Correlation of Diffusion Tensor and Dynamic Perfusion MR Imaging Metrics in Normal-Appearing Corpus Callosum: Support for Primary Hypoperfusion in Multiple Sclerosis

BACKGROUND AND PURPOSE: Hypoperfusion of the normal-appearing white matter in multiple sclerosis (MS) may be related to ischemia or secondary to hypometabolism from wallerian degeneration (WD). This study evaluated whether correlating perfusion and diffusion tensor imaging (DTI) metrics in normal-appearing corpus callosum could provide support for an ischemic mechanism for hypoperfusion.

MATERIALS AND METHODS: Fourteen patients with relapsing-remitting MS (RRMS) and 17 control subjects underwent perfusion MR imaging and DTI. Absolute measures of cerebral blood volume (CBV), cerebral blood flow (CBF), and mean transit time (MTT) were calculated. Mean diffusivity (MD) and fractional anisotropy (FA) maps were computed from DTI data. After visual coregistration of perfusion and DTI images, regions of interest were placed in the genu, central body, and splenium of normal-appearing corpus callosum. Pearson product-moment correlation coefficients were calculated using mean DTI and perfusion measures in each region.

RESULTS: In the RRMS group, CBF and CBV were significantly correlated with MD in the splenium ($r = 0.83$ and $r = 0.63$, respectively; both $P < .001$) and in the central body ($r = 0.86$ and $r = 0.65$, respectively; both $P < .001$), but not in the genu ($r = 0.23$ and 0.25, respectively; both $P$ is nonsignificant). No significant correlations were found between MTT and DTI measures or between FA and any perfusion measure in the RRMS group. No significant correlations between diffusion and perfusion metrics were found in control subjects.

CONCLUSION: In the normal-appearing corpus callosum of patients with RRMS, decreasing perfusion is correlated with decreasing MD. These findings are more consistent with what would be expected in primary ischemia than in secondary hypoperfusion from WD.
Materials and Methods

Patients
Approval for this study was obtained from the Institutional Board of Research Associates, and informed consent was obtained from all patients. Fourteen patients with clinically definite RRMS23,23 consecutively met inclusion criteria for this study. There were 6 male patients (median age, 43.6 years; range, 33.1–49.4 years) and 8 female patients (median age, 46.9 years; range, 36.7–60.0 years). Median age for all 14 patients was 45.7 (range, 33.1 to 60.0 years). All patients were receiving immunomodulating therapy with either interferon al-a (Avonex; Biogen Idec, Cambridge, Mass) or Copolymer 1 (Copaxone; Teva, Petach Tikva, Israel). All patients were in a stable phase of disease, and no patients were receiving systemic corticosteroids during the study or in the 3 months preceding the study.

For comparison, 17 control patients were selected with no history of cerebrovascular disease, cardiovascular disease, evidence of small vessel ischemic disease, ischemic stroke, or substantial intracranial pathology on MR imaging. There were 6 male patients (median age, 49.5 years; range, 30.4–60.8 years) and 11 female patients (median age, 35.3 years; range, 23.0–65.7 years); the median age for all 17 control subjects was 39.0 (range, 23.0–65.7 years).

MR Imaging
MR imaging was performed on a 1.5T Siemens clinical scanner. Patients with RRMS and control subjects each underwent conventional MR imaging, DTI, and DSC-MR imaging within the same imaging session. Localizing sagittal T1-weighted images of the brain were obtained, followed by contiguous 3-mm axial dual echo fast spin-echo proton attenuation and T2-weighted images (TR, 3400 ms; effective TE, 17.119 ms; NEX, 1), and contiguous 5-mm thick axial FLAIR images [TR, 9000; effective TE, 110 ms; TI, 2500 ms; matrix, 512 × 256; FOV, 220 × 220 mm2]. DTI data were then obtained, followed by DSC-MR imaging and postcontrast axial (TR, 600 ms; TE, 14 ms; NEX, 1) T1-weighted images.

DTI
Axial DTI data were acquired with a pulsed gradient, double spin-echo, echo-planar imaging method (TR, 4400 ms; TE, 100 ms; matrix, 200 × 128; FOV, 220 × 220 mm2; 20-5-mm contiguous sections; b = 1000 s/mm2). The pixel size for DTI is 1.72 × 1.72 mm2. The double spin-echo technique is optimized to minimize eddy currents and geometric image distortion.24 All axial imaging was acquired oriented parallel to the anterior commissure-posterior commissure plane. Diffusion weighting (b = 1000 s/mm2) was applied along noncollinear directions, (Gx, Gy, Gz) = [(1,0,0), (1,0,1), (0,1,1), (-1,0,0), (-1,0,1), (0,1,−1)]. For each of these 6 gradient directions, 4 acquisitions were averaged. One image without diffusion weighting (b = 0) was acquired without averaging. This produced 7 images for each section.

DSC-MR Imaging
A series of 60 gradient-echo echo-planar images were acquired at 1-second intervals during the first pass of a standard dose (0.1 mmol/kg) bolus of gadopentetate dimeglumine (Magnevist; Berlex Laboratories, Wayne, NY). Seven 5-mm thick sections were acquired, positioned from the T2-weighted images to ensure coverage of a major vessel such as the anterior or middle cerebral arteries, as well as the corpus callosum. All patients had an 18- or 20-gauge intravenous catheter placed in the antecubital fossa for the purpose of contrast administration. The first 10 acquisitions were performed before contrast agent injection to establish a precontrast baseline. At the 10th acquisition, gadopentetate dimeglumine (0.1 mmol/kg) was injected with a power injector (Medrad, Indianola, Pa) at a rate of 5 mL/s, immediately followed by a bolus injection of saline (total of 20 mL at 5 mL/s). The specific imaging parameters are: TR, 1000 ms; TE, 54 ms; FOV, 230 × 230 mm2; section thickness, 5 mm; matrix, 128 × 128; in-plane voxel size, 1.8 × 1.8 mm; intersection gap, 0%–30%; flip angle, 30°; signal intensity bandwidth, 1470 Hz/pixel. The methods for acquiring perfusion data from a set of DSC enhanced echo-planar images have been described previously.25–28

Image Processing and Evaluation
Data processing was performed off-line using a Sun Ultra 10 workstation with programs developed in-house using the C and IDL (RSL, Boulder, Colo) programming languages. Six maps of the apparent diffusion coefficient were computed, from which the diffusion tensor was calculated. From this tensor, eigenvalues and eigenvectors were derived. The FA and the MD trace were computed from these measures for each voxel by standard algorithms.29 MD and FA values were calculated on a pixel-by-pixel basis to form maps for each section.

Absolute cerebral blood volume (CBV), cerebral blood flow (CBF), and mean transit time (MTT) were calculated using the method of Rempp et al.26,27 In brief, tissue concentration is found from the change in relaxation rate that occurs during bolus passage. The tissue concentration that results from an idealized, instantaneous bolus is then found by deconvolving the actual tissue concentration with the arterial input function (AIF). Deconvolution was carried out by singular value decomposition.27,28 The AIF was found using an automated method similar to that described by Rempp et al.26,30 The minimum signal intensity, corresponding to the bolus peak, is found in each pixel within the head. The average signal intensity drop and average bolus arrival time are then calculated for all pixels. Pixels where the bolus arrives early and where the signal intensity drop is larger than average are assumed to be within arteries. The AIF is found by averaging the signals from all such pixels.

After visual coregistration of the FA map, MD map, and perfusion MR images, 2 regions of interest (ROIs) were placed in the central body and in bilateral areas of genu and splenium of the corpus callosum, as shown in Fig 1. ROIs were fixed in size (radius = 1 image pixel, 1.8 mm) and placed so as to avoid arterial or venous structures on the perfusion images. Furthermore, the ROIs were placed after visual coregistration with the axial T2-weighted and FLAIR images to ensure lesions were not included in the ROI. This generated measures of FA, MD, CBV, CBF, and MTT for each of the ROIs.

Statistical Analysis
The strength of the association between DTI and perfusion measures within each area of the corpus callosum was estimated for both the RRMS group and control group using Pearson product-moment correlation coefficients. Because each subject provided multiple assessments of each metric within each area (each measure was assessed at multiple locations within each area), there were statistical dependencies (within-subject correlations) among observations that had to be accounted for in the statistical analysis. Consequently, mixed-model least-squares regression was used to assess the statistical significance of the associations between DTI and perfusion measures and to test whether the correlation between a given pair of metrics (eg, CBV with FA) in a given area was different for patients than it is for controls, while adjusting for the correlation among observations provided by the same subject and the potential confounding effects of age and sex.
The mixed model included subject age and sex as fixed numeric and classification factors, respectively, assumed that the observations derived for a given metric were correlated or independent when derived for the same subject or different subjects, respectively, and allowed the variance of the metric to differ across both subject groups and areas of the corpus callosum.

Results
The results for correlations between MD and perfusion measures are shown in Table 1. In RRMS patients, CBF was significantly correlated with MD in the central body (r = 0.86; P < .0001) and splenium (r = 0.83; P < .0001) of the corpus callosum. CBV was also significantly correlated with MD in the central body (r = 0.63; P = .0046) and splenium (r = 0.65; P < .0001) of the corpus callosum. In the genu of the corpus callosum, no significant correlations were found between MD and either CBF or CBV, and no significant correlations were present between MD and MTT in any of the areas of corpus callosum of patients with RRMS. The control group did not demonstrate significant correlations between MD and any perfusion measures in any of the areas of corpus callosum examined. The results for correlations between FA and perfusion measures are shown in Table 2. No significant correlations were present between FA and any of the perfusion metrics in any of the areas of corpus callosum of both patients with RRMS and control subjects.

The P values listed in Tables 1 and 2 were determined from the mixed-model analysis to account for within-subject correlations. These P values were used to test whether the correlation between a given pair of measures is different for patients with RRMS than it is for control subjects. Indicating that the significant correlations found in Table 1 were different between the RRMS group and the control group, significant interactions were found between MD and CBV in the central body (P = .037) and splenium (P = .018), and between MD and CBF in the central body (P = .045) and splenium (P = .017). A significant interaction was also found between FA and CBF within the central body of the corpus callosum body (P = .047). This significant interaction occurred because the correlation was positive among patients with RRMS, negative among control subjects, and moderate in magnitude within both groups (ie, although neither correlation was significant individually, the difference between them was large). No other significant interactions were found between the RRMS and control groups with respect to correlations between perfusion and DTI metrics.

Discussion
Numerous histopathologic and biochemical studies have demonstrated that in addition to the characteristic discrete white matter lesions found in patients with MS, there is diffuse abnormality involving grossly NAWM. Findings in the NAWM include decreased myelin-specific protein, diffuse astrogliosis, and infiltration by macrophages and T lymphocytes. Although MS has prototypically been characterized as a demyelinating disease, axonal damage, including axonal transection and decreased fiber attenuation, has been demonstrated in both lesions as well as within the NAWM.

Advanced MR imaging studies have demonstrated abnormalities in areas of white matter that are not visualized on conventional MR imaging. MR spectroscopy studies have found metabolic alterations within the NAWM. N-acetylaspartate is a neuronal marker that is commonly decreased within MS lesions and has also been found to be decreased in regions of NAWM, signifying diffuse axonal loss or dysfunction. Loss of macromolecular organization, such as when myelin is fragmented or destroyed, influences the transfer of magnetization from macromolecules to free water, and this magnetization can be measured using MR as the magnetization transfer ratio (MTr). In patients with MS, the MTr is significantly reduced in areas of NAWM, supporting the concept of diffuse parenchymal macrostructural abnormality.

Using DSC enhanced perfusion MR imaging, Law et al demonstrated significant hypoperfusion in the NAWM of RRMS patients compared with control subjects; however, a specific mechanism for the hypoperfusion could not be suggested on the basis of that study. Fundamentally, there are 2 possible causes for NAWM hypoperfusion, which can be categorized as primary or secondary. In the first scenario, a primary vascular pathologic lesion results in decreased perfusion in the NAWM with consequent ischemic parenchymal injury. In the second scenario, axonal damage in MS lesions leads to WD of axons traversing distant areas of white matter, resulting in decreased axonal attenuation, regional hypometabolism, and secondary hypoperfusion of the NAWM. The distinction between these mechanisms has potentially important implications, because ischemia may be an early and potentially reversible finding, whereas hypometabolism from axonal degeneration would represent advanced and irreversible disease.

DTI has proved to be a valuable tool for investigating the integrity of white matter microstructure that cannot be assessed by conventional MR imaging and has been used to study both ischemia and WD. Parameters such as the MD...
averaged over 3 orthogonal directions measure the magnitude of diffusion of water molecules, whereas diffusion anisotropy indices, such as FA, indicate the degree of deviation from isotropic diffusion of water molecules. In studies of acute ischemic stroke, MD has been found to be decreased in the immediate setting, related to acute cellular swelling. The MD later increases toward normal values and finally becomes elevated in the chronic phase, thought to represent destruction of membrane integrity and progression toward tissue necrosis. Fractional anisotropy progressively decreases in the white matter in the setting of ischemia, again attributed to ongoing tissue destruction.

Studies have demonstrated DTI changes related to WD in the setting of ischemic stroke. Thomalla et al found decreases in FA reflecting early WD in the cerebral peduncle of the affected side as early as 2–16 days after ischemic stroke, whereas maps of the orientationally averaged diffusivity did not reveal obvious changes. Months to years after ischemic stroke, FA becomes chronically decreased, and MD increases slightly along the pyramidal tract on the affected side below the primary lesion. This is thought to be related to fibrosis and atrophy of affected connected fiber tracts.

In this study, we correlated DTI and perfusion MR metrics in the normal-appearing corpus callosum of patients with MS and control subjects to further elucidate the basis for hypoperfusion in MS. We chose the corpus callosum as the focus for evaluation because it is the most highly organized interhemispheric structure in the brain, providing a sensitive area for evaluation of subtle changes in DTI measures. In the MS group, we found highly significant large-magnitude correlations between perfusion measures and MD, specifically that decreasing CBF was associated with decreasing MD (increased diffusion restriction) and not significantly correlated with FA. We interpret these findings as support for primary hypoperfusion (ischemia) in MS. In the setting of primary ischemia, areas with the lowest perfusion would be expected to have the most restricted diffusion (the most decreased mean diffusivity), with possibly only slight decreases in FA, consistent with the findings of this study. In secondary hypoperfusion related to WD, the most hypometabolic/hyperperfused areas should in theory have the lowest axonal fiber attenuation and would thus be expected to have the most increased MD and most decreased FA. The mixed model analysis demonstrated that the significant correlations between DTI and perfusion measures are not present in control subjects, suggesting that they are related to the disease process.

This study does not refute the presence or importance of WD in MS; however, it suggests that hypoperfusion is due to alternative (primary vascular) pathologic conditions. Indeed, considerable evidence supports the idea that WD does occur in MS and specifically within the corpus callosum. Recent evidence from a quantitative postmortem study of patients with MS demonstrated a significant reduction of axonal attenuation and volume in areas of corpus callosum that appeared grossly normal. Several studies of diffusion properties of the NAWM in MS have demonstrated findings of increased MD and decreased FA, as seen in WD from ischemic stroke. However, it is important to consider that the end point of tissue destruction of disparate pathologic conditions could lead to a similar pattern of DTI findings. Although studies have shown that DTI abnormalities in the NAWM correlate with DTI abnormalities within lesions, this may simply

Table 1: Pearson correlation coefficients for the association of MD with perfusion measures within each area of the corpus callosum of patients with RRMS and control subjects

<table>
<thead>
<tr>
<th>Location</th>
<th>CBV</th>
<th>P</th>
<th>CBF</th>
<th>P</th>
<th>MTT</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>RRMS (n = 14)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genus</td>
<td>0.25</td>
<td>.87</td>
<td>0.23</td>
<td>.89</td>
<td>0.08</td>
<td>.73</td>
</tr>
<tr>
<td>Body</td>
<td>0.53</td>
<td>.045</td>
<td>0.86</td>
<td>&lt;.0001</td>
<td>0.12</td>
<td>.61</td>
</tr>
<tr>
<td>Splenium</td>
<td>0.11</td>
<td>.74</td>
<td>0.04</td>
<td>.64</td>
<td>0.01</td>
<td>.72</td>
</tr>
</tbody>
</table>

Control (n = 17) | | | | | | |
| Genus | -0.09 | 0.74 | 0.01 | 0.72 | 0.31 | 0.07 |
| Body | -0.01 | 0.54 | 0.04 | 0.54 | 0.03 | 0.33 |
| Splenium | -0.03 | 0.54 | 0.004 | 0.82 | 0.004 | 0.82 |

Note: MD, mean diffusivity; RRMS, relapsing-remitting multiple sclerosis; CBF, cerebral blood flow; CBV, cerebral blood volume; MTT, mean transit time. Statistical significance accepted at P < .05.

Table 2: Pearson correlation coefficients for the association of FA with perfusion measures within each area of the corpus callosum of patients with RRMS and control subjects

<table>
<thead>
<tr>
<th>Location</th>
<th>CBV</th>
<th>P</th>
<th>CBF</th>
<th>P</th>
<th>MTT</th>
<th>P</th>
</tr>
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<tbody>
<tr>
<td>RRMS (n = 14)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genus</td>
<td>-0.38</td>
<td>.07</td>
<td>-0.24</td>
<td>.45</td>
<td>0.02</td>
<td>.99</td>
</tr>
<tr>
<td>Body</td>
<td>0.44</td>
<td>0.08</td>
<td>0.28</td>
<td>0.09</td>
<td>0.31</td>
<td>0.19</td>
</tr>
<tr>
<td>Splenium</td>
<td>-0.04</td>
<td>0.74</td>
<td>0.01</td>
<td>0.90</td>
<td>0.01</td>
<td>0.99</td>
</tr>
</tbody>
</table>

Control (n = 17) | | | | | | |
| Genus | -0.02 | 0.77 | -0.05 | 0.80 | 0.01 | 0.87 |
| Body | -0.06 | 0.67 | -0.17 | 0.26 | 0.08 | 0.59 |
| Splenium | -0.01 | 0.73 | -0.05 | 0.82 | 0.12 | 0.56 |

Note: FA, fractional anisotropy; RRMS, relapsing-remitting multiple sclerosis; CBF, cerebral blood flow; CBV, cerebral blood volume; MTT, mean transit time. Statistical significance accepted at P < .05.
reflect an underlying primary pathologic condition responsible for both lesions and NAWM damage and does not necessarily indicate that WD is primarily from lesions. In theory, diffuse ischemic injury in MS could also lead to WD of axons in the NAWM. In fact, microscopic ischemic injury in the corpus callosum, given its central location and high axonal fiber attenuation, could lead to extensive bihemispheric WD.

Further arguing against WD from lesions as the basis for NAWM hypoperfusion is that abnormalities in the NAWM have been described early in the course of disease in patients with MS without substantial lesion loads. De Stefano et al. showed that cerebral N-acetylaspartate/creatinine and MTr values are diffusely decreased in patients with MS with early disease, no significant disability, and low demyelinating lesion load, suggesting that axonal injury begins very early in the course of MS. Metabolic abnormalities have been detected in the corpus callosum using MR spectroscopy at the earliest stage of clinically isolated syndrome suggestive of MS, before atrophy and lesions are detected. It could be argued that rather than WD from distant lesions, decreased perfusion in NAWM could be related to widespread parenchymal damage below the resolution of conventional MR imaging from a nonischemic etiology. While diffuse nonischemic parenchymal injury could lead to hypoperfusion secondary to hypometabolism and similar findings of increased MD and decreased FA within the NAWM, the expected correlations between perfusion and DTI measures would be similar to that found in WD, with the most hypometabolic/hypoperfused areas containing the most parenchymal damage demonstrating highest MD and lowest FA.

Considerable histopathologic evidence supports a primary vascular pathologic lesion in MS. Studies have described perivascular inflammatory changes such as lymphocytic infiltration and edematous onion-skin changes of vein walls in NAWM lacking adjacent parenchymal inflammation, suggesting that MS could represent a form of subacute or chronic vasculitis. Vascular occlusion in MS was described on histopathologic examination by Putnam in the 1930s and later by Wakefield et al., who demonstrated fibrin deposition and thrombosis of vessels in the absence of cellular infiltration, suggesting that thrombosis of small veins and capillaries could represent an ischemic basis for disease. More recent studies have demonstrated the presence of extensive oligodendrocyte apoptosis and preferential loss of myelin-associated glycoprotein, which is suggestive of hypoxic-ischemic-type tissue injury.

It is not clear why correlations between perfusion and DTI metrics were found in the body and splenium of the corpus callosum, but not in the genu; however, this could be related to differences in fiber composition of the corpus callosum. Thin fibers seem to be most susceptible to injury in MS, and there is a higher attenuation of thin fibers in the splenium than in the genu. Although vascular changes in MS would be expected to diffusely affect the corpus callosum, it is possible that larger fibers in the genu would be more resistant to ischemic injury.

Although we interpret our findings of significant correlations between DTI and perfusion measures as significant, there are limitations to this study. Visual coregistration of the perfusion images and DTI maps can lead to errors from mis-registration. Although changes in MD would not be as subject to variability over small distances, FA is highly dependent on the area of corpus callosum sampled, and slight differences in ROI placement could partly explain a lack of correlation between perfusion measures and FA in both MS and control groups. Another limitation of the study related to visual coregistration is the inability to measure DTI and perfusion parameters in a blinded fashion, because simultaneous placement of corresponding ROIs was required. However, measurements from the perfusion analysis were not visible when placing ROIs on the DTI images and were obtained simultaneously for all ROIs so as to limit bias from manipulation of ROI placement.

The perfusion algorithm used has inherent limitations and is accurate only if there is negligible delay and dispersion in the bolus between the arteries where the AIF is measured and the tissue of interest. Delays and dispersion introduce errors into the calculation of perfusion parameters; however, these errors should be minimal, because the AIF is estimated close to the site of the perfusion measurements. The effect of the disease process itself on the AIF is also not known, though at this time it seems that there is relative sparing of the major arteries in the vasculitic process and hence the AIF may not be greatly affected in MS. Finally, all patients in the MS group were undergoing chronic immunomodulating therapy, possibly decreasing the inflammatory component of the disease process within the NAWM and affecting both DTI and perfusion measures.

The results of this study correlating DTI and perfusion changes are more consistent with what would be expected in primary ischemia than in secondary hypoperfusion from WD. Further investigation, including larger and longitudinal studies, are warranted to additionally support the concept of ischemic injury in MS. A better understanding of the role of ischemia could aid in predicting clinical course, monitoring response to therapy, and designing novel targets for therapeutic intervention for MS.

Conclusions

In areas of normal-appearing corpus callosum of patients with RRMS, decreasing perfusion is correlated with decreasing MD and is not significantly correlated with FA. These findings support the concept of primary ischemia in MS rather than secondary hypoperfusion as a result of WD.

References


