Changes within the “normal” cerebral white matter of multiple sclerosis patients during acute attacks and during high-dose cortisone therapy assessed by means of quantitative MRI

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SUMMARY Changes in the apparently unaffected cerebral white matter of multiple sclerosis (MS) patients were studied during acute attacks as well as during high-dose prednisolone therapy. Serial MR scans of patients with a clinically definite diagnosis were performed on four defined occasions: before an episode, within three days after its onset, after 10 days of therapy as well as four weeks later. Thirteen patients agreed to cooperate in forming a MRI data base and to be rescanned immediately after the onset of an acute relapse. Within one year, six patients had such episodes, one of them had a second bout. Both T1 and T2 relaxation times within the apparently normal white matter were significantly prolonged in all cerebral lobes compared to a control group of healthy volunteers. During the acute attacks as well as during therapy the T1 values remained as before. The T2 values were elevated only in two out of six cases during the episode. After therapy a considerable clinical improvement was seen in all cases, but a significant T2 decrease as a possible effect of cortisone was noted in only one case. We conclude that the prolonged relaxation times T1 and T2 within the apparently normal cerebral white matter of MS patients are the result of a number of molecular events differing considerably among individual patients and that serial measurements of these relaxation times do not consistently change during an acute relapse and do not reflect clinical improvement after high dose prednisolone therapy.

According to a number of studies1-3 the measurement of the relaxation times T1 and T2 within the visually normal cerebral white matter in cases of multiple sclerosis (MS) has offered some interesting possibilities for the clinical use of quantitative magnetic resonance imaging (MRI). It was shown that the apparently normal cerebral white matter has longer T1 and T2 relaxation times than normal controls. These findings demonstrate in vivo what has been concluded from earlier neuropathological reports4 that MS is not a disease restricted to focal areas of demyelination but rather involving the entire cerebral white matter.

The clinical significance and the biophysical nature of this finding have mostly remained unclear to date. In another study5 the normal cerebral white matter of two groups of MS patients, differing only in the duration of their disease, were investigated by means of quantitative MRI. Only patients with a long-standing course of the disease (>5 years) showed a T2 prolongation in the apparently normal periventricular white matter compared with the group with a shorter disease duration (<5 years) or with another group of healthy controls. The conclusion from this finding was that the prolongation of T2 develops gradually and is more pronounced in the later stages. Although the specificity of T1 and T2 measurements within the apparently normal cerebral white matter has not yet been established, there is reasonable hope for quantitative MRI to be used as a sensitive paraclinical method for documenting the progression of the disease. In another study6 quantitative MRI has shown some potential for assessing treatment and in helping to understand the mechanism of action of therapeutic agents in multiple sclerosis patients.
The purpose of this study was to ascertain whether such changes in relaxation times of the apparently normal cerebral white matter occur during an acute relapse and whether these measurements are subject to changes during high-dose cortisone therapy.

Patients and methods

Thirteen patients with a clinically definite diagnosis of MS (Poser Criteria) and a recent inception of the illness (mean: 2-7 years, range: 0-5-3-5 years) as well as with mild physical impairment (Expanded Disability Status Scale (Kurtzke): Grade 0–2) agreed to cooperate in forming a data base and to be rescaned immediately after the onset of a bout. All patients had a remitting and relapsing course of the disease and had shown at least one bout with a cerebral localisation. In all patients the cerebrospinal fluid examination had revealed oligoclonal bands as well as elevated intrathecal IgG production. In addition, the disseminated character of the disease was demonstrated by means of conventional MRI showing high-signal lesions on the spinecho sequences that were regarded as characteristic. In no case were the MR-scans normal. No patient had had a bout in the preceding three months and only patients without long-time medication (cortisone and azathioprin) were accepted. All patients were regularly seen in the MS outpatient service of the Vienna Neurological University Clinic. All episodes were diagnosed by unequivocal clinical signs. Within one year six patients had had episodes, one of them a second bout. The clinical and MRI data of these patients are summarised in table 1.

The first examination was performed when the patient entered the study. Another scan was made within three days after the onset of the episode but before therapy was started. The third examination was performed on the day following a nine-day high-dose cortison therapy (1 g of prednisolone over five days, 500 mg to the seventh day, and 250 mg to the ninth day) and the final fourth examination was performed four weeks after the onset of the episode. The control group consisted of 18 healthy normal volunteers (nine males and nine females; mean age: 37 years, range: 23–49 years).

In all serial examinations care was taken to achieve exact repositioning with reference to the bicommissural plane. All examinations were performed on a 0.5 Tesla unit (Gyroscan) with a standard heat coil, using spin-echo sequences (TR 2000 ms, TE 50/100 ms) with contiguous slices, a slice factor of 1:1, and a slice thickness of 8 mm. Routine scans were displayed on a 256 × 256 matrix.

In addition, calculated images were obtained in two different planes, one directed through the bicommissural plane, the second immediately above the corpus callosum parallel to the bicommissural plane through the upper frontal and upper parietal lobe. In the chosen planes IR and SE-sequences were obtained as well as a mixed mode sequence with the chosen parameters: SE repetition time 710 ms, IR repetition time 2290 ms, inversion delay 310 ms, echo time 50 ms, slice thickness 8-0 mm, number of measurements two, scan resolution 128. The techniques by which the calculations were achieved have been described previously. Methods of minimising error and increasing reproducibility of relaxation time measurements have been considered. To assure the long-term reproducibility of our measurements, measurements with CuSO4 standard probes were performed on repeated occasions showing an overall accuracy for T1 and T2 within a range of 10%. The calculated images were stored on magnetic tape for off-line measurements.

Measurements were performed with a standard cursor device (0-2 square centimetre). The region of interest (ROI) was always chosen on the IR scan, which is thought to be most sensitive for showing plaques. Care was taken to clearly place the ROI outside a plaque region within the normally appearing white matter. The ROI was then kept constant for the measurements. T1 and T2 measurements were taken from the upper parietal lobe, the upper frontal lobe, the temporal-parietal region, the temporal lobe, and the frontobasal lobes. For each patient or control a total of 20 measurements were taken. Statistical evaluation was performed by means of analysis of variance.

Results

A total of six patients with seven episodes were evaluated. In one patient the quantitative follow up data could not be evaluated because of technical reasons.

Quantitatively, the base examinations all showed elevated T1 and T2 values of the apparently normal

<table>
<thead>
<tr>
<th>Case</th>
<th>Sex</th>
<th>Age/years</th>
<th>Clinical findings during episodes</th>
<th>Cranial MRI-lesion visualised during bout</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>65</td>
<td>Spinal paraparesis, numbness D5 downwards, cerebellar tremor</td>
<td>Unchanged lesion left internal capsule, periventricular lesion adjacent to left lateral ventricle</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>29</td>
<td>Sensorimotor paraes right arm</td>
<td>Periventricular lesions on lateral and fourth ventricles increase in size</td>
</tr>
<tr>
<td>3</td>
<td>M</td>
<td>38</td>
<td>Bilateral internuclear ophthalmoplegia, cerebellar ataxia</td>
<td>Large lesion in upper frontal and upper parietal white matter bilaterally</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>18</td>
<td>Sensory paroes left hand and reduction of finger tapping</td>
<td>Paramedian left midbrain lesion. Increase in size of bilateral lesions in the upper frontal white matter</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>34</td>
<td>Bilateral internuclear ophthalmoplegia, increase of spastic atactic signs on lower extremities</td>
<td>Unchanged bilateral periventricular lesions adjacent to lateral ventricles</td>
</tr>
<tr>
<td>6a</td>
<td>F</td>
<td>30</td>
<td>Spastic-atatic gait disturbance</td>
<td>Unchanged</td>
</tr>
<tr>
<td>6b</td>
<td>F</td>
<td>30</td>
<td>Increase of spastic atactic gait disturbance. Peripheral facial palsy</td>
<td>Unchanged</td>
</tr>
</tbody>
</table>
Changes within the "normal" cerebral white matter of multiple sclerosis patients

Changes within the "normal" cerebral white matter of multiple sclerosis patients (table 2). During the series of clinical and therapeutic events (base examination, relapse, treatment phase, follow up) the patient group did not show significant changes for either T1 or T2 values. This was true for all brain regions measured (upper frontal, upper parietal, temporoparietal, parietal, or frontobasal).

All patients showed marked clinical improvement after therapy. No consistent changes were found for T1 values and only individual patients showed T2 changes compared with their values obtained at the base examination (fig la and b). In two out of six patients there was a significant T2 elevation during the episode. One of them showed a return to the base value after therapy with a second increase at the follow up examination. One patient showed an elevated T2 throughout the serial study following the base examination which did not change during therapy or the four weeks after. In the other four patients the onset of an episode did not significantly affect the T2 value compared to the base value. In two cases no T2 changes were measured throughout the serial study, in one further patient there was a significant T2 prolongation after therapy and in another patient the only significant change measured was a T2 decrease following a second episode at the time of the second follow up examination.

Discussion

This study shows that T1 and T2 measurements of the visually normal cerebral white matter of MS patients

Table 2 Measurements of relaxation times T1 and T2 within the visually normal cerebral white matter of MS patients as compared with normal healthy controls show a significant prolongation in all measured regions (ANOVA ranges: * = p < 0.05; † = p < 0.01; § = p < 0.001).

<table>
<thead>
<tr>
<th></th>
<th>T1- values</th>
<th>T2- values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Normals (n = 18)</td>
<td>Patients (n = 13)</td>
</tr>
<tr>
<td>Upper parietal</td>
<td>454 (20)</td>
<td>490 (14)†</td>
</tr>
<tr>
<td>Upper frontal</td>
<td>426 (10)</td>
<td>484 (39)§</td>
</tr>
<tr>
<td>Temporoparietal</td>
<td>444 (16)</td>
<td>487 (21)§</td>
</tr>
<tr>
<td>Temporal</td>
<td>433 (15)</td>
<td>511 (38)§</td>
</tr>
<tr>
<td>Frontobasal</td>
<td>414 (20)</td>
<td>464 (47)§</td>
</tr>
</tbody>
</table>

Figs 1a and 1b Changes of T1- and T2- values within the visually unaffected cerebral white matter of six patients with multiple sclerosis. A total of seven acute relapses were documented. BD = base data (before the onset of an episode), B = within three days after beginning of the bout, before therapy, AT = after treatment (on the day following a nine-day high-dose cortisone therapy), FU = follow up (four weeks after the beginning of the bout). ANOVA ranges see figures.
do not change in a uniform way during a clinical relapse (fig 2). In addition, such measurements do not predictably change in response to high dose cortisone therapy in our cases. Although it could not be demonstrated in all patients, in individual cases, such measurements reflect changes of the macromolecular environment. In one patient (case 1) an increase of T2 in the normal white matter could possibly indicate an increase of intercellular water, which was shown to be reversible by cortisone. In the further course a second

Figs 2a–d   Serial MR-examination (SE 100/2000) of corresponding cuts through the upper frontal and parietal lobes of case 2. Further quantitative evaluation of T1 and T2 values within the visually unaffected white matter of the brain did not show a significant change between the scans at the time before or after the acute relapse although all values were elevated in comparison to normal controls. a: base examination. b: on the day following the onset of an acute relapse. c: on the day following a nine-day high-dose cortisone therapy. d: four weeks later.
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increase of T2 was shown four weeks after therapy which could reflect the beginning of astrogial proliferation within the entire white matter of the brain. But in other patients such measurements were totally unpredictable or did not change at all. Also, the failure to find changes of T1 during a clinical relapse weighs against the assumption that an increase in the concentration of extracellular water is the sole biochemical factor contributing to the change of T2. Arguably the degree of elevation of T1 is positively correlated with the total amount of water in a lesion, whereas the degree of T2 elevation also depends on the total amount of water as well as on the macromolecular environment of the water. Changes on the macromolecular level within the visually normal white matter has not yet been fully investigated. One quantitative neuropathological study has shown, however, that the main fragments of demyelination (myelin-associated glycoprotein, basic myeloprotein, and proteolipoprotein) cannot be traced within the normally appearing white matter of MS brains, whereas within the plaque- and periplaque area they can be found in abnormally high amounts. It therefore cannot be assumed that demyelination contribute to the T2 values within the normal white matter.

In one other study multiple sclerosis patients have been investigated using quantitative MRI immediately before as well as three to seven days after a five day treatment with intravenous methylprednisolone. Similarly in our study no significant change of T2 values was found within the apparently normal cerebral white matter in the pre- and post treatment measurements, but a significant decrease of T1 was noted within the "normal" white matter which the authors ascribed to the antioedematous effect of cortisone. Although the patients in that study are comparable to our group of patients (in terms of age, disability status and applied cortisone dose), we failed to find such a significant change of T1 values in our group. We are inclined to explain this difference through the activity status of the disease. We included only patients with an acute relapse and only these patients received cortisone therapy in our series, whereas Kesselring et al applied cortisone only to clinically stable patients. We therefore conclude that, in acute relapses of multiple sclerosis, additional macromolecular changes play a role in the intercellular compartment of the apparently unaffected white matter of the brain thus influencing the T1 values. Such changes may seem to be effective in the relapsing phase and not in clinically stable patients, where the effect of high-dose cortisone is mainly reflected in a decrease of T1 within the "normal" white matter. This might explain why our T1 measurements did not significantly change in response to high-dose cortisone therapy in spite of a clear-cut remission achieved in all our patients.

Measurements of T1 and T2 within the plaque areas have not been evaluated in this study. Other reports show that neither age nor activity of the plaque can be reliably assessed, although a prolongation of T1 and T2 is frequently seen within an active plaque. However, the ranges of the values obtained show considerable overlap, so that no conclusions can be drawn from single measurements within plaque areas. Therefore, serial measurements of plaque areas with and without application of Ga-DTPA can be considered the most reliable biological marker reflecting the activity of the disease.

References