Changes in the normal appearing brain tissue and cognitive impairment in multiple sclerosis

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Abstract

Objectives—To assess (a) whether the changes in the normal appearing brain tissue (NABT), as revealed by magnetisation transfer (MT) histogram analysis, correlates with cognitive dysfunction in patients with multiple sclerosis and (b) the relative contribution of these changes by comparison with that of multiple sclerosis lesions visible on conventional MRI.

Methods—Dual echo, T1 weighted and MT scans of the brain were obtained in 12 patients with multiple sclerosis with cognitive impairment and in seven without cognitive impairment. Lesion loads were assessed from T2 and T1 weighted scans. To create MT histograms of the NABT, multiple sclerosis lesion outlines from dual echo scans were superimposed automatically and nulled out from the co-registered and scalp stripped MTR maps. Average lesion MT ratio (MTR) and brain size were also measured.

Results—T2 and T1 lesion loads were significantly higher and the average lesion MTR and brain size were significantly lower in the group of cognitively impaired patients. Patients with cognitive deficits also had significantly lower average MTR and peak location of the NABT histogram. Logistic regression analysis showed that 68% of the total variance was explained by average NABT-MTR alone. A multivariable regression model showed that NABT-MTR was the only factor that significantly correlated with cognitive impairment in these patients (p=0.001).

Conclusions—The extent of abnormalities which go undetected when using conventional MRI is relevant in determining cognitive impairment in multiple sclerosis.

Keywords: multiple sclerosis; magnetic resonance imaging; magnetisation transfer imaging

Several studies have shown that in patients with multiple sclerosis, the extent of white matter abnormalities visible on conventional brain MRI, their regional distribution, and their intrinsic nature, as assessed on T1 weighted and magnetisation transfer imaging (MTI), all correlate well with cognitive impairment. In general, the degree of such correlations is higher than that found with physical disability.

Lesion load estimates from conventional MRI are known not to give a complete picture of the burden of disease. Pathological studies showed that abnormalities can be detected in the brain tissue outside macroscopic multiple sclerosis lesions. These abnormalities include diffuse astrocytic hyperplasia, patchy oedema, perivascular cellular infiltration, and abnormally thin myelin and axonal damage. Such pathological abnormalities modify the relative proportions of mobile and immobile protons of the diseased tissue and, therefore, it is not surprising that MTI is able to show microscopic damage outside multiple sclerosis lesions visible on conventional T2 weighted scans.

More recently, two studies showed that the overall disease burden (macroscopic lesions and subtle abnormalities in the normal appearing brain tissue (NABT)), as assessed by MTI histograms of the entire brain, is correlated with the presence and severity of cognitive impairment in patients with multiple sclerosis. As in these studies the whole brain tissue was used to create MTI histograms, the relative contributions of macroscopic lesions and of subtle NABT abnormalities to the cognitive decline in multiple sclerosis were not assessed. In the present study, we created MTI histograms of the NABT to answer the following questions: (a) Is the damage of the NABT, as shown by the MTI histogram analysis, relevant in determining cognitive dysfunction in patients with multiple sclerosis? and (b) If so, which is the relative contribution of these changes in comparison to that of macroscopic sclerosis lesions visible on conventional MRI?

Patients and methods

PATIENTS

We studied 19 patients (eight men and 11 women). Of these, 12 patients had cognitive impairment and seven had no evidence of cognitive impairment (see below). Patients were similar in age, level of education, disease duration, and neurological disability, as assessed by the expanded disability status scale (EDSS) (table 1). According to the Lublin and Reingold criteria, 11 patients (four without and...
Table 1: Main demographic and clinical data from patients with multiple sclerosis with and without cognitive impairment

<table>
<thead>
<tr>
<th></th>
<th>Unimpaired</th>
<th>Impaired</th>
<th>pValue*</th>
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<tbody>
<tr>
<td>Mean age (SD)</td>
<td>37.7 (8.6)</td>
<td>37.6 (7.0)</td>
<td>NS</td>
</tr>
<tr>
<td>Mean duration of the education (SD) (y)</td>
<td>11.6 (3.8)</td>
<td>11.1 (4.2)</td>
<td>NS</td>
</tr>
<tr>
<td>Mean disease duration (range) (y)</td>
<td>15.0 (10-20)</td>
<td>11.2 (4-18)</td>
<td>NS</td>
</tr>
<tr>
<td>Mean EDSS (range)</td>
<td>4.5 (3.0-7.0)</td>
<td>4.5 (2.0-6.5)</td>
<td>NS</td>
</tr>
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</table>

For statistical analysis, see text.

NEUROPSYCHOLOGICAL TESTS

Neuropsychological tests exploring the executive functions (Weigl test, Dual task test, Wisconsin card sorting test (WCST), Stroop Colour/word interference test (Stroop test), Hanoi Towers test, verbal fluency test) and verbal and spatial memory (digit span test, short tale test, and Corsi span and supraspan test) were always obtained within 48 hours from the acquisition of the MRI. These tests are routinely used in the assessment of cognitive impairment in multiple sclerosis and are extensively described elsewhere. Patients’ results on the Weigl test, verbal fluency test, digit span test, short tale test, and Corsi span and supraspan test were considered abnormal according to previously published criteria based on normative values obtained from a large group of Italian speaking subjects. For the other tests, patient performances were considered abnormal when they were equal to or above the 90th percentile (Stroop test, Hanoi towers test) and equal to or below the 5th percentile (WCST, dual task test) of the results obtained from a group of 20 sex and age matched healthy controls. Patients who had normal results in tests exploring the two cognitive domains (executive and memory functions) were classified as cognitively unimpaired, the others (patients with deficits of the executive functions, memory, or both) were classified as cognitively impaired. For each patient, the results from all neuropsychological tests were then scored, using a standardised method based on a comparison with the percentile distribution of values from normal controls. These scores ranged from 0 to 4, where grade 0 means a very poor performance and grade 4 means a normal performance. Individual test scores were then summed to provide a composite cognitive score for each patient.

IMAGE ACQUISITION

On a 1.5 Tesla scanner, each patient underwent dual echo turbo spin echo (TSE) (TR=3300, TE=16/98, echo train length (ETL)=6) scans. Twenty four axial 5 mm thick contiguous interleaved slices were obtained, with rectangular 188x250 mm field of view and 194x256 image matrix, thus obtaining approximately a 1x1 mm inplane resolution. MTI and T1 weighted magnetisation prepared rapid acquisition gradient echo (MP-RAGE) scans were also obtained in the same session. For MTI, 2D gradient echo images (TR=600, TE=12, flip angle=20°) with and without a saturation pulse were obtained. The saturation pulse was an off resonance gaussian radiofrequency pulse centred 1.5 kHz below the water frequency, with a duration of 16.4 ms, a bandwidth of 250 Hz, and a power intensity of 3.4x10⁻³ Tesla. The same acquisition parameters as for the TSE scans were used for MTI scans, except for the number of slices, which was 20. The set of slices for MTI was positioned to obtain the same central 20 slices as for the other acquisition schemes. From the two sets of images— that is, without (Mo) and with (Ms) saturation pulse and after their coregistration (see next paragraph)—quantitative MT ratio (MTR) images were derived pixel by pixel according to the following equation: MTR=(Mo−Ms) / Mo×100, where Mo is the mean signal intensity for a given pixel without the saturation pulse and Ms is the mean signal intensity for the same pixel when the saturation pulse is applied. Signal intensities in the calculated images represented the MTR value. For MP-RAGE scans (TR=10, TE=4, TI=700, flip angle=10°, number of acquisitions=1), a 3D sagittal slab (194x256x160 image matrix, 250 mm field of view) covering the entire brain was acquired. This sequence gives the advantage that the acquired 3D images can be reconstructed in any plane and with different slice thicknesses and it has been shown that it enables a similar load of hypointense lesion to be detected as conventional T1 weighted scans do. The original data were then reformatted to obtain 24 axial contiguous 5 mm thick slices, with the same orientation and offsets as the corresponding TSE slices. Patients were positioned in the scanner using published guidelines for multiple sclerosis studies.

IMAGE ANALYSIS

For the MRI data analysis, multiple sclerosis lesions were first outlined on proton density (PD) and MP-RAGE hard copies by the consensus of two experienced observers, who were unaware of the patients’ clinical characteristics and cognitive test results. T2 weighted scans were always used to increase confidence in lesion detection. For T1 weighted MP-RAGE scans, a conservative approach was used. Only areas with a signal intensity between those of the CSF and grey matter and with corresponding abnormalities on PD and T2 weighted images were considered as hypointense lesions. Lesion volume measurements were then performed by a trained technician, who was also unaware of patients’ clinical characteristics and cognitive test results.
Table 2 T1 and T2 lesion loads, average lesion MTR and brain volumes from patients with multiple sclerosis with and without cognitive impairment

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<tbody>
<tr>
<td>Median T2 lesion load (range) (ml)</td>
<td>8.4 (2.4–18.4)</td>
<td>38.4 (4.7–73.6)</td>
<td>0.002</td>
</tr>
<tr>
<td>Median T1 lesion load (range) (ml)</td>
<td>1.4 (0.1–6.2)</td>
<td>6.7 (1.2–47.2)</td>
<td>0.01</td>
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<tr>
<td>T1/T2 ratio</td>
<td>0.12 (0.03–0.34)</td>
<td>0.38 (0.06–0.71)</td>
<td>0.03</td>
</tr>
<tr>
<td>Average lesion MTR (SD) (%)</td>
<td>35.1 (2.0)</td>
<td>32.1 (3.0)</td>
<td>0.006</td>
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For statistical analysis, see text.

Table 3 MT histogram derived measures of the normal appearing brain tissue (NABT) from patients with multiple sclerosis with and without cognitive impairment

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<th>Impaired</th>
<th>p Value*</th>
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<tbody>
<tr>
<td>Average MTR (SD) (%)</td>
<td>41.6 (0.5)</td>
<td>39.8 (0.9)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Peak height</td>
<td>103.0 (9.1)</td>
<td>92.5 (12.7)</td>
<td>0.07</td>
</tr>
<tr>
<td>Peak location</td>
<td>37.1 (0.9)</td>
<td>34.7 (1.2)</td>
<td>&lt;0.0001</td>
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*Student’s t test for unpaired data.
with the T2 (r=−0.78, p<0.0001) and the T1 (r=−0.60, p=0.006) lesion loads. The percentages of the total variance explained by each individual factor included in the model were as follows: average NABT-MTR=68%, T1 lesion load=38%, T2 lesion load=37%, average lesion MTR=35%, and brain size=21%. The multivariable regression model showed that average NABT-MTR (p<0.001) was the only factor that was significantly correlated with cognitive impairment in our patients. The other tested variables (T1 and T2 lesion loads, average lesion MTR, NABT histogram peak height and location, and brain size) were all removed from the model. Significant correlations were also found between the composite cognitive score and average lesion MTR (r=0.72, p<0.001), average NABT-MTR (r=0.75, p<0.0001), peak height of the NABT-MTR histogram (r=0.65, p=0.004), and peak location of the NABT-MTR histogram (r=0.77, p<0.0001).

**Discussion**

Conventional MRI is very sensitive in detecting multiple sclerosis lesions and their changes over time.17 For this reason MRI is widely used to monitor multiple sclerosis evolution, both natural or modified by treatment.37 38 However, it is known that conventional MR techniques, among other limitations, are not able to detect the full extent of changes seen pathologically in multiple sclerosis.3 They include subtle abnormalities in the NABT.3 14 Although multiple sclerosis related changes in the NABT have to be subtle, otherwise they would result in visible areas of focal abnormalities on conventional scanning, they may be so extensive as to have some influence on the evolution of the disease. The following points support the concept that the extent of the microscopic changes in the NABT may contribute to the development of disability in multiple sclerosis. Firstly, the MTR of the white matter away from multiple sclerosis lesions and the MTR in the white matter surrounding such lesions from patients with secondary progressive multiple sclerosis are lower than those of the corresponding regions from patients with less disabling relapsing-remitting multiple sclerosis.19 30 Secondly, white matter outside macroscopic multiple sclerosis lesions from patients with secondary progressive multiple sclerosis has an N-acetyl aspartate to creatine ratio on average 8.2% lower than the that from patients with relapsing-remitting multiple sclerosis.38 In patients with relapsing-remitting multiple sclerosis a progressive reduction of this ratio is detectable over time and its decrease correlates strongly with changes in physical disability.40 Although changes in the brain tissue which appears normal on conventional scanning are more likely to have an impact on cognitive functioning of multiple sclerosis patients rather than on physical disability, no previous study has assessed the contribution of such changes to the impairment of cognition.

The present study confirms that the extent of the lesions seen on conventional MRI and the severity of tissue disorganisation within these lesions are relevant in determining cognitive impairment in patients with multiple sclerosis.1 4 This study also clearly shows that NABT changes do make a significant contribution to multiple sclerosis clinical manifestations. Clearly, MTI is not the only MR technique that detects such subtle changes in the normal appearing white matter of the patients we studied. Therefore, it is likely that other quantitative MR techniques, such as the analysis of T1 and T2 relaxation times, diffusion measurements and imaging, and MR spectroscopy, might also provide useful information. That widespread NABT damage is relevant for cognitive decline in multiple sclerosis is in agreement with the results of previous PET studies,41 42 suggesting that cognitive decline in multiple sclerosis may be the result of widespread damage to deep brain structures, leading to functional disconnection of different cortical areas. This finding also agrees with the relatively low correlations found by previous studies between regional MR abnormalities and deficits in specific cognitive domains.4 The recent demonstration that significant axonal loss occurs in the white matter outside macroscopic multiple sclerosis lesions provides further evidence that changes which go undetected on conventional scanning may be functionally relevant.15 16

Because normal appearing white matter represents a large part of the tissue included in NABT MT histograms, we think that microscopic white matter abnormalities were the major contributing factor to the observed differences between the MT histograms of the two groups of patients. Nevertheless, it is likely that lesions in or adjacent to the cerebral cortex, which can be imaged using fast fluid attenuation inversion recovery sequences,43 44 may have been missed in the present study and may also have contributed to the MT histogram changes seen for NABT and, as a consequence, to cognitive impairment.5 The role of pathological changes in the basal ganglia is likely to be minor, if present at all, due to the low frequency of clinical or MRI involvement of these structures in multiple sclerosis.45–47 We can only speculate about the nature of the subtle changes potentially responsible for the cognitive impairment seen in our patients. Small focal abnormalities independent of larger lesions visible on conventional scans or damage to axons transversing large focal lesions and
resulting in wallerian degeneration in areas away from such abnormalities may both contribute to cognitive impairment. The results of the present study allow us to propose that these abnormalities are not mutually exclusive. The larger amount of macroscopic lesions seen in cognitively impaired patients and the correlations found between NABT changes and T1 and T2 lesion loads suggest the role of secondary axonal degeneration. However, the larger contribution of NABT-MTR than lesion extent and severity to the presence of cognitive impairment, as shown by the multivariable analysis, suggests that subtle focal abnormalities other than secondary axonal degeneration are also relevant.

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