Axonal Damage in Multiple Sclerosis Patients with High versus Low Expanded Disability Status Scale Score

Steven D. Brass, Sridar Narayanan, Jack P. Antel, Yves Lapierre, Louis Collins, Douglas L. Arnold

ABSTRACT: **Background:** The pathophysiological basis for differences in disability in patients with multiple sclerosis is unclear. **Methods:** We used magnetic resonance imaging to examine whether differences in disability in cohorts of multiple sclerosis patients with similar T2-weighted lesion volume and disease duration were associated with a more destructive disease process in the more disabled patients. **Results:** The benign and severely disabled groups had similar brain atrophy metrics and similar decreases of the neuronal marker, N-acetylaspartate, in the normal appearing white matter (NAWM), and chronic T1-weighted hypointense lesions (black holes) and atrophy. **Conclusion:** The dissociation of spinal cord atrophy and cerebral atrophy between these two groups suggests that the difference between the more benign and more disabled groups cannot be explained by a more aggressive pathological process that is affecting the entire neuroaxis in a homogeneous fashion.

RÉSUMÉ: Comparaison du dommage axonal chez les patients atteints de sclérose en plaques ayant un score élevé ou un score bas à l’échelle étendue du statut d’invalidité. **Introduction:** La physiopathologie des différences dans le degré d’invalidité chez les patients atteints de sclérose en plaques est mal connue. **Méthodes:** Nous avons utilisé l’imagerie par résonance magnétique pour évaluer si les différences observées dans le degré d’invalidité chez des cohortes de patients atteints de sclérose en plaques dont le volume des lésions à l’examen pondéré en T2 et la durée de maladie étaient similaires, étaient associées à un processus morbide plus destructeur chez les patients les plus invalides. **Résultats:** Que l’invalidité soit légère ou grave, la mesure de l’atrophie cérébrale et la diminution du marqueur neuronal N-acétyl aspartate dans la matière blanche du cerveau, à la spectroscopie par résonance magnétique in vivo, étaient similaires dans les deux groupes. La cohorte ayant le degré d’invalidité le plus élevé avait plus d’atrophie au niveau de la moelle épiplère. **Conclusion:** La dissociation entre l’atrophie de la moelle épiplère et l’atrophie cérébrale chez ces deux groupes de patients suggère que la différence entre le groupe ayant une invalidité légère et celui ayant une invalidité plus marquée ne peut s’expliquer par un processus pathologique plus agressif touchant de façon homogène tout l’axe neural.

Methods

Study population

Data from 25 patients with clinically definite MS were reviewed: 13 had benign MS (EDSS ≤ 3.0, disease duration ≥10 years) and 12 had disabling MS (EDSS = 5 - 8.5 inclusively, disease duration ≥ 10 years). We obtained the benign and disabled groups by retrospectively looking back at the Montreal Neurological Institute MS-magnetic resonance spectroscopy (MRS) database of patients who had previously undergone combined MRI/MRS exams. We then identified the disabled group by selecting patients who had T2W lesion loads and disease duration that were similar to those of the benign group, but a higher EDSS score (5 - 8.5). The group of patients with disabling MS was composed of two patients with relapsing-remitting disease and 10 patients with secondary-progressive MS. A group of 15 normal volunteers served as control subjects. Radiological data were examined to determine T2W lesion volume, NAA:creatine ratio (NAA/Cr), spinal cord cross-sectional area, and demographics in all three cohorts (Table). Informed consent was obtained from all subjects.

Proton MRI and MRS of brain

Combined MRI and MRS examinations of the brain were obtained in a single session for each examination using a scanner operating at 1.5 T (Phillips Gyroscan). A transverse dual echo, turbo spin-echo sequence [repetition time (TR), 2075 milliseconds; echo times 32 and 90 milliseconds, 256 x 256 matrix, 1 signal average, 250-mm field of view] yielding proton density-weighted and T2WI images with 50 contiguous slices was acquired parallel to the line connecting the anterior and posterior commissures. The MRIs were used to select an intracranial volume of interest (VOI) for spectroscopy measuring approximately 90 mm anteroposteriorly by 20 mm cranio-caudally by 90 mm left to right. This VOI was centred on the corpus callosum to include mostly white matter and some mesial cortex of both hemispheres. Two-dimensional spectroscopic images were obtained using a PRESS sequence (TR, 2000 milliseconds; TE 272 milliseconds; 250-mm field of view; 32 x 32 phase encoding steps; 1 signal average per step) as previously described.

Measurement of spinal cord atrophy

High resolution T1-weighted imaging of the spinal cord was used to accurately assess spinal cord cross-sectional area at the level of C2 as previously described by Narayanan et al.

Measurements of the brain to intracranial capacity ratio

The brain to intracranial capacity ratio (BICCR) is calculated from the volume of automatically segmented cerebrospinal fluid, grey matter, white matter, and lesions, as previously described by Collins et al. The BICCR is the ratio obtained by dividing the sum of the volumes of (grey matter + white matter + lesion) by the sum of the volumes of (grey matter + white matter + lesion + cerebrospinal fluid).

Data analysis

N-acetylaspartate values were expressed relative to intravoxel Cr to compensate for machine-dependent variations of signal intensity over the VOI. The NAA/Cr ratios for all the voxels in the spectroscopic VOI (excluding edge voxels containing spectra that were artfactually distorted) were then averaged to obtain a summary NAA/Cr ratio for each subject. Central brain NAA/Cr values and spinal cord cross sectional areas were compared across groups using analysis of variance followed by pairwise post hoc comparison using the Tukey honestly significant difference procedure to account for multiple comparisons. Data were considered significant at the 0.05 level.

Table: Demographics of the Cohort.

<table>
<thead>
<tr>
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<th>Controls (n=15)</th>
<th>Benign MS (n=13)</th>
<th>Disabled MS (n=12)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years) Mean (SD)</td>
<td>39 (8)</td>
<td>39 (5)</td>
<td>45 (9)</td>
</tr>
<tr>
<td>EDSS Median (Range)</td>
<td>-</td>
<td>1.5 (0-2.5)</td>
<td>7 (5-8.5)</td>
</tr>
<tr>
<td>Disease Duration (years) Mean (SD)</td>
<td>-</td>
<td>15 (5)</td>
<td>17 (5)</td>
</tr>
<tr>
<td>Cerebral T2W LV (cc) Mean (SD)</td>
<td>-</td>
<td>5.4 (3.5)</td>
<td>8.7 (4.8)</td>
</tr>
<tr>
<td>% VOI Occupied by Lesion Mean (SD)</td>
<td>-</td>
<td>2.4 (1.9)</td>
<td>3.2 (1.8)</td>
</tr>
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SD = standard deviation; EDSS = Expanded Disability Status Scale; VOI = volume of interest

was widespread and uniform throughout the neuraxis. We therefore predicted that the more disabled cohort would demonstrate evidence of more spinal cord injury as determined by the extent of magnetic resonance imaging (MRI)-defined spinal cord atrophy. To assess this, we measured the density of the neuronal marker compound, NAA in deep central white matter of the brain, as well as global brain atrophy. If differences in the destructiveness of MS pathology are responsible for differences in disability and the pathology is homogeneous within an individual, then patients with greater disability would be expected to have lower density of NAA and greater cerebral atrophy for a given cerebral lesion volume.

Since the benign group had relatively low cerebral lesion volumes, matching the groups for cerebral lesion volume had the effect of selecting for low cerebral lesion volume in both the benign and disabled groups. This meant that > 95% of the spectroscopic volume of interest (VOI) used for spectroscopy was composed of NAWM. Technical limitations do not allow us to derive information about the NAA density in lesions that comprise such a small proportion of brain volume. Therefore, our measurements of NAA in deep central brain essentially reflect the situation in NAWM. Since NAA density in lesions and NAWM decreases nonlinearly over the course of MS, we matched patients for disease duration, as well.
RESULTS

A retrospective review was carried out on 11/13 benign patients and 11/12 disabled patients with respect to clinical presentation and course of illness. Among the benign group, only 1/11 had relapses involving motor dysfunction compatible with myelopathy whereas 10/11 had sensory related relapses. In the disabled group, 8/11 had relapses involving motor dysfunction compatible with spinal cord involvement whereas 3/11 had recurrent events involving brainstem rather than clearcut motor dysfunction in the limbs. Cerebral NAA/Cr ratios, BICCR values and the spinal cord cross-sectional areas for the three cohorts are summarised in the Figure. The mean cerebral NAA/Cr ratio in a volume of interest centred on the corpus callosum containing mostly normal appearing white matter was not different in the patients with benign MS compared to disabling MS (p=0.44). Despite having low T2W lesion volume, both the benign and disabled group had low brain NAA/Cr compared to the normal control group (p<0.003). The BICCR values did not differ among the benign, disabled and control groups (p>0.16). The disabled group had significantly smaller mean spinal cord cross-sectional area at the level of C2 than the benign group (p=0.006) and controls (p<0.001). Cord cross-sectional area did not differ between the benign and the control groups (p>0.45).

DISCUSSION

Our study compared cohorts of patients with benign and disabling multiple sclerosis with similar cerebral lesion loads and disease durations in order to determine the factor responsible for differences in their disability. The cytodestructive process in the brains of the disabled group of patients was not found to be any more severe than that of the benign group, based on measures of cerebral NAA/Cr and BICCR, neither of which showed statistically significant differences between the benign and disabled groups. The low cerebral NAA/Cr in both our benign and disabled groups, compared to control subjects, indicates that significant axonal pathology was present in both. Given that T2W lesions occupied only 2.4% and 3.2% of the VOI in the benign and disabled groups, respectively, the observed decreases of central brain NAA/Cr in both patient groups must reflect primarily reduction of NAA/Cr in normal-appearing white matter.8

The difference in disability between the benign and disabled patients in this cohort appear to be due to the development of spinal cord pathology rather than differences in the degree of cerebral tissue injury. A retrospective review of the clinical course of patients demonstrated that patients in the disabled group acquired disability from progressive disease rather than from one single destructive attack in the spinal cord. Four patients of the high disability cohort had early relapses with motor involvement compatible with spinal cord involvement contributing to the progressive disability. Among the low disability group, only one patient presented with myelopathy whereas others had mainly sensory symptomatology. When comparing both groups, patients who suffered from attacks involving spinal cord-related motor dysfunction were more likely to go on to secondary progressive disease and become disabled than those patients who had purely sensory findings likely pointing to a more benign course of disease. This is consistent with previous prognostic guides in multiple sclerosis which demonstrated that early onset sensory symptoms versus a course involving residual motor signs is one of the factors associated with a more benign prognosis.10 The severely disabled cohort had more spinal cord atrophy, suggesting that more spinal cord injury had occurred in the more disabled cohort. This reinforces the importance of relapses and lesion location13,14 as causes of disability as opposed to widespread, homogeneous destructive pathological processes that are more severe throughout the central nervous system of patients with greater disability. The amount of tissue destruction was dissociated between the brain and spinal cord in our patient groups. Further imaging and histological studies will be required to establish whether this dissociation is a result of the quantity of spinal cord lesions or heterogeneity of the disease process at different sites of the neuroaxis.

REFERENCES

1. Thompson AJ, Kermode AG, MacManus DG, et al. Patterns of


