Reduced magnetisation transfer ratio in cognitively impaired patients at the very early stage of multiple sclerosis: a prospective, multicenter, cross-sectional study

J H Faiss,1 D Dähne,1 K Baum,2 R Deppe,3 F Hoffmann,3 W Köhler,4 A Kunkel,1 A Lux,5 M Matzke,5 I K Penner,6 M Sailer,5 U K Zettl7

ABSTRACT

Objectives: Cognitive impairment belongs to the core symptoms in multiple sclerosis (MS) and can already be present at the very early stages of the disease. The present study evaluated cognitive functioning after the first clinical presentation suggestive of MS and brain tissue damage in a non-lesion focused MRI approach by using magnetisation transfer imaging (MTI).

Setting and participants: 47 patients (15 men and 32 women; mean age: 31.17 years) after the first clinical event suggestive of MS were recruited in six different MS centres in Germany and underwent a neuropsychological test battery including tests for attention, memory and executive function as well as depression and fatigue. MTI and conventional MRI measures (T1/T2 lesion load) were assessed. In addition, Magnetisation Transfer Ratio (MTR) maps were calculated. Primary outcome measure was the investigation of cognitive dysfunction in very early MS in correlation to MRI data.

Results: 55.3% of patients with MS failed at least one test parameter. Specifically, 6% were reduced in working memory, 14.9% in focused attention, 25.5% in figural learning and up to 14.9% in executive function. When the sample was subdivided into cognitively impaired and preserved, MTR scores within the cognitively impaired subgroup were significantly lower compared with the preserved group (t(43) =2.346, p=0.02*). No significant differences between the two groups were found in T2-weighted and T1-weighted lesion volume.

Conclusions: After the first MS-related clinical event, 55.3% of patients showed distinct cognitive deficits. Cognitively impaired patients had significantly lower whole brain MTR, but no differences in focal brain lesion volumes supporting the idea that early cognitive deficits may be related to diffuse loss of brain tissue integrity.

INTRODUCTION

The prevalence of cognitive deficits in multiple sclerosis (MS) ranges between 43% and 65%, depending on various research settings and characteristics of study samples.1–2 It is assumed that the MS-induced inflammatory demyelination and axonal damage may contribute to the cognitive decline. Other contributing factors are advanced age, low IQ or educational level and depression or fatigue.3–8 Cognitive dysfunction has a profound effect on maintaining employment, daily-living activities, social life, ability to drive and benefits from inpatient rehabilitation.1–3–8

Whereas intelligence, language, semantic memory and attention span are widely preserved, complex attention, information

For numbered affiliations see end of article.

CrossMark

Downloaded from http://bmjopen.bmj.com/ on May 10, 2022 by guest. Protected by copyright.
processing speed, verbal and visuospatial memory and executive functions are frequently involved. More specifically with regard to memory functions, explicit and episodic memory as part of long-term memory as well as short-term “working memory” are commonly affected.1

Reports about cognitive dysfunction already present at the stage of the first clinical event suggestive of MS are growing.9–18 Previous work shows that patients with MS in the early stage of the disease performed significantly lower on neuropsychological assessment compared to healthy controls.3 4 9–12 19 20 Often studies used only short neuropsychological test batteries, which were not sensitive enough to detect impairment in the study population of clinically isolated syndrome (CIS) or patients with early MS.19 20 Furthermore, methodological problems like a diffuse definition of neuropsychological impairment, lack of consideration of depression and fatigue as covariates or lack of healthy controls to detect practice effect reduce the informative value of such studies.

Clinical trials on neuroimaging in patients at the point of first clinical event in correlation to cognitive dysfunction are rare.19 20 In other MS subtypes, cross-sectional studies with conventional MRI demonstrated conflicting results between cognitive dysfunction and cerebral lesion load in T2-weighted and T1-weighted sequences including corpus callosum lesions,21 22 juxtacortical lesions,23 and lesions in the prefrontal cortex.24 More advanced MRI techniques such as Magnetisation Transfer Transfer Ratio (MTR) allow detection of brain tissue involvement correlating with myelin and axonal density.25 26 MTR is a sensitive parameter to quantify the integrity of myelinated white matter in patients with MS including demyelination and remyelination even in the absence of axonal loss. This technique has been used in the past to assess the global burden of occult diseases using histograms or overall mean MTR values. The decrease of the mean MTR of cortical and subcortical regions was previously related to poorer performance on cognitive tests19 27 and impaired sustained attention and concentration28 29 in relapsing-remitting MS. The aim of our study was to corroborate the previous reports of this underestimated clinical finding in early disease stage and to substantiate the cognitive deficits by using MRI in a conventional versus a more advanced non-lesion focused way that potentially better reflects the underlying clinicopathological mechanism especially very early in the disease progress.

METHODS
Participants
Forty-seven patients (15 men and 32 women; mean age: 31.17 years) after the first clinical event suggestive of MS were recruited in six different MS centres for the study (Halle: N=11, Hennigsdorf: N=6, Magdeburg: N=5, Rostock: N=10, Teupitz: N=7 and Wernsdorf: N=8). Patients were included into the study directly after the diagnosis of MS or CIS was established and after checking the inclusion and exclusion criteria. Major exclusion criteria were current alcohol or substance abuse, history of head injury or any other medical condition affecting cognition. Furthermore participants were excluded if they had severe motor or visual impairments that might interfere with cognitive testing. All patients were treatment-naive. Disease modifying drugs prior to study inclusion were defined as exclusion criteria.

MRI
MRI was conducted for all patients at the MS centre, Magdeburg. MRI was performed using a neuro-optimised 1.5-T GE Signa Horizon LX scanner with the standard quadrature head coil (General Electric, Milwaukee, Wisconsin, USA). Protocol for MTI consisted of a PD-weighted SE sequence (TR 2600, TE 20, 256×256) with and without a preparing saturation pulse (1200 Hz off-resonance, 1180 flip-angle, 16 ms). Forty-eight slices of 3 mm thickness aligned along the AC-PC line were acquired. Image postprocessing included calculation of MTR maps and an intersequence correction of movement with the automated image registration package rigid body model (AIR).30

T1-weighted sequences (3 mm slices, FOV 250, TR 700 ms) were performed without and with 0.1 mmol/kg bodyweight Gd-DTPA (3 mm slices, TR 3000 ms, TE100 ms). Additionally, T2-weighted images were assessed (3 mm slices, TR 3000 ms, TE100 ms). Lesion volume quantification was performed on T2-weighted and on T1-weighted images using a semiautomated, local contour technique. T1 lesions were identified as hypointense areas on the T1-weighted scans and confirmed on a T2-weighted scan as lesions. The semiquantitative lesion load measurement was performed using a highly reproducible31 32 threshold technique based on the local environmental intensity of the lesion (Displimage software package supplied by Dave Plummer, University College London, UK).

By application of an additional off-resonance HF impulse during a proton density-weighted imaging sequence the magnetisation of the immobile protons is partially saturated. Driven by relaxation processes, magnetisation is transferred from the mobile proton pool to the immobile one. The resulting signal attenuation in the mobile pool gives a signal reduction by imaging the pulse sequence with MT pulse. The signal reduction depends on tissue properties (especially the content of different macromolecules) and image sequence parameters. In two independent imaging sequences with (MT) and without (noMT) a saturation pulse the magnetisation transfer can be expressed as an MTR calculated voxel by voxel: MTR=(noMT−MT)/noMT in percent. Calculations were performed with MATLAB (MathWorks). Lesion and cortex were included to obtain an overall measure for brain parenchymal integrity.

Neuropsychological assessment
Neuropsychological assessment was performed 90–180 days after the first clinical event because at the point of
neuropsychological testing all patients had to be relapse-free and without corticosteroid treatment. The interval between neuropsychological assessment and MRI was no longer than 14 days. Neuropsychological tests were performed by neuropsychologists. The used test procedure was selected by means of high objectivity (independence of rater). Before study start, neuropsychologists were instructed to use the same test order and instructions to patients.

The neuropsychological battery (table 1) was administered in two test parts with a 15 min break in between. The sequential order of tests within each test part was kept constant. The battery comprised attention tasks that are part of a computerised Attention Test Battery (TAP33) and the Symbol Digit Modalities Test (SDMT34), the Verbal Learning and Memory Test (VLMT35), the Visual Learning and Memory Test for Neuropsychological Assessment (DCS-a Wolfram version36) and measures of executive functions, such as the Wisconsin Card Sorting Test (WCST, Nelson version37) and the Regensburger Word Fluency Test (RWT, animals, S-words, G/R change38). In addition, a recognition vocabulary test was used to assess the premorbid intelligence (german vocabulary scale, WST39). Finally, participants completed self-reported instruments for depression (German Depression Scale, ADS-L40), fatigue (the Modified Fatigue Impact Scale, MFIS41) and quality of life (Functional Assessment of Multiple Sclerosis, FAMS12). The MFIS consists of 21 statements regarding fatigue, cut-off was a total value above 22 points. Depressive symptoms were assessed using the ADS-L. Cut-off was a total value above 23 points. Further, all participants were examined by means of the Multiple Sclerosis Functional Composite (MSFC43) and the Expanded Disability Status Scale (EDSS). All patients gave written informed consent prior to inclusion in the study. The present study was conducted in accordance with the declaration of Helsinki and has been approved by the appropriate ethics committee of Brandenburg. Details that might disclose the identity of the participants were omitted.

### Data Evaluation and Analysis

The presented study was an exploratory pilot study with no sample size calculation. Individual performance on each of the neuropsychological tests was evaluated against standardised data. Test scores 2 SDs below the normative sample (ie, PR<2.3, T<30) were considered to reflect impaired performance. Statistical analyses were calculated with the Statistical Package for the Social Sciences (SPSS Inc, Chicago, Illinois, USA) and by an independent biometrical institute. Patients with and without cognitive dysfunction were compared using t tests or Mann-Whitney U tests for independent samples. Pearson’s χ² tests were used to compare the observed frequencies of gender, educational level and handedness between the two groups (cross-over-tables). Correlations between cognitive parameters and clinical parameters were calculated by means of non-parametric rank correlations (Spearman) or product–moment correlations (Pearson). The non-parametric Mann-Whitney test was used for comparisons of lesion volumes of the cognitively impaired and unimpaired patients with MS. Mean MTR values were compared using t tests for independent samples. A significance α level of 0.05 for minimum was predetermined.

### Results

#### Population

Table 2 summarises demographic and clinical characteristics of all participants and of the two subgroups (cognitively impaired/cognitively preserved). Mean disease duration of the overall group was 1.56 months (±1.25) at the point of study inclusion. At the time of neuropsychological testing and MR investigation (90–180 days after study inclusion), 77% of the patients were classified as CIS and 23% as RRMS (relapsing remitting multiple sclerosis) according to McDonalds criteria.45 Regarding demographic and clinical parameters between the two subgroups no significant differences were observed (table 2).

#### Cognitive Functioning

By means of cognitive testing our sample could be divided into the following two subgroups: 55.3% (N=26) of patients who failed at least one test (≥2 SD of the normative data) and 44.7% (N=21) of patients who were cognitively unimpaired. Twenty-eight per cent of the whole sample failed in one test parameter, 17% failed in two test parameters and 11% failed in at least three test parameters. Results of the neuropsychological testing are presented in table 3.

#### Attention

Compared with the normative sample, 14.9% of patients failed for divided attention and 6.4% for information processing speed and working memory measured by the Paced Auditory Serial Addition Test (PASAT as part of MSFC).
Memory
Verbal learning and memory measures were preserved in nearly all patients. In contrast, 25.5% of the patients showed disturbed figural memory performance.

Executive functions
Patients of 8.5–14.9% showed deficits in verbal fluency but only 4.3% of patients made significantly more perseverative errors in the WCST compared to the normative sample.

Intellectual ability
A recognition vocabulary test (WST) was used to estimate premorbid intellectual abilities to consider the possible positive effect of ‘cognitive reserve’ on test performance. In our study population premorbid intelligence was averaged and between the subgroups (cognitively impaired and preserved patients) no significant differences were obtained (WST: z-norm: M=0.03±0.59).

Depressive mood
Depression scores obtained from the ADS-L revealed that 11% of our patients showed clinically relevant depression. In the cognitively impaired subgroup 4.3% were depressed but cognitive impairment and depression scores were not significantly associated (t(45)=1.51, p=0.14).

Fatigue
Approximately 36% of patients showed evidence of clinically relevant fatigue. Twenty-three per cent of the cognitively impaired subgroup and 14.9% of the cognitively preserved subgroup were fatigued. t Test analysis showed no significant differences regarding the occurrence of fatigue between the two subgroups (t(45)=1.01, p=0.32).

Disability measurement
Median EDSS score in the overall group was 1.5 (range 0–4). Cognitively impaired patients did not show significant differences concerning EDSS (t(43)=−0.125, p=0.90) and MSFC scores (t(45)=0.14, p=0.89) when compared to unimpaired patients.

Magnetisation transfer imaging and cognition
Whole brain MTR was significantly lower in the cognitively impaired subgroup compared to the preserved patients (t(43)=2.346, p=0.02*). Regarding T1-weighted and T2-weighted lesion load no significant differences between the two subgroups were detectable. No significant correlations were found between MTR and EDSS (r=−0.09, p=0.55).

DISCUSSION
The results indicate that after the first MS-related clinical event, up to 55.3% of patients showed distinct cognitive deficits in the domains of attention, figural learning and executive functions. According to the classification by Amato et al.4 who suggested an allocation due to severity of cognitive dysfunction: 28% of the patients were cognitively mildly disabled, 17% were moderately disabled and 11% were severely disabled. These results correspond to the prevalence rates usually obtained in more advanced MS stages.4 46 47

Table 2 Demographic, clinical and MRI characteristics

|                    | All patients | Patients with cognitive deficits | Patients without cognitive deficits |
|--------------------|--------------|----------------------------------|-----------------------------------|
| Patient number     | 47           | 26 (55.3%)                       | 21 (44.7%)                        |
| Gender m/f         | 15/32        | 10/16                            | 5/16                              |
| Mean age in years  | 31.17±8.89   | 30.54±9.20                       | 31.95±8.64                        |
| Education in years (number) | <10 years: 3  | <10 years: 0                      | <10 years: 3                      |
|                    | 10 years: 25 | 10 years: 15                      | 10 years: 10                      |
|                    | >10 years: 16 | >10 years: 9                      | >10 years: 7                      |
| Handedness right/left | 40/3        | 22/2                             | 18/1                              |
| Verbal-IQ (WST: z score) | 0.03±0.59   | 0.01±0.68                        | 0.07±0.46                         |
| Mean disease duration (month) | 1.56±1.25   | 1.41±0.85                        | 1.74±1.59                         |
| Disease course†    | CIS: 76.6% (N=36), RRMS: 23% (N=11) | CIS: 76.9% (N=20), RRMS: 23.1% (N=6) | CIS: 76.2% (N=16), RRMS: 23.8% (N=5) |
| Median EDSS (min/max) | 1.5 (0/4.0)  | 1.5 (0/3.5)                      | 1.5 (0/4.0)                       |
| MSFC (M/SD)        | 0.56±0.33    | 0.55±0.32                        | 0.57±0.34                         |
| T1-weighted lesion load‡ (mean/median) | 1332.07/492.00 | 1380.00/558.00          | 1275.00/474.00                    |
| T2-weighted lesion load (mean/median)‡ | 2793.65/1798.50 | 2949.84/1734.00 | 2607.71/1863.00                  |
| Mean MTR (M/SD)iii | 47.69±0.97   | 47.39±0.91                       | 48.03±0.94                        |

Significant differences between patient groups are marked and p values are displayed below: †t(44)=−0.186, p=0.854 ns; ‡t(44)=−0.365, p=0.717 ns; ‡t(43)=2.346, p=0.02, MTR data based on 45 patients, 1 patient did not complete the MRI investigation. †Disease course at the time of neuropsychological testing and MR investigation. ‡The T1-weighted lesion load is related to the volume of T1 hypointense lesions. CIS, clinically isolated syndrome; EDSS, Expanded Disability Status Scale; MTR, magnetisation transfer ratio; MSFC, Multiple Sclerosis Functional Composite.
The proportion of observed cognitive disorders in MS is very inconsistent in the literature due to a lack of standards in definition of cognitive impairment and in neuropsychological tests used. In this study a very conservative criterion for cognitive impairment (at least 2 SDs below the appropriate norm) was applied. The selected neuropsychological tests are well established tools providing high methodological standards for retest-reliability and validity. The SDMT and PASAT, one of the most widely used brief repeatable battery of neuropsychological tests (BRB-N 48), were used to investigate sustained and complex attention as well as information-processing speed and working memory. Although in this study cohort performance on the SDMT did not account for the subdivision into cognitively impaired and preserved patients, the recently represented Brief International Assessment of Cognition for MS (BICAMS), which takes 15 min to complete and comprises the SDMT, a learning test (The California Verbal Learning Test) and a visuospatial memory test (The Brief Visuospatial Memory Test-Revised) can be recommend for use in daily clinical practice unless extensive cognitive analysis is not possible.49 Although, no significant correlations between depression or fatigue and cognitive impairment were observed, both were considered as parameters which can interfere with cognitive performance. The lack of correlations between cognition and fatigue or depression in contrast to results in other studies 50 51 could be explained by the short disease duration and disease course of our study population. In the literature, the relationship between psychological status and cognition is still under discussion.52–54

| Cognitive tests | All patients M (SD)* | 2SD (%) | Patients with cognitive deficits M (SD) | Patients without cognitive deficits M (SD) |
|-----------------|----------------------|---------|---------------------------------------|-------------------------------------------|
| **Attention**   |                      |         |                                       |                                           |
| Tonic alertness | 252.87 (±44.83)      | 4.3     | 256.27 (±47.63)                       | 248.67 (±41.88)                           |
| Phasic alertness| 241.49 (±41.54)      | 2.1     | 247.50 (±48.69)                       | 234.05 (±30.00)                           |
| Divided attention|                     |         |                                       |                                           |
| Speed           | 689.39 (±78.01)      | 14.9    | 712.64 (±84.66)                       | 660.60 (±58.85)                           |
| Mistakes        | 1.04 (±1.33)         | 2.1     | 1.23 (±1.37)                          | 0.81 (±1.29)                              |
| Omissions       | 1.26 (±1.63)         | 6.4     | 1.73 (±1.99)                          | 0.67 (±0.73)                              |
| Cognitive flexibility | 756.62 (±147.23)  | 2.1     | 770.29 (±148.18)                      | 739.69 (±147.86)                          |
| SDMT            | 59.96 (±10.07)       | 2.1     | 56.96 (±10.10)                        | 63.67 (±8.92)                             |
| PASAT (MSFC)    | 47.87 (±9.86)        | 6.4     | 46.85 (±10.38)                        | 49.14 (±9.26)                             |
| **Memory**      |                      |         |                                       |                                           |
| DCS             |                      |         |                                       |                                           |
| Immediate recall| 26.23 (±11.29)       | 6.4     | 28.65 (±11.30)                        | 23.24 (±10.81)                            |
| Learning score  | 22.86 (±9.58)        | 25.5    | 19.12 (±10.20)                        | 27.50 (±6.34)                             |
| **Executive function** |            |         |                                       |                                           |
| WCST-Nelson†    | 14.95 (±20.19)       | 4.3     | 17.27 (±20.95)                        | 12.07 (±19.32)                            |
| RWT             |                      |         |                                       |                                           |
| Animals         | 37.64 (±11.04)       | 14.9    | 34.15 (±11.17)                        | 41.95 (±9.45)                             |
| S-words         | 21.11 (±6.73)        | 8.5     | 19.58 (±7.13)                         | 23.00 (±5.81)                             |
| G/R-change      | 19.81 (±5.90)        | 10.6    | 17.38 (±5.58)                         | 22.81 (±4.92)                             |
| **Depression**  |                      |         |                                       |                                           |
| ADS-L           | 11.23 (±7.72)        | 10.6‡   | 9.73 (±7.20)                          | 13.10 (±8.10)                             |
| **Fatigue**     |                      |         |                                       |                                           |
| MFIS            | 19.72 (±15.62)       | 36.2§   | 17.65 (±14.00)                        | 22.29 (±17.43)                            |
| **Premorbid IQ**|                      |         |                                       |                                           |
| WST             | 29.87 (±4.29)        | 0       | 29.46 (±4.98)                         | 30.38 (±3.28)                             |

*Mean and SD. †per cent perseverative errors. ‡per cent clinically relevant depression. §per cent clinically relevant fatigue. ADS-L, German version of the Center for Epidemiological Studies Depression Scale; DCS, Visual Learning and Memory Test for Neuropsychological Assessment; MFIS, Modified Fatigue Impact Scale; MSFC, Multiple Sclerosis Functional Composite; PASAT, Paced Auditory Serial Addition Test; TAP, Attention Test Battery; RWT, Regensburger Word Fluency Test; VLMT, Verbal Learning and Memory Tests; SDMT, Symbol Digit Modalities Test; WST, German vocabulary scale.
The extent of physical disabilities as measured by EDSS and MSFC was not correlated with cognition, which is often described in the literature.\textsuperscript{30–34} This can be explained by the fact that EDSS is not sensitive in determining cognitive disorders and physical and cognitive impairment occur independently from each other during the disease course. Our study population was only mildly disabled measured by EDSS and had a short disease duration. Cognition might be a sensitive marker especially at the onset of the disease. In addition no significant associations were found between MTR and EDSS. However, cognitively impaired patients had significantly lower MTR scores. MTR decrease may be an early sign for tissue changes related to impaired cognitive function while physical disability may be absent. Recently, reductions of cortical MTR have also been found to correlate with cognitive status in physically mildly disabled patients in the later stages of the disease.\textsuperscript{35–36} In contrast lesion load parameters, such as T2 lesion load, showed no association with overall cognitive decline in the patients, although white matter lesion formation is a hallmark in the pathology of MS. The dissociation of focal inflammatory lesions in the white matter from the clinical manifestations of relapses, disability and cognitive performance has been previously reported.\textsuperscript{21–25} In a recent study Deloire et al\textsuperscript{37} found that diffuse brain damage measured by normal-appearing brain tissue MTR at the onset of the disease was the main predictor of cognitive changes over 7 years. Consequently a less lesion-focused view of MS especially in the workup of cognitive deficits is necessary. Since cognitive performance is heavily dependent on the integrity of neuronal network a lesion independent assessment of tissue integrity may at best reflect cognitive deficits. This study supports the hypothesis that cognitive decline in early MS is rather associated with impaired integrity of brain tissue as a result of generalised myelin destruction and axonal loss than with focal white matter pathology. Our results also indicate that non-conventional MRI techniques, for example, MTR, with increased specificity to more destructive aspects of MS pathology are able to reflect cognitive disturbances even at earliest stages of MS (mean disease duration 1.56 month). The evolution of pathological changes in brain tissue as depicted by a lower MTR in the overall cognitively impaired group may even start prior to the first clinical event.\textsuperscript{51} On the basis of our findings that show significant correlations between cognitive performance and decreased MTRs suggesting diffuse brain tissue changes in patients with early MS,\textsuperscript{62–64} the cognitive status should be included in treatment decisions, independent of physical disability as a marker for disease severity and progression. Neuropsychological tests may reveal clinically significant cognitive disability from the first presentation suggestive of MS.

**References**

1. Amato MP, Zippoli V, Portaccio E. Cognitive changes in multiple sclerosis. *Expert Rev Neurother* 2008;8:1585–96.
2. Rao SM, Leo GJ, Bernardin L, et al. Cognitive dysfunction in multiple sclerosis. I. Frequency, patterns, and prediction. *Neurology* 1991;41:685–91.
3. Patti F. Cognitive impairment in multiple sclerosis. *Mult Scler* 2009;15:2–B.
4. Amato MP, Ponziani G, Pracucci G, et al. Cognitive impairment in early-onset multiple sclerosis. Pattern, predictors, and impact on everyday life in a 4-year follow-up. *Arch Neurol* 1995;52:168–72.
5. Beaty WW, Paul RH, Wilbanks SL, et al. Identifying multiple sclerosis patients with mild or global cognitive impairment using the Screening Examination for Cognitive Impairment (SEFCI). *Neurology* 1995;45:718–23.
6. Langdon DW, Thompson AJ. Multiple sclerosis: a preliminary study of selected variables affecting rehabilitation outcome. *Mult Scler* 1999;5:94–100.
7. Rao SM, Leo GJ, Ellington L, et al. Cognitive dysfunction in multiple sclerosis. II. Impact on employment, social functioning. *Neurology* 1991;41:692–6.
8. Schultheis MT, Garay E, Deluca J. The influence of cognitive impairment on driving performance in multiple sclerosis. *Neurology* 2001;56:1089–94.

9. Achiron A, Barak Y. Cognitive impairment in probable multiple sclerosis. *J Neurol Neurosurg Psychiatry* 2003;74:443–6.

10. Schulz D, Kopp B, Kunkel A, et al. Cognition in the early stage of multiple sclerosis. *J Neurol* 2006;253:1002–10.

11. Simioni S, Ruffieux C, Bruggmann L, et al. Cognition, mood and fatigue in patients in the early stage of multiple sclerosis. *Swiss Med Wkly* 2007;137:496–501.

12. Zavidovin R, Sepic J, Nasuelli D, et al. A longitudinal study of brain atrophy and cognitive disturbances in the early phase of relapsing–remitting multiple sclerosis. *J Neurol Neurosurg Psychiatry* 2001;70:773–80.

13. Achiron A, Barak Y. Cognitive changes in early MS: a call for a common framework. *J Neurosci* 2006;26:47–51.

14. Audoin B, Duong MVA, Ranjeva J-P, et al. Magnetic resonance study of the influence of tissue damage and cortical reorganization on PASAT performance at the earliest stage of multiple sclerosis. *Hum Brain Mapp* 2005;24:216–28.

15. Feuillet L, Reuter F, Audoin B, et al. Early cognitive impairment in patients with clinically isolated syndrome suggestive of multiple sclerosis. *Mult Scler* 2007;13:1004–10.

16. Fredrikson S, Wickklein E, Scherer P, et al. Cognitive performance in early multiple sclerosis (MS): baseline data for CogniMS, a multinational longitudinal study. *Neurology* 2008;70(Suppl 1):PO4.174.

17. Glanz B, Holland C, Gauthier S, et al. Cognitive dysfunction in patients with clinically isolated syndrome (CIS): a newly diagnosed multiple sclerosis. *Mult Scler* 2007;13:1004–10.

18. Potagas C, Giogkaraki E, Koutsis G, et al. Cognitive impairment in different MS subtypes and clinically isolated syndromes. *J Neurol Sci* 2008;267:100–6.

19. Deloire MS, Assen C, Hamel D, et al. MRI predictors of cognitive outcome in early multiple sclerosis. *Neurology* 2011;76:1161–7.

20. Khalil M, Enzinger C, Langkammer C, et al. Cognitive impairment in relation to MRI metrics in patients with clinically isolated syndrome. *Mult Scler* 2011;17:173–80.

21. Camp SJ, Stevenson VL, Thompson AJ, et al. Cognitive function in primary progressive, transitional progressive multiple sclerosis: a controlled study with MRI correlates. *Brain* 1999;122:1341–8.

22. Rovaris M, Comi G, Filippi M. MRI markers of destructive pathology in early multiple sclerosis. *Mult Scler* 2006;24:111–16.

23. Lazeron RH, Langdon DW, Filippi M, et al. Neuropsychological impairment in multiple sclerosis patients: the role of (juxta)cortical lesions on FLAIR. *Mult Scler* 2000;6:280–9.

24. Foong J, Quaghebeur G, et al. Executive function in multiple sclerosis. The role of frontal lobe pathology. *Brain* 1997;120:15–26.

25. van Waesberghe JH, Kamphorst W, De Groot C, et al. Fatigue in multiple sclerosis: relationship to working memory capacity. *Ann Neurol* 2004;56:407–15.

26. Rovaris M, Filippi M, Minicucci L. Cortical/subcortical disease burden and cognitive impairment in patients with multiple sclerosis. *AJNR Am J Neuroradiol* 2000;21:402–8.

27. Deloire MS, Salort E, Bonnet M, et al. Cortical/subcortical disease burden associated with fatigue in patients with multiple sclerosis. *Mult Scler* 2005;11:1221–3.

28. Summers MM, Fiskniku LK, Anderson VM, et al. Cognitive impairment in relapsing-remitting multiple sclerosis can be predicted by imaging performed several years earlier. *Mult Scler* 2006;12:197–204.

29. Woods RP, Mazzotti JC, cherry SR, et al. MRI-PET registration algorithm. *J Comput Assist Tomogr* 1997;21:356–40.

30. Plummer D. DispImage: a display and analysis tool for medical images. *Riv Neuroradiol* 1992;5:485–98.

31. Boringa JB, Lazeron RH, Reuling IE, et al. Validation of the functional assessment of multiple sclerosis quality of life instrument. *Mult Scler* 2007;13:1004–10.

32. Anderson AK, Splid PE, Andersen H, et al. Fatigue and processing speed are related in multiple sclerosis. *Eur Neurol* 2010;70:1218–21.

33. Arnett PA, Higginson CI, Voss WD, et al. Correlation of self-assessed fatigue and alertness in multiple sclerosis. *Mult Scler* 2010;16:313–20.

34. Arnett PA, Higginson CI, Voss WD, et al. Depression in multiple sclerosis: relationship to working memory capacity. *Neuropsychology* 1999;13:546–56.

35. Arnett PA, Higginson CI, Voss WD, et al. Depression mood in multiple sclerosis: relationship to capacity-demanding memory and attentional functioning. *Neuropsychology* 1999;13:434–46.

36. Foong J, Rozewicz L, Quaghebeur G, et al. Neuropsychological deficits in multiple sclerosis after acute relapse. *J Neurol Neurosurg Psychiatry* 1998;64:529–32.

37. Reuter F, Audoin B, Filippi M, et al. Pulsed methylprednisolone induces a reversible impairment of memory in patients with relapsing-remitting multiple sclerosis. *Acta Neurol Scand* 1998;97:366–9.

38. Flachenecker P, Meissner H. Fatigue in multiple sclerosis presenting as acute relapse: subjective and objective assessment. *Mult Scler* 2008;14:274–7.

39. Bellmann-Strobl J, Wuerfel J, Aktas O, et al. Poor PASAT performance correlates with MRI contrast enhancement in multiple sclerosis. *Neurology* 2009;73:1624–7.

40. Filippi M, Rocca MA, Benedict RHB, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

41. Feinstein A, Kitsos P, Miller DH, et al. Increasing normal–appearing grey and white matter magnetization transfer ratio and myelin in postmortem multiple sclerosis brain. *Ann Neurol* 2004;56:407–15.

42. Feinsteins A, Kurkjeff F, Amato MP, et al. Correlation of measurement of change in MRI lesion volume in multiple sclerosis: a comparison of computer assisted techniques. *J Neurol Neurosurg Psychiatry* 1998;65:42–7.

43. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

44. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

45. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

46. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

47. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

48. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

49. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

50. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.

51. Feinsteins A, Kurkjeff F, Amato MP, et al. The contribution of MRI in assessing cognitive impairment in multiple sclerosis. *Neurology* 2002;55:2128–34.