Reduced white matter connectivity in the corpus callosum of children with Tourette syndrome

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Background: Brain imaging studies have revealed anatomical anomalies in the brains of individuals with Tourette syndrome (TS). Prefrontal regions have been found to be larger and the corpus callosum (CC) area smaller in children and young adults with TS compared with healthy control subjects, and these anatomical features have been understood to reflect neural plasticity that helps to attenuate the severity of tics.

Method: CC white matter connectivity, as measured by the Fractional Anisotropy (FA) index from diffusion tensor images, was assessed in 20 clinically well-defined boys with Tourette syndrome and 20 age- and gender-matched controls.

Results: The hypothesis that children with TS would show reduced measures of connectivity in CC fibers was confirmed for all subregions of the CC. There was no significant interaction of TS and region. Reductions in FA in CC regions may reflect either fewer interhemispheric fibers or reduced axonal myelination. FA values did not correlate significantly with the severity of tic symptoms. Group differences in measures of connectivity did not seem to be attributable to the presence of comorbid ADHD or OCD, to medication exposure, or group differences in IQ.

Conclusion: Our findings of a reduced interhemispheral white matter connectivity add to the understanding of neural connectivity and plasticity in the brains of children who have TS.

Keywords: Tourette syndrome, brain development, brain imaging.

Tourette syndrome (TS) is characterized by the presence of chronic motor and phonic tics that fluctuate in severity (ICD 10/DSM IV)(American Psychiatric Association, 1994). The phenotypic spectrum of tics can range from simple movements or sounds to more complex and debilitating motor behaviors and vocalizations. Recent studies confirm the importance of genetic factors that predispose individuals to developing TS (Abelson et al., 2005; Pauls, 2003). Moreover, a genetically mediated relationship between TS and obsessive-compulsive disorder (OCD) has been established through family-genetic (Eapen, Pauls, & Robertson, 1993; Pauls, Leckman, Towbin, Zahner, & Cohen, 1986; Pauls, Raymond, Stevenson, & Leckman, 1991) and molecular genetic studies (Cuker, State, King, Davis, & Ward, 2004; State et al., 2003). TS children in clinical and epidemiological settings have high rates of comorbid OCD (Termine et al., 2006); attention-deficit/hyperactivity disorder (ADHD) and depression (Kurlan et al., 2002) are also commonly seen. The typical natural history of tic symptoms in TS is an increase in severity from childhood to early adolescence and, in most cases, an attenuation during or after puberty (Leckman et al., 1998; Pappert, Goetz, Louis, Blasucci, & Leurgans, 2003). More than 40% of all children with TS are free of tics by the age of 18 (Leckman et al., 1998). This natural history suggests that the brain develops mechanisms during adolescence that help to modulate the severity of tics (Spessot, Plessen, & Peterson, 2004).

Recent structural and functional brain imaging studies have improved our understanding of brain adaptation and neural plasticity (Pascual-Leone, Amedi, Fregni, & Merabet, 2005). In TS, reports of abnormalities in brain morphology have focused on the basal ganglia as an important part of corticostriato-thalamo-cortical (CSTC) circuits (Alexander, DeLong, & Strick, 1986), which are implicated in the underlying pathophysiology of TS (Peterson et al., 1993, 1998, 2003). Anatomical abnormalities of the basal ganglia in childhood have also been found to predict the severity of tics in early adulthood (Bloch et al., 2005). Cortical regions, on the other hand, seem to play an important role in the modulation and suppression of tics (Fredericksen et al., 2002; Peterson et al., 1998; Peterson, Staib et al., 2001; Serrien,
of corticical volume in children and adults with TS (Peterson, Staib et al., 2001) have revealed larger dorsal prefrontal cortex (DPFC) volumes in children with TS compared with control children, and the magnitude of this volumetric enlargement was associated with fewer tic symptoms. Adults with persistent TS, however, did not show this enlargement of prefrontal volumes, and in fact their volumes were smaller than in control adults, suggesting that their prefrontal cortices did not adapt successfully to the presence of tics during adolescence. In light of prior anatomical studies demonstrating a normal reduction in prefrontal gray matter density during childhood and adolescence (Sowell et al., 2003), we have speculated that the normal regressive processes of childhood and adolescence, including the pruning of axons, dendrites, and synapses in prefrontal regions, are slowed in individuals who successfully adapt to the presence of tics, presumably because these brain regions are activated prominently (Peterson et al., 1998) by the frequent need for children to suppress tics in social situations.

The overall midsagittal cross-sectional area of the corpus callosum (CC) has been reported to be smaller in subjects with TS (Peterson et al., 1994; Plessen et al., 2004) and to be correlated inversely in children with both symptom severity and prefrontal volumes (Plessen et al., 2004). These studies have thus suggested that interhemispheric axons in children with TS, together with prefrontal cortices, successfully reorganize in order to enhance regulatory control of motor and phonic tics. This interpretation is consistent with the known functional characteristics of the CC, which involves primarily transfer of information between the two hemispheres, as well as modulation of attention (Hugdahl, 1998), and inhibition of cortical activity (Borojejerdi, Topper, Folty, & Meincke, 1999) and plastic reorganization of the brain (Werhahn, Mortensen, Kaelin-Lang, Borojejjerdi, & Cohen, 2002). Transcallosal inhibition presumably involves GABAergic inhibitory interneurons, as the CC itself consists of glutamatergic excitatory fibers (Carr & Sesack, 1998; Conti & Manzoni, 1994).

Interhemispheric fiber tracts within the CC can be studied directly using diffusion tensor imaging (DTI), a magnetic resonance imaging (MRI)-based technique that measures the constrained diffusion of water, or the degree of anisotropy, within neuronal tissue (Ramnani, Behrens, Penny, & Matthews, 2004). It has been shown that DTI reliably detects white matter tracts and can be regarded as a tool to study connectivity between cortical regions (Basser, Mattiello, & LeBihan, 1994; Basser, Pajevic, Pierpaoli, Duda, & Aldroubi, 2000; Pierpaoli et al., 2001). The parameter used most widely to characterize the degree of anisotropy and degree of structural organization in the brain, particularly in clinical applications of DTI, is Fractional Anisotropy (FA), an index that ranges from 0 (isotropic diffusion, characteristic of the diffusion of unconstrained water within CSF) to 1 (complete anisotropic diffusion and maximal organization of tissue) (Hasan, Alexander, & Narayana, 2004). The FA has been shown to correlate with histological markers of myelination in newborns, toddlers, and adults (Winberger et al., 1995; Huppi et al., 1998; Klingberg, Vaidya, Gabrieli, Moseley, & Hedehus, 1999), which likely contributes to the increasing speed of neural transmission with advancing development (Paus et al., 1999). Based on our prior report of a smaller CC area in children with TS (Plessen et al., 2004), the present study investigated the hypothesis that a smaller CC in TS subjects is the consequence of either fewer interhemispheric fibers or reduced myelination, which should result in reduced FA values within the CC. By further subdividing the CC into five anatomically discrete regions, we investigated differences in the FA index for different CC regions and possible interactions with a diagnosis of TS.

Methods

Subjects

TS subjects were recruited from the Department of Child and Adolescent Psychiatry at the Haukeland University Hospital, University of Bergen, Norway and from outpatient clinics in the greater Bergen area in the western part of Norway. All met DSM-IV criteria for a diagnosis of TS (American Psychiatric Association, 1994). Healthy control children (HC) were recruited by randomly contacting local schools in the same geographic area. The controls were matched for age and gender with the patient group. Written informed consent was obtained from all participants and the study was approved by Regional Committee for Medical Research Ethics, West-Norway.

Exclusion criteria for the control group included a lifetime history of Tic Disorder, OCD, ADHD, or a current DSM-IV Axis I disorder. Additional exclusion criteria for both groups were epilepsy, head trauma with loss of consciousness, former or present substance abuse, or an IQ below 70, as measured with the WISC-III (Wechsler, 1996).

Psychiatric diagnoses were established using the Schedule for Affective Disorders and Schizophrenia for School-Age Children–Present and Lifetime version (Kauffman et al., 1997) and a best-estimate consensus procedure that considered all available study materials (Leckman, Sholomskas, Thompson, Belanger, & Weissman, 1982). OCD symptoms were quantified using the Children’s Yale-Brown Obsessive Compulsive Scale (Goodman et al., 1989; Scahill et al., 1997), and the severity of tics was rated with Yale Global Tic Severity Scale (YGTTSS) (Leckman et al., 1989). Socioeconomic status (SES) was estimated by measuring the parental level of education in four categories, dependent on their school and higher education (JAAACP, 2005).

Sample size was limited to the first 20 subjects in each group who were referred consecutively to the study and who met the criteria for inclusion. Only two girls with TS were referred to the study, and they experi-
enced severe tics during the scanning session that produced imaging artifacts requiring exclusion of these subjects. Thus, the reported data consist of two male groups: 20 TS and 20 HC boys, 9 to 17 years of age. The groups were of comparable age (TS = 13.6 years, ±1.9; HC = 13.4 years, ±2.4; t = −3; p = .77) and SES. The diagnostic interview revealed that five of the subjects in the TS group in addition had combined-type ADHD, four others had comorbid OCD, and none had both conditions. Participants were right-handed as measured by the Edinburgh handedness inventory (Oldfield, 1971), except for two left-handed individuals in each group.

At the time of MRI scanning, nine subjects in the TS group were taking medication, either neuroleptics (n = 4), alpha agonists (n = 2), selective serotonin uptake inhibitors (n = 1), or stimulants (n = 2). No subjects were on combinations of medications. HC subjects were free of any psychotropic medications. Tic severity at the time of scan in the TS group was 11.4 ± 2.9 for motor and 9.2 ± 3.4 for phonic tics, and at for lifetime-worst ever was 15.5 ± 5.1 for motor and 13.7 ± 5.5 for phonic tics (possible range 0 to 25 in each category). The groups differed in verbal IQ (TS = 94.4 ± 11.4; HC = 104.4 ± 10.5; t = 2.9; p < .006), performance IQ (TS = 95.6 ± 10.8; HC = 106.1 ± 12.1; t = 2.9; p < .006), and full-scale IQ (TS = 94.5 ± 10.2; HC = 105.7 ± 9.2; t = 3.6; p < .001).

MRI scanning and image-analysis

MR-images were acquired on a Siemens Symphony, 1.5 Tesla scanner. Head positioning was standardized using canthomeatal landmarks. T1-weighted, sagittal 3D volume MP-Rage images were acquired for all subjects, with repetition time (TR) = 1910 ms, echo time (TE) = 3.93 ms, flip angle (FA) (θ) = 4°, image matrix = 256 × 192, field of view (FOV) = 256 mm, slice thickness = 1 mm with 176 contiguous slices. The DTI data were acquired with single shot spin echo EPI sequence in axial and in sagittal acquisition, with TR = 4000 ms, TE = 96 ms, 128 × 128 matrix and 240 mm FOV, at a nominal resolution of 1.875 × 1.875 × 4 mm³. The diffusion weighting was b = 1000 s/mm² in 6 noncolinear directions (8 averaged means per slice), in addition to one reference (b = 0) image per slice. We focused on acquiring data of high quality using behavioral relaxation techniques and eventually repeating series containing movement artifacts.

MR image analysis for overall CC area, using the T1-weighted MP-Rage images, was performed using Analyze 6.0 software (Rochester Minnesota) on a Linux Workstation. Raters were blind to subject characteristics and group assignments. Each MR dataset was realigned to the midsagittal slice, which was identified using local curvature maxima. To compare FA values for CC subregions that carry fiber tracts belonging to different cortical regions, the CC was subdivided into seven segments according to a previously published method (Whitfield, 1989), except for summing the two most anterior and two most posterior regions (Luders et al., 2005) (see Figure 1). Also, to enhance the stability of FA measures further, mean FA values from the CC regions (Moeller et al., 2005) from three midsagittal slices (midsagittal and the two adjacent parasagittal slices 4 mm bilaterally off the midline) were summed into a single midsagittal FA index, after assuring a normal distribution by visual inspection. Table 1 shows mean values and SD for FA measures for each group.

The main analyses reported here were complemented by a co-registration and spatial normalization of the DTI images, which were intended to confirm findings based on FA values averaged across all voxels in each CC subregion. We further confirmed our findings of differences in average FA values between groups through a comparison of FA values at each pixel of the CC in a common template space. To compute group-average FA values at all voxels, we co-registered, normalized, and reoriented the DT image of each subject into the common space by a two-step procedure. In the first step, we co-registered and nonlinearly warped the DT image of each subject to its anatomical MR image (Xu, Mori, Shen, van Zijl, & Davatzikos, 2003). In the second step, we co-registered and normalized the MR image of each subject to the reference image (Bansal, Staib, Whitman, Wang, & Peterson, 2005). The reference image was an anatomical MR image of a randomly chosen single health subject. We concatenated the two nonlinear deformations estimated in the two steps and used the concatenated deformation to co-register and normalize the DT image of each subject into the template space. Tensors were correctly reoriented into the template space by using the method of Procrustean estimation (Xu et al., 2003). Once DT images for each subject had been normalized and reoriented into the common template space, we computed the average FA values at each pixel in the template space. We then displayed and compared visually the gray-scale encoded average FA values for the two groups.

Statistical analyses

All statistical procedures were performed in SAS v. 8.2 (SAS Institute Inc., Cary NC) or SPSS v. 13 (SPSS, 1999). The a priori hypothesis was tested using a mixed-model regression analysis (PROC MIXED) with repeated measures over a spatial domain (the five regions of the CC) with a first order autoregressive (AR1) model of the covariance structure. The model included the within-subjects factors ‘CC region’ with 5 levels. ‘Diagnosis’ (TS or controls) was a between-
Subjects factor. ‘Age’ was included as a covariate in the model. In addition to the independent variables described above, all two and three-way interactions of ‘group’ (TS and HC), ‘region’, and ‘age’ were tested. Interactions that were not statistically significant were hierarchically eliminated via backward stepwise regression, with the constraint that the model at each step had to be hierarchically well formulated (i.e., all possible lower order terms had to be included in the model, regardless of their statistical significance) (Morrell, Pearson, & Brant, 1997). Whole-brain volume was not included in the model, as it has been shown that FA values are not influenced by brain volume (Schulte, Sullivan, Muller-Oehring, Adalsteinsson, & Pfefferbaum, 2005).

Potential confounds. In order assess the effects of comorbidity and medication on our findings in the TS group, the analysis was repeated while restricted to all subjects with TS, excluding those with comorbid ADHD (remaining *n* = 15) or OCD (remaining *n* = 16), and excluding those on medication (remaining *n* = 11), compared with the original control sample (*n* = 20). Because the TS and HC groups significantly differed in IQ, this also was considered as a covariate in the mixed model.

Association with symptom severity. Correlations of FA values with symptom severity in the TS group were investigated by calculating the Pearson correlation coefficient *r* for each CC subregion, and by visual inspection of scatterplots of FA values and symptom severity. Partial correlations were performed by using age and IQ as a covariate.
Results

The test for fixed effects in a mixed model revealed a significant main effect of group ($F_{1,38} = 7.60; p < .009$), demonstrating the hypothesized differences in FA values for the CC in individuals with TS compared to controls (see Table 2).

Other covariates in the final model were region of the CC ($F_{4,155} = 9.74; p < .001$) and age ($F_{1,38} = 2.17; p = .15$). Age of the subjects was conservatively retained in the model, due to biologic plausibility. Group differences were not affected whether age was retained in the model or not. No significant interaction between TS and subregion of the CC (at the point of elimination n.s.) was detected, indicating that the group influence on FA values did not vary according to CC subregion (see also Figure 2). Also, the group-by-region-by-age interaction was not significant (at the point of elimination n.s.), indicating that the findings were stable across the age range of children studied.

Post hoc analyses. A post-hoc assessment of the origin of this difference between groups, using a test of differences in least-square means, indicated that FA values in the TS group were lower than in controls (mean FA .80 vs. .82; $t_{38} = 2.76; p < .009$).

Potential confounding variables. The model remained stable, even when subjects with comorbid diagnoses of ADHD ($n = 5$) and OCD ($n = 4$) and those taking medication ($n = 9$) were excluded from the analyses subsequently (Table 2). Also IQ as a covariate did not show a significant influence on FA values, whether verbal (at the point of elimination n.s.), performance (at the point of elimination n.s.) or full-scale IQ (at the point of elimination n.s.). Moreover, when excluding subjects with comorbid ADHD, verbal IQ no longer differed significantly between the two groups, whereas the finding of lower FA values in the TS group remained stable (see Table 3).

Associations with tic severity. Correlations of tic severity with FA values in each of the five CC regions were not significant (region 1 $r = 0.06$; region 2 $r = 0.25$; region 3 $r = 0.07$; region 4 $r = 0.28$; region 5 $r = 0.14$). However, symptom severity did correlate positively, even if not at a level of statistical significance, with FA values in the four anterior CC subregions (see Figure 3).

Coregistration and normalization of DTI images. In normalized space, FA maps for the TS group showed smaller FA values in the CC (visualized as lower image intensity within the CC in Figure 4), and hence confirmed the findings derived using the mean values without prior normalization.

To visualize FA differences further, fiber-tracking was performed by placing one region of interest in the anterior CC of one randomly chosen individual in each group (Figure 5).

Discussion

White matter microstructure in TS children showed less directional organization (lower FA values) compared with control children, within all subregions of the CC. Correlations of FA values with measures of current tic severity were not significant. Reduction of FA values in the TS group confirmed our a priori hypothesis. We verified findings of lower FA values by also presenting group FA values in the CC with the DTI maps spatially normalized to a template.
Lower FA values in the TS group likely represent a reduction of the number of axons, reduction in their degree of myelination, or a disruption of their structural organization within the white matter of the CC. It is not possible to determine the underlying biological ultrastructure of the group.

### Table 3

<table>
<thead>
<tr>
<th>Measure</th>
<th>TS vs. HC (n= 40)</th>
<th>TS(ADHD excluded) vs. HC (n = 35)</th>
<th>TS (OCD excluded) vs. HC (n = 36)</th>
<th>TS (Not medicated) vs. HC (n = 31)</th>
</tr>
</thead>
<tbody>
<tr>
<td>df</td>
<td>t-value</td>
<td>p</td>
<td>df</td>
<td>t-value</td>
</tr>
<tr>
<td>Verbal IQ</td>
<td>38</td>
<td>2.89</td>
<td>.006</td>
<td>33</td>
</tr>
<tr>
<td>Performance IQ</td>
<td>38</td>
<td>2.91</td>
<td>.006</td>
<td>33</td>
</tr>
<tr>
<td>Full-scale IQ</td>
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<td>3.63</td>
<td>.001</td>
<td>33</td>
</tr>
<tr>
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<td>.014</td>
<td>31</td>
</tr>
<tr>
<td>Perceptual Organisation Index</td>
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<td>n.s.</td>
<td>31</td>
</tr>
<tr>
<td>Freedom From Distractability Index</td>
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<td>n.s.</td>
<td>31</td>
</tr>
<tr>
<td>Processing Speed Index</td>
<td>36</td>
<td>2.41</td>
<td>.021</td>
<td>31</td>
</tr>
</tbody>
</table>

### Figure 3

Correlation FA values with symptom severity of current motor symptoms in the TS group

### Figure 4

Co-registered DTI and anatomical images for the HC (a) and TS (b) group. The averaged CC FA map is shown on the co-registered midsagittal slice and intensities in white matter are visualized.

(Figure 4). Lower FA values in the TS group likely represent a reduction of the number of axons, reduction in their degree of myelination, or a disruption of their structural organization within the white matter of the CC. It is not possible to determine the underlying biological ultrastructure of the group.
difference with the present investigation. However, given extensive prior evidence for the presence of compensatory neuroplasticity within prefrontal cortices of children with TS, we suspect that the most likely cause of reduced FA values in the CC in TS children is the presence of fewer callosal axons connecting the cerebral hemispheres, although we cannot exclude other explanations at this point. Fewer interhemispheric axons would likely reduce transcallosal inhibition of cortical neurons and thereby enhance prefrontal functioning and ultimately support frontostriatal control of tic symptoms in children with TS. Although the neurotransmitter specifications of most transcallosal axons are glutamatergic and therefore excitatory, the CC exerts primarily inhibitory effects on cortical functioning that seem to be mediated by synapses of CC axons onto GABAergic inhibitory interneurons within cortical gray matter (Chen, 2004). Fewer transcallosal axons presumably provide less inhibitory influence on prefrontal cortices, and therefore would enhance the net output of prefrontal cortices which are thought to regulate tic behaviors. Also, in healthy individuals there is growing evidence for a high degree of interhemispheric inhibition of voluntary movements, possibly to counteract the production of mirror movements, yet it still remains to be determined which cortical regions contribute to the interhemispheric inhibitory interactions (Duque et al., 2005). The importance of inhibitory processes for neuroplasticity has been recognized lately (Kandler, 2004). This model is consistent with prior reports of cortical hyperexcitability in children with TS, which presumably is attributable to a reduction in cortical GABAergic interneurons (Ziemann, Paulus, & Rothenberger, 1997). Thus the CC seems to be a component of a larger compensatory neuroregulatory system in TS involving prefrontal control on motor activity within the basal ganglia portions of CSTC circuits.

These postulated neuroplastic effects within the CC and frontal cortices are presumably mediated through altered rates of pruning during CNS development. Pruning of axons within the CC and of axons, dendrites, and synapses within prefrontal cortices is a prominent feature of normal CNS development (Hua & Smith, 2004). Given their smaller CCs that seemingly contain fewer axons, normal axonal pruning is likely accelerated in children with TS. Their larger prefrontal cortices, an exaggeration of the higher prefrontal gray matter density present in younger compared with older healthy children (Sowell et al., 2003), suggests that normal pruning in frontal cortices is reduced in children with TS. These alterations in rates of pruning during development likely reflect the effects of an activity-dependent plasticity that is induced by the intense activation of prefrontal cortices that accompanies the suppression of tic behaviors (Peterson et al., 1998), a frequent and nearly ubiquitous occurrence for most children with TS. Another possible model to understand a reduction of interhemispheric connectivity could be the direct involvement of CC fibers in tic suppression. Glutamatergic neurons within the CC do not only mediate inhibitory functions, thus we cannot exclude that a lack of modulatory excitatory impulses to the motor cortices directly may be involved in tic suppression. The fact that FA reduction is found in all regions of the CC could also point to the involvement of the CC in tic generation as opposed to a local compensatory phenomenon. A recent DTI study (Huang et al., 2005), however, suggests that cortical fibers originating from one cortical region spread widely along the midsagittal CC axis. Especially frontal fibers spread from the most anterior region of the CC along the whole callosal body. Hence, the reduction of FA values along all regions may still be contributed to compensatory mechanisms primarily located in the prefrontal cortices.

It should, however, be noted that almost all the cited studies, including the present study, dealing with brain morphology in TS had a cross-sectional study design, which only allows limited conclusions concerning developmental trajectories and cannot infer about causation or about the temporal sequencing in the development (Kraemer, Yesavage, Taylor, & Kupfer, 2000; Peterson, 2003). The suggested models should thus be further tested in longitudinal studies. Moreover, the integration of multimodal MR imaging techniques in the same sample, e.g., the inclusion of volumetric prefrontal

Figure 5 Fibre-tracking in two representative children, (a) a healthy child; (b) a child with TS. The seeds for fiber tracking were placed on an axial slice within the CC.
and basal ganglia together with DTI measurements and eventually functional MRI studies in larger samples, would help to further confirm or refute the suggested models.

Although the cross-sectional areas of the CC did not differ significantly between TS and HC groups in this sample, overall average size of the CC was smaller in the TS group, consistent with our findings in a much larger sample of TS children. There was no significant group-by-region interaction that pointed to any of the CC regions being specifically involved in the TS condition.

Correlations of FA values with symptom severity were not statistically significant. Positive associations of FA values with current tic severity in two CC regions tend to support our hypothesis that lower FA values in the TS group reflect a compensatory, neuroplastic response that helps to modulate tic severity. The absence of significant correlations of FA values with age may attest to the stability of the observed findings in the examined age range.

Although IQ measures were significantly lower in the TS children, covariance analyses provided evidence that IQ differences across groups were not responsible for the lower FA value in the TS group. When excluding subjects with comorbid ADHD, verbal IQ was no longer significantly different between the TS and HC groups, consistent with prior evidence that comorbidity with ADHD entails academic difficulties in individuals with TS (Abwender et al., 1996; Erenberg, Cruse, & Rothner, 1987; Peterson, Pine, Cohen, & Brook, 2001; Schuerholz, Baumgardner, Singer, Reiss, & Denckla, 1996).

Comorbidity with ADHD or OCD, and medication use did not seem to contribute to group differences in FA values. When these effects were controlled statistically, and when subjects with these conditions were separately excluded from the statistical analyses, findings of significantly lower FA values in the TS group persisted, despite the loss of statistical power that inclusion of fewer subjects entailed (see Table 2).

This is, to our knowledge, the first DTI investigation of individuals with TS. It confirms the hypothesized reduction in anatomical connectivity of the CC in children with TS. Reduced connectivity in the CC provides further supporting evidence for the presence of neuroplastic responses in the CC and other regions in the brains of these children, presumably in the service of enhancing prefrontal functioning and the regulation of tic behaviors.

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