Multiparametric magnetic resonance imaging analysis of the corticospinal tract in multiple sclerosis

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Background/purpose: Muscle weakness is an important feature of multiple sclerosis and is responsible for much of the disability associated with that condition. Here, we describe the quantitative magnetic resonance imaging (MRI) attributes of the major intracerebral motor pathway – the corticospinal tract – in multiple sclerosis. To do so, we develop an intuitive method for creating and displaying spatially normalized tract-specific imaging data.

Methods: In 75 individuals with multiple sclerosis and 29 healthy controls, the corticospinal tracts were reconstructed from diffusion tensor imaging at 3 T. Multiple MRI indices – T2 relaxation time; fractional anisotropy; mean, longitudinal, and transverse diffusivity; and magnetization transfer ratio – were examined within the reconstructed tracts. Spatially normalized tract profiles were created to compare, across subjects, the variation in MRI index as a function of tract position.

Results: Each index’s tract profile had a characteristic shape. Individual subjects had markedly abnormal tract profiles, particularly at lesion sites. On average, tract profiles were different between patients and controls, particularly in the subcortical white matter and corona radiata, for all indices examined except for fractional anisotropy. Magnetization transfer ratio was further decreased in subjects with secondary progressive disease. Tract asymmetry was increased in multiple sclerosis compared to controls.

Conclusion: Multiparametric MRI allows rapid detection, localization, and characterization of tract-specific abnormalities in multiple sclerosis. Tract profiles bridge the gap between whole-brain imaging and the interrogation of individual, functionally relevant subsystems.

Keywords: Multiple sclerosis; Corticospinal tract; Pyramidal tract; Diffusion tensor imaging; Magnetization transfer imaging; Tract profiles

Introduction

Multiple sclerosis (MS) is a central nervous system disorder, with protein manifestations and multiple subtypes, that can lead to a wide spectrum and degree of disability. Among the functional systems that it frequently affects is the pyramidal motor system, resulting in weakness of one or more limbs or the facial muscles. Abnormalities in this system contribute to Kurtzke’s Expanded Disability Status Score (EDSS) (Kurtzke, 1983) and can result in clinically relevant disability. Because both its function and location within the brain are well defined, the pyramidal system serves as a model for the detection and clinicoradiologic assessment of functional disability in MS.

In this paper, we discuss the magnetic resonance imaging (MRI) characteristics of a major portion of the pyramidal system, the corticospinal tract (CST), in MS. In doing so, we have two primary objectives: (1) to facilitate the detection and monitoring of CST-specific abnormalities in individual patients; and (2) within an MS cohort, to assess the typical degree and locations of those abnormalities. In the first instance, we hope to be able to evaluate objectively the state of an individual’s disease and ultimately to monitor its progression over time, or, hopefully, its recovery after specific therapy. In the second instance, we expect to gain a deeper understanding of the ways in which MS can affect the motor system and ultimately to track the effects of novel therapeutic agents on a population level.
The imaging appearance of the CST in MS has been studied previously, although not over its entire intracranial course or for a wide variety of MRI indices in the same subject. MS plaques can affect the CST, causing typical changes of increased T2-weighted signal, decreased T1-weighted signal, and enhancement following contrast administration. There are MRI abnormalities even in portions of the CST that are not overtly affected by MS plaques, including increased T2-weighted signal, which has been interpreted to reflect Wallerian degeneration (Simon et al., 2000). With diffusion tensor imaging (DTI), a newer quantitative MRI technique, MS plaques typically demonstrate increased mean diffusivity (MD) and decreased fractional anisotropy (FA) (Tievsky et al., 1999; Werring et al., 1999). Similar abnormalities have also been reported along the CST specifically (Lin et al., 2007; Wilson et al., 2003). Additionally, levels of N-acetyl aspartate, a measure of axonal integrity, are decreased within the internal capsule, through which the CST passes (Lee et al., 2000). However, the degree to which these MRI abnormalities relate to axonal pathology along the CST remains unclear (DeLuca et al., 2004).

MRI abnormalities along the CST have also been studied in other neurological diseases. In neuromyelitis optica, a neuroimmunological disorder related to MS, decreased FA and increased MD have been observed in the cerebral peduncles (Lin et al., 2006). Similar patterns have been described in other neurological diseases, including stroke (Werring et al., 2000) and amyotrophic lateral sclerosis (Toosy et al., 2003). In the latter, the magnetization transfer ratio (MTR), which is influenced by protons bound to macromolecules such as myelin, has been reported to be decreased in the CST but not in nearby white matter (Tanabe et al., 1998).

In this paper, we use DTI with fiber tracking (Mori et al., 1999, 2005; Mori and van Zijl, 2002) to identify the intracerebral portion of the CST. This technique, while still in relative infancy and not in general clinical use, is emerging as a powerful tool for assessing pathway-specific abnormalities in neurological disease. Within the identified tracts, we measure various quantitative MRI indices derived from DTI and additional acquisitions (T2 and magnetization transfer) that are anatomically coregistered to the DTI data. We calculate each index as a function of position within the tract, referring to plots depicting their spatial variation as “tract profiles.” Tract profiles allow localization of focal tract abnormalities in individual subjects. We also consider the asymmetry between the right and left tracts.

This study builds upon previous work (Reich et al., 2006) that describes our methods for identifying, characterizing, and detecting asymmetry within the CSTs of healthy volunteers. In this paper, we focus on the presentation and discussion of data from our subjects with MS.

Materials and methods

Magnetic resonance imaging

We studied 75 individuals with MS: 43 individuals with relapsing remitting MS (RRMS; median age: 40; range: 24–60; 30 women and 10 men); 22 individuals with secondary progressive MS (SPMS; median age: 50; range: 40–67; 13 women and 9 men); and 10 individuals with primary progressive MS (PPMS; median age: 54; range: 44–67; 6 women and 4 men). We also studied 29 healthy controls (median age: 33; range: 22–63; 20 women and 9 men). All imaging protocols were accepted by the Institutional Review Boards at Johns Hopkins University and the Kennedy Krieger Institute, where scanning was done, and subjects were required to document their informed consent to participate in the study.

Imaging was done on a 3 T MRI scanner (Philips Medical Systems, Best, The Netherlands). A full description of our scanning protocol, as well as detailed results form our cohort of healthy volunteers, can be found in a previous publication (Reich et al., 2006). Note that the healthy controls in the present study include those subjects and 9 additional ones who were scanned after the original study was submitted for publication.

DTI on all subjects used spin echo, single shot, echo planar sequences and a sensitivity encoding (SENSE) reduction factor of 2.5. We obtained axial diffusion weighted images in 32 non-coplanar gradient directions with a nominal b-value of 700 s/mm². We also obtained a scanner average of 5 minimally diffusion-weighted scans with b = 33 s/mm². We collected data in isotropic 2.2 mm voxels, reconstructing to an in-plane resolution of 0.83 mm, and covered nearly the entire brain from the cervicomedullary junction to just below the vertex. We repeated the 3 min and 38 s sequence twice to improve the signal-to-noise ratio, and all data were used in the tensor calculation.

We coregistered all images (from the DTI acquisitions as well as from the other acquisitions described below) to the first minimally diffusion-weighted scan using the Automatic Image Registration (AIR) algorithm (Woods et al., 1992) with a 6-parameter rigid-body transformation, and we corrected the gradient directions for the rotational component of the transformation. We then estimated the diffusion tensor in the standard fashion (Basser et al., 1994) and calculated maps of fractional anisotropy (FA), mean diffusivity (MD), longitudinal diffusivity (λL), and transverse diffusivity (λT) (Basser and Pierpaoli, 1999). We also created color-coded maps from the principal eigenvectors, weighted by the fractional anisotropy (Pajevic and Pierpaoli, 1999). These analyses were performed in DtiStudio (Jiang et al., 2006), as well as with custom software purpose – written in Matlab (The Mathworks, Natick, MA).

Additional MRI acquisitions were obtained on subsets of the subjects scanned with DTI as the full protocol could not be obtained on every subject due to scanning time constraints and subject discomfort. On 66 individuals with MS and 26 controls, we acquired axial, double echo, turbo spin echo scans (TR=4158 ms; 3 min and 27 s) to visualize proton density (TE=28.2 ms) and T2 weighted (TE=80 ms) images, and to estimate absolute T2 relaxation time. From 63 subjects with MS and 24 controls, we acquired echo planar axial 3D spoiled gradient echo images, with and without 1.5 kHz off-resonance preparation, for the calculation of MTR (TR=65 ms, TE=15 ms, 3 repeats, 1 min and 41 s each) (Smith et al., 2006). Further scan details are provided elsewhere (Reich et al., 2006).

Fiber tracking

We used the DTI datasets to obtain 3D reconstructions of the CSTs using the fiber association by continuous tractography (FACT) method (Mori et al., 1999, 2005; Mori and van Zijl, 2002). Briefly, tracts were reconstructed using FA and the principal eigenvector of the diffusion tensor. Whole-brain seeding was used with predefined thresholds of FA and principal eigenvector turning.
angle (0.13 and 40°, respectively). The relatively low FA cutoff is permissive in that it fails to reject many spurious fibers, but the choice is necessary because FA is typically decreased within MS plaques (Tievsky et al., 1999; Werring et al., 1999), and we wanted to be able to track through those plaques to the extent possible.

To compensate for this choice and to reject spurious fibers, we chose multiple restrictive ROIs to limit the reconstructed CSTs to their known anatomical course, as described in a previous publication (Reich et al., 2006). One of us (DSR) drew axial ROIs around the entire caudal medulla; the ipsilateral anterior pons; and the ipsilateral cerebral hemisphere at the level of the subcortical white matter. We selected for further analysis the fibers that passed through all of these ROIs, traversing the entire intracranial CST from the cortex to the medulla. Rare spurious fibers passing through the contralateral anterior and bilateral posterior pons, and other fibers that clearly fell outside the major portion of the CST, were then manually excluded.

**Tract profiles**

We used tract profiles to depict the variation in each MRI index (FA, MD, λ₁, λ₂, MTR, and T2) as a function of location along the CST. This was done simply by plotting the mean of the index, within a specified spatial window, against the axial slice number (a convenient choice for the CST, which generally runs perpendicular to the axial plane). In order to determine how a given subject compares to other subjects, we applied a spatial normalization procedure to the tract profiles. Such a normalization is necessary because brain and tract shape change from one person to the next, so that axial scan positions are not constant across subjects.

The spatial normalization proceeded as follows. We divided each CST into six segments (of approximately equal length) according to anatomical landmarks defined on axial MRI sections of the color-coded principal eigenvector maps. These six segments correspond approximately to the subcortical white matter, corona radiata, internal capsule, midbrain, pons, and medulla, which are denoted by abbreviations in the figures. Due to inconsistencies in defining the imaging volume across subjects, data from the medulla were only obtained for 56 (75%) of the subjects with MS and 18 (62%) of the controls.

The seven boundary points of the six segments of the CST are illustrated in Fig. 1. Each of the segments was approximately 2 cm long, with a wide range of variation. Segments were then divided into 20 equal subsegments, each about 1 mm long. MRI indices were boxcar averaged within a sliding window 4 subsegments wide and then plotted against subsegment number to obtain the tract profile. Profiles from each subject were compared to the average profile obtained from controls.

**Tract profile statistics**

Within each segment of the CST, and across the entire tract, we compared median MRI indices between subjects with MS and controls using multiple linear regression analysis, accounting for the contributions made by age, sex, and number of reconstructed fibers per tract. Subject group and sex were considered categorical variables in the regression model, whereas age and number of reconstructed fibers were considered continuous variables. The significance level for each variable was the $p$-value corresponding to the partial Pearson correlation coefficient for that variable, holding the other variables constant. Statistical analyses were performed in Stata (StataCorp LP, College Station, TX).

To assess whether a particular median MRI index within any segment or across the entire tract was abnormal, we compared that index to the distribution obtained from controls. For this purpose, we used repeated results from the controls if more than one scan was available. The number of scans per control was as follows: 1 scan $(n=15)$; 2 scans $(n=9)$; 3 scans $(n=4)$; 4 scans $(n=1)$. In order to avoid bias due to repeated measurements from...
the same individual, results from subjects scanned more than once were weighted by the inverse of the number of scans for that subject (Taylor et al., 1996). A result was considered abnormal if it was below the 5th or above the 95th percentile of results from controls. Every individual contributed two tracts, from the left and right CSTs, so the number of indices in each control distribution was twice the number of individuals. Because of the exploratory nature of this work, we made no specific correction for multiple comparisons, and we simply report significance as p values.

Asymmetry analysis

To quantify the differences between the right and left CSTs, we used an asymmetry index, described in a previous publication (Reich et al., 2006), and defined (for the jth subsegment) as:

\[ A_j = \frac{I_{R,j} - I_{L,j}}{I_{R,j} + I_{L,j}} \]

where \( I_{R,j} \) is the index of interest derived from the jth subsegment of the right CST and \( I_{L,j} \) is the corresponding index from the jth subsegment of the left CST. \( A_j \) can range from -1 (maximum asymmetry with the index on the right equal to 0) to 1 (maximum asymmetry with the index on the left equal to 0); \( A_j = 0 \) corresponds to equality between the indices on the two sides. Total tract asymmetry was defined as the root mean square asymmetry across the entire tract, \( \sqrt{\sum A_j^2} \), and ranges from 0 to 1. We assessed the significance of tract asymmetry by the procedures described in the previous paragraph, comparing the observed asymmetry value to the distribution obtained from healthy controls.

Results

Sample tract profiles

We find a range of abnormalities in the CSTs of individuals with MS, as illustrated in Fig. 2 for three representative examples. Each example highlights abnormalities in one or two of the MRI indices. The first subject (panel A) is a 41-year-old woman with RRMS with no clinical weakness on either side. Her MRI reveals many confluent areas of T2-weighted signal abnormality, some of which are seen in the FLAIR image. She had no lesions that enhanced with gadolinium at the time of examination. Both CSTs pass through areas of T2-weighted hyperintensity as they traverse the periventricular white matter. These areas correspond to loci of increased transverse diffusivity (λ⊥; second image, panel A). Bilaterally increased λ⊥ is seen to better advantage in the tract profile (graph, panel A), which shows the variation in that index as a function of normalized distance along the tract. The area of greatest abnormality (red arrow) is in the periventricular white matter, which is a typical site of MS plaques.

The images in panel B come from a 32-year-old woman with RRMS with bilateral extremity weakness. This individual has multi-level tract abnormalities. In the top row of panel B, FLAIR and MTR images through the brainstem are shown, together with the MTR tract profile; in the bottom row of panel B, the second image and tract profile depict mean diffusivity (MD). There is an area of focally decreased MTR in the midbrain (dashed arrow) as well as a long segment of decreased MTR in the corona radiata and subcortical white matter. MD is normal at the level of the midbrain and focally abnormal at the junction of the corona radiata and subcortical white matter (solid arrow).

Panel C shows data from a 52-year-old man with PPMS, manifested primarily by spasticity and impaired ambulation. His MRI reveals marked periventricular signal abnormality without evidence of enhancement following gadolinium administration. In the corona radiata, we find a segment of elevated FA (arrow), more on the right than on the left. This elevation is due to increased longitudinal diffusivity (λ‖; bottom row) with normal or minimally decreased transverse diffusivity (λ⊥; not shown). Unchanged or even mildly elevated FA in this region is a common and somewhat surprising finding in our subjects, which is addressed further in the Discussion.

Average tract profiles

Just as it enables rapid localization of the portions of the CST that are abnormal in each individual with MS, the tract profile analysis also reveals patterns of abnormality across our cohort of subjects with MS. This is shown in Fig. 3, in which average tract profiles from subjects with MS are compared to average tract profiles from controls. Across the entire tract, there are significant differences in MD, λ⊥, λ‖, and MTR. The difference between subjects with MS and controls is most pronounced in the periventricular zone – which includes portions of the segments labeled as subcortical white matter and corona radiata – where, on average, MS patients have significantly elevated MD, λ‖, λ⊥, and absolute T2, and significantly decreased MTR. The region of elevated λ‖ is more diffuse than the region of elevated λ⊥. Perhaps surprisingly, FA is not significantly decreased in the MS CST.

Overall, we find very few significant differences among average tract profiles for the three MS subtypes. This is illustrated, for RRMS and SPMS, in Fig. 4; differences are even smaller for the other pairwise group comparisons. Although there is a barely significant difference between the two groups for MD across the entire CST, the most prominent finding is lower MTR in the internal capsule and midbrain in SPMS. MTR in this portion of the tract is slightly decreased in all subjects with MS compared to controls (Fig. 3), but not significantly so. Notably, however, there are no significant subgroup differences in the periventricular zone, where most of the abnormalities that distinguish subjects with MS from controls are found.

Abnormality of individual tract profiles

Fig. 5A demonstrates that between 20% and 50% of MS CSTs are significantly abnormal for each MRI index except for FA, where the percentage is close to the 5% expected by chance alone. Normal cutoffs are derived nonparametrically from the control data as described in Materials and methods. Note that the size of our control population does not allow us to control this analysis for age, sex, or number of reconstructed fibers, unlike the group analysis of Figs. 3 and 4. As for the group results, most of the abnormalities are in the periventricular zone, but a substantial number of subjects have decreased MTR and elevated MD more distally along the tract.

Asymmetry

MS is an asymmetric disorder, typically affecting both sides of the brain and spinal cord, but in different ways and to different
To test the hypothesis that the disease can disrupt the structural symmetry between the right and left sides of the brain, we calculate a total tract asymmetry index as described in Materials and methods. This index ranges from 0 (symmetry) to 1 (maximal asymmetry). Overall, median asymmetry is low, ranging from 0.02 to 0.09 in MS (Fig. 5B). Multi-way ANOVA, controlling for age, sex, and average number of reconstructed fibers across the two tracts, demonstrates significantly increased tract asymmetry in MS for $\lambda_\perp (p=0.004)$, $T2 (p=0.01)$, and MTR ($p=0.004$), but not for $\lambda_\parallel$, $FA$, $MD$, or $\lambda_\parallel$. Fig. 5C demonstrates that, except for $\lambda_\parallel$, between 20% and 50% of subjects with MS have significantly asymmetric tract indices compared to the distribution of controls.

**Discussion**

Our multiparametric analysis of the values of 6 MRI indices within the corticospinal tracts of a cohort of subjects with MS and healthy controls reveals significant abnormalities in individuals...
and, across the population, typical areas of abnormality within the tract. Our work also represents, to our knowledge, the most comprehensive, quantitative description of the range of MRI abnormalities found in a specific functional system within the brains of MS patients. We examined the CSTs because of its functional importance, the ease with which its course is reconstructed with DTI, and the ability to obtain a clinical assessment of its dysfunction. Due to the methodological issues involved in accurately characterizing clinical dysfunction and comparing that dysfunction with imaging abnormalities, we defer analysis of the relationship between MRI findings and motor weakness to future work.

**Tract profiles**

The most useful methodological contribution of this work is in the development of spatially normalized tract profiles that depict the variation in MRI indices as a function of position along a tract of interest, in this case the CST. By interpolating between anatomically well-defined boundary points within the tract – identified separately in each individual – we achieve a one-to-one correspondence among tract positions in different individual or in different scans of the same individual. This enables a quantitative description of the normal range of variation of MRI indices parametric in tract position. An individual’s deviation from this normal range and the locations of any abnormalities become readily apparent in plots such as those of Fig. 2. Because tract profiles for each index have characteristic shapes and because many of the abnormalities are focal, averaging MRI indices across the whole tract is likely to have only limited utility.

Although plots of MRI index vs. tract position have been published (Lin et al., 2006; Partridge et al., 2005; Stieltjes et al., 2001; Virta et al., 1999; Xue et al., 1999), to our knowledge no prior study covers as many indices over as long a section of the CST. Since it is unlikely that any single MRI index has a unique pathological correlate (e.g., demyelination vs. axonal disruption), the ability to examine multiple MRI indices simultaneously is an important step in acquiring a more complete description of the damage done to brain tissue by neurological disease. Accomplishing this with limited scan times requires high-field MRI scanners and sequence optimization. In this vein, it is worth noting that the scans described
increase sensitivity for detection of abnormality in disease, both within individuals and across populations.

Average vs. individual tract profiles

It is important to realize that because MS affects different individuals in different ways, the average tract profiles shown in Fig. 3 are artificial constructions. They are very useful for detecting the locations along the tract that tend to be abnormal in MS; for the CST, these locations are the subcortical white matter and corona radiata for most indices and nearly the entire brain for MTR, particularly in SPMS. Where a lesion is present in an individual, the tract profile will be much more abnormal than the average MS tract at that location, as demonstrated in Fig. 2. Thus, average profiles underestimate the degree of tract involvement by including relatively unaffected tracts and by blurring the tract profiles of individuals with lesions at different locations along the tract.

A related issue involves the distinction between MS lesions and the so-called “normal appearing white matter,” which has been demonstrated in many studies to be subtly abnormal in MS (Evangelou et al., 2000; Loewner et al., 1995; Werring et al., 1999). The tract profiles in our study do not distinguish between these different tissue types, but visual inspection suggests that most of the abnormalities that fall outside the normal range are due to lesions. It would be possible, with careful tissue segmentation, to construct separate profiles for lesional and normal appearing tissue, and such an exercise is likely to be informative.

Individual MRI indices

For most of the MRI indices discussed here, the direction of change in CST profiles of MS patients, reflected in the average profiles of Fig. 3, parallels the direction of change for that index within MS plaques and also within so-called “normal-appearing” white and gray matter. Thus, we find increased MD (Tievsky et al., 1999; Werring et al., 1999), increased T2, and decreased MTR (Dousset et al., 1992). FA does not conform to this pattern and is discussed further below.

Values of selected DTI indices (most commonly, MD and FA) along the CST have been reported in a previous study that used regions of interest at various axial positions (Toosy et al., 2003), with results in healthy controls similar to those reported here. A previous publication from our group demonstrates differences in multiple MRI indices in the hemispheric vs. brainstem portions of the CST (Reich et al., 2006). In MS, tract-specific analysis of the CST, using DTI, has demonstrated increased MD and mildly decreased FA along the tract, although indices as a function of tract position were not reported (Wilson et al., 2003).

Over a more restricted segment of the CST and set of MRI indices, tract profiles with similar shapes to those described here were found in subjects with neuromyelitis optica (NMO) and in healthy controls (Lin et al., 2006). However, we did not find that subjects with MS had decreased FA, increased MD, or \( \lambda_\perp \) within the cerebral peduncles, as was found in NMO. Whether this is due to differences between MS and NMO, or to technical differences between the two studies (e.g., 3 T vs. 1.5 T MRI scanner and 32 vs. 6 diffusion gradients), remains to be determined.

We find that MTR is the MRI index for which the overall tract profile is abnormal in the greatest number of individuals with MS (36%). MTR has long been known to be a sensitive MRI marker of
MS-related pathology (Dousset et al., 1992; Filippi and Rocca, 2004). In our results, average MTR is diminished over nearly the entire course of the intracranial CST, although the difference only achieves statistical significance in the rostral portion of the tract. In MS, MTR is sensitive to damage to both axons (van Waesbergh et al., 1999) and myelin, with perhaps a greater relative contribution from myelin (Schmierer et al., 2004). Our finding that MTR is significantly lower in subjects with progressive compared to relapsing MS (Fig. 4) is consistent with the notion that MTR is affected by factors other than demyelination – although demyelination is prominent even in PPMS (Lucchinetti et al., 2000).

Although FA within MS plaques and normal-appearing white matter has been shown to be decreased relative to control populations (Tievsky et al., 1999; Werring et al., 1999), we find, on average, no significant differences along the CST. In some cases (see Fig. 2C), FA is slightly elevated in the corona radiata and internal capsule, and this trend is also visible in the average tract profiles of Fig. 3. Within individual plaques, we find that FA is decreased as expected (not shown), although not to the same extent that MD and the directional diffusivities are increased, or that MTR is decreased.

Unchanged FA in the CST – particularly in the regions where there are significant changes in MD, T2, and MTR – seems at first surprising, but it is best explained on technical grounds. FA is a measure of the degree to which the 3D probability density function of water diffusion within a single voxel is oblong (FA closer to 1) or spherical (FA closer to 0). Even under normal circumstances, the measured FA is expected to decrease where local tract curvature is high relative to voxel size. At those sites, each voxel contains axons that are turning sharply, which leads to a more spherical shape of the diffusion probability density function and therefore a lower FA. The periventricular zone, in which MRI indices are most abnormal in MS, is close to the point of highest tract curvature, where the fibers of the CST converge to enter the internal capsule. Thus, the primary determinant of FA in this region is probably tract curvature, which dominates effects due to disease. Indeed, under some circumstances, FA might even increase in these regions in MS – if, for example, there is a decrease in local tract curvature due to brain atrophy, a prominent result of MS (Rudick et al., 1999).

Our results indicate that mean diffusivity (MD), as well as its longitudinal ($\lambda_\parallel$) and transverse ($\lambda_\perp$) components, are on average increased in MS, with the abnormality in $\lambda_\parallel$ extending over a longer segment of the tract. (By itself, a region of elevated $\lambda_\parallel$ with unchanged $\lambda_\perp$ would yield increased FA.) As with other MRI index abnormalities, increased diffusivity is most prominent in the corona radiata and subcortical white matter. Within MS plaques (see Fig. 2), diffusivity can be markedly increased, consistent with multiple previous reports (Tievsky et al., 1999; Werring et al., 1999).

### Asymmetry

Tract asymmetry indices comparing the right and left CST profiles are increased in MS relative to controls (Figs. 5B–C) but are still overall close to their minimum value of 0. Increased asymmetry reflects the fact that MS is for the most part a patchy disease, despite multiple findings of MRI abnormalities in the so-called normal appearing white matter. In practice, however, little additional information is gained by considering asymmetry in addition to the values of the MRI indices themselves. Moreover, decreasing asymmetry was not associated with chronic, progressive disease, as might perhaps be expected (data not shown).

### Limitations

Our statistical analysis included no corrections for multiple comparisons, primarily because the goal of this study was to explore the value of tract profile analysis and to determine the MRI indices that are most abnormal along the CST in MS. Needless to say, future projects that use these methods in the context of longitudinal studies or clinical trials will need to pay close attention to this issue. In addition, the control population was relatively small and had some differences in baseline characteristics – particularly, younger age – from the MS cohort. Our group analysis partially controls for age differences, which can have a substantial impact on DTI indices (Jones et al., 2006).

Another limitation of our analysis is that coregistration of DTI images with images from other sequences, including MT and T2, is imperfect in certain regions of the brain. Visual inspection indicates that for the CST, the area most strongly affected is the medulla, which probably accounts for some of the variability observed in the tract profiles in that region on the T2 scans (see Fig. 3). Distortions were similar for the DTI and MT datasets because both used echo planar readout schemes. Coregistration in the periventricular zones did not present a problem. Finally, the tractography method leads to substantial variability in the number of reconstructed fibers, which can also affect the measured MRI indices. More robust and automated fiber tracking methods would help to alleviate this problem.

### Conclusion

The magnetic resonance imaging appearance of the corticospinal tracts in multiple sclerosis is abnormal both within individuals and across the MS population. Spatially normalized tract profiles allow easy identification of the location of those abnormalities and offer the possibility of specific, quantitative assessment of individual functional systems in the brain. Future work will examine the correlations between tract-specific abnormalities and functional disability.

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