (SD)        | Range        |
|----------------------|------------------|--------------|
| **PLIC**             |                  |              |
| FA                   | 0.37 (0.02)      | 0.34–0.43    |
| ADC \(10^{-3}\ mm^2/s\) | 1.06 (0.05)     | 0.95–1.23    |
| Length (mm)          | 59.8 (8.2)       | 38.5–80.6    |
| **CC genu**          |                  |              |
| FA                   | 0.37 (0.04) ***  | 0.31–0.55    |
| ADC \(10^{-3}\ mm^2/s\) | 1.32 (0.10) *** | 1.10–1.56    |
| Length (mm)          | 46.7 (10.1) ***  | 26.8–79.6    |
| **CC splenium**      |                  |              |
| FA                   | 0.40 (0.04)      | 0.34–0.51    |
| ADC \(10^{-3}\ mm^2/s\) | 1.36 (0.11) *   | 1.10–1.69    |
| Length (mm)          | 59.3 (13.1)      | 27.5–80.2    |

* FA fractional anisotropy, ADC apparent diffusion coefficient
***: significant difference \(p<0.001\) between genu and splenium of the CC

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### Table 3

|                      | GA | Birth weight | HC at birth | PMA at MRI | Weight at MRI | HC at MRI |
|----------------------|----|--------------|-------------|------------|---------------|-----------|
| **PLIC**             |    |              |             |            |               |           |
| FA                   | 0.02 | −0.07        | −0.02       | 0.48 ***   | 0.30 **       | 0.29 **   |
| ADC                  | 0.06 | 0.12         | 0.05        | −0.65 ***  | −0.48 ***     | −0.42 ***  |
| Length               | −0.13 | −0.09        | −0.03       | 0.42 ***   | 0.30 **       | 0.34 **   |
| **CC genu**          |    |              |             |            |               |           |
| FA                   | −0.03 | −0.15        | −0.35 **    | 0.17       | 0.07          | 0.10      |
| ADC                  | 0.13 | 0.23 *       | 0.20        | −0.53 ***  | −0.22 *       | −0.14     |
| Length               | −0.14 | −0.10        | −0.04       | 0.46 ***   | 0.35 ***      | 0.34 **   |
| **CC splenium**      |    |              |             |            |               |           |
| FA                   | −0.001 | −0.08        | −0.16       | 0.24 **    | 0.17          | 0.10      |
| ADC                  | −0.06 | 0.05         | −0.02       | −0.35 ***  | −0.19         | −0.21     |
| Length               | 0.09 | 0.05         | 0.33 *      | 0.43 ***   | 0.42 ***      | 0.52 ***  |

HC head circumference, GA gestational age, PMA post-menstrual age, * significant association: \(p<0.05\), ** significant association: \(p<0.01\), *** significant association: \(p<0.001\)
Counsell et al. randomly recruited 38 preterm infants and compared these with 8 healthy term born controls. The preterm infants were divided into a group with normal appearing WM and a group with DEHSI. Infants with overt WM injury were excluded from their study. The difference with our results may again be explained by the fact that we studied a group of unselected VPTI. It may also be hypothesised that the wide age range at imaging in our study group plays a role. However, as the results in the subgroup of infants undergoing imaging before 46 weeks were comparable to those in the whole group, we feel this will not have influenced our results. Possibly the unequal distribution of infants with normal/mildly, moderately or severely abnormal WM has played a more important role. However, when comparing the 18 subjects with normal/mildly abnormal WM with the 15 with severe WM injury we found no difference between these two groups. In a recent study Cheong et al. found a difference in FA in the PLIC in VPTI with extensive WM injury compared with VPTI with normal or focal WM injury using DTI region of interest measurements. In their study 39 infants had no WM signal intensity abnormalities, 59 infants had focal and 13 infants extensive WM abnormalities. The categorisation of WM injury we used is very similar to that of Cheong et al., but they included a large number of infants with normal appearing WM whereas that group was relatively small in our study.

Furthermore, in accordance with the study of Counsell et al. we did not identify any categorical clinical factors associated with DTI parameters.
For the CC we found significantly different DTI values between the genu and splenium. In the genu FA and ADC values were significantly lower, indicating that at this young age the genu is less developed than the splenium. In terms of development similar findings have been reported by Barkovich, who pointed out that the myelination of the splenium precedes that of the genu on conventional MRI [9].

One of the main limitations of our study is that, due to clinical circumstances not all VPTI were imaged around TEA. Therefore, although most were imaged before 46 weeks PMA (69/84), a wide age range existed. On the other hand, this enabled us to study the relation between PMA and DTI values. Another limitation of the present study is that we did not perform DTI in normal term infants. Therefore we were unable to compare the DTI values of the VPTI with those of a healthy control group.

We performed fibre tractography with commercially available software and with given default settings. Still, one of the limitations of fibre tracking is that in other studies different parameters or different tracking programs may be used, complicating the direct comparison of fibre tracking values between studies. Finally, as we performed fibre tractography of fibres passing through PLIC and CC, we only calculated DTI values in the WM tracts passing through these structures. We may therefore have missed a possible association between DTI parameters and either normal or injured WM.

In conclusion, our data show in VPTI imaged around or within 5 months of TEA, a strong association between DTI parameters of the PLIC and CC and PMA at imaging, but not between these values and GA. No association was found between DTI parameters of the PLIC and CC and the degree of WM injury or categorical clinical factors.

Although it has been reported that WM injury may result in atrophy of the PLIC and CC later on in life [3–7], our results indicate that in our cohort of VPTI, at this young age, the development of the PLIC and CC is ongoing and independent of the degree of prematurity or diffuse WM injury.

### Table 4
Stepwise linear regression models for DTI parameters (PLIC and CC), including the strongest contributing factors

| R     | p     | Predictors          | β    |
|-------|-------|---------------------|------|
| PLIC  |       |                     |      |
| FA    | 0.51  | <0.001              | PMA at MRI | 0.002 |
| ADC   | 0.63  | <0.001              | PMA at MRI | -0.008 |
| Length| 0.31  | < 0.01              | HC at MRI | 1.27   |
| CC genu |      |                     |      |
| FA    | 0.37  | 0.02                | HC at birth | -0.008 |
| ADC   | 0.55  | <0.001              | PMA at MRI | -0.013 |
| Length| 0.33  | <0.01               | PMA at MRI | 0.80   |
| CC splenium |     |                     |      |
| FA    | 0.23  | 0.04                | PMA at MRI | 0.002 |
| ADC   | 0.41  | <0.001              | PMA at MRI | -0.01  |
| Length| 0.69  | <0.001              | PMA at MRI | 1.52   | HC at birth | 3.02   |

**PLIC** postmenstrual age, **HC** head circumference

For the CC we found significantly different DTI values between the genu and splenium. In the genu FA and ADC values were significantly lower, indicating that at this young age the genu is less developed than the splenium. In terms of development similar findings have been reported by Barkovich, who pointed out that the myelination of the splenium precedes that of the genu on conventional MRI [9].

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### Table 5
DTI parameters for PLIC and CC and categorised WM injury (n=84)

|          | Normal-mild (n=18) | Moderate (n=51) | Severe (n=15) | Overall p value | Normal-Mild vs. Severe | Normal-Mild vs. Moderate | Moderate vs. Severe |
|----------|--------------------|-----------------|---------------|-----------------|------------------------|------------------------|---------------------|
| PLIC     |                    |                 |               |                 |                        |                        |                     |
| FA       | 0.38 (0.02)        | 0.37 (0.02)     | 0.37 (0.03)   | 0.27            | 1.00                   | 0.34                   | 1.00                |
| ADC      | 1.05 (0.05)        | 1.07 (0.05)     | 1.07 (0.04)   | 0.64            | 1.00                   | 0.73                   | 1.00                |
| Length   | 59.4 (8.0)         | 60.0 (7.0)      | 59.6 (11.9)   | 0.96            | 1.00                   | 0.76                   | 1.00                |
| CC genu  |                    |                 |               |                 |                        |                        |                     |
| FA       | 0.38 (0.05)        | 0.37 (0.04)     | 0.38 (0.04)   | 0.58            | 1.00                   | 1.00                   | 1.00                |
| ADC      | 1.30 (0.11)        | 1.32 (0.09)     | 1.33 (0.10)   | 0.65            | 1.00                   | 1.00                   | 1.00                |
| Length   | 45.6 (11.1)        | 47.4 (9.6)      | 45.6 (11.1)   | 0.73            | 1.00                   | 1.00                   | 1.00                |
| CC splenium |                |                 |               |                 |                        |                        |                     |
| FA       | 0.40 (0.04)        | 0.39 (0.03)     | 0.41 (0.03)   | 0.54            | 1.00                   | 1.00                   | 0.83                |
| ADC      | 1.32 (0.09)        | 1.36 (0.12)     | 1.39 (0.11)   | 0.16            | 0.17                   | 0.60                   | 0.84                |
| Length   | 62.7 (12.2)        | 59.3 (12.7)     | 55.2 (15.2)   | 0.27            | 0.31                   | 1.00                   | 0.84                |

**PLIC** posterior limb of the internal capsule, **CC** corpus callosum, **WM** white matter.

WM injury was categorised: normal/mildly abnormal, moderately abnormal and severely abnormal.
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