Deficient learning and memory in early and middle phases of multiple sclerosis

IGOR GRANT,* W IAN MCDONALD,† MICHAEL R TRIMBLE,† EILEEN SMITH,‡ ROBERT REED*

From the Psychiatry Service, San Diego V.A. Medical Center and Department of Psychiatry, University of California at San Diego School of Medicine, La Jolla, California, USA,* the Institute of Neurology and National Hospitals, Queen Square and Maida Vale,† and the National Hospitals (Maida Vale),‡ London, UK

SUMMARY Forty-three patients with multiple sclerosis showed disturbances in short-term memory, learning, and delayed recall which were associated with years of active disease (average was 4·5 years), age, presence of flare-up, but not steroid/ACTH treatment. Unrecognised memory loss might be prevalent early in the natural history of multiple sclerosis and deserves neuropsychological assessment.

Cognitive decline commonly occurs in the later stages of multiple sclerosis.1 In the past three decades some of the qualitative and quantitative features of cognitive change have been delineated through neuropsychological inquiry. Many investigations of intelligence in multiple sclerosis have concluded that patients with demyelination show disproportionate drops in performance IQ while maintaining relatively good verbal skills.2 More comprehensive neuropsychological studies, utilising procedures such as the Halstead-Reitan Battery revealed patients with multiple sclerosis to have marked motor and perceptual-motor integrative deficits, generally intact verbal-language skills, and variable decrements in abstracting ability.3–10

A century has passed since Charcot first observed “enfeeblement of memory” in his patients11 and Gowers wrote of their “failure of memory”.12 Despite this, the neuropsychological research reviewed above has been singularly lacking in systematic assessment of memory. Surridge13 noted from structured psychiatric examination that approximately two-thirds of his patients showed intellectual decline, with amnesia being a central problem. Jambor’s14 related psychometric study found deficits in sentence learning and delayed recall of nonverbal information. These observations were based on a group of patients about half of whom had the disease 6–10 years, and a third, 11 years or longer.

Beatty and Gange15 examined a group of patients who were ill for approximately nine years. They found deficiencies in learning word lists and in delayed recall of a short story. Staples and Lincoln16 showed that multiple sclerosis patients with severe disability had worse learning and delayed recall both on verbal and nonverbal tasks, when compared to a matched sample of patients with muscular dystrophy.

These few systematic studies appear to confirm clinical observations and patient self-reports of impaired memory in association with established and advanced multiple sclerosis. The purpose of this study has been to expand our understanding of memory pathology in multiple sclerosis by: (1) examining a more complete range of memory processes than has been attempted heretofore; (2) considering whether memory pathology occurs commonly at a relatively early stage in the natural history of multiple sclerosis.

Method

SUBJECTS

(a) Multiple Sclerosis Group (N = 43)

Patients with multiple sclerosis were inpatients and outpatients at the National Hospitals, Queen Square and Maida Vale. Because some of these patients were also selected to be participants in a related study (life events and multiple sclerosis, to be reported separately) which called for enrolling patients in the early phase of their disease, the major-
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Healthy Comparison Group (N = 28)
This sample consisted of normal subjects who were comparable in age, sex distribution, and estimated intelligence to the patient group. There were 17 women and 11 men whose average age was 34.8 ± 10.3 (range 18-54), and whose Vocabulary scale score (from the Wechsler Adult Intelligence Scale, or WAIS) was 12.3 ± 2.7. Their vocabulary performance was comparable to that of the multiple sclerosis patients (11.9 ± 2.3; F(1,69) = 0.52, ns), and placed both groups into an average intelligence (est. IQ = 110) bracket.

NEUROPSYCHOLOGICAL TESTS
WAIS Vocabulary (18)
This subtest from the Wechsler Adult Intelligence Scale requires subjects to define words whose frequency of occurrence ranges from common to infrequent. Beyond reflecting language skills, performance on Vocabulary is highly correlated with overall intelligence.

TRAIL MAKING TEST (19)
This component of the Halstead-Reitan Battery has two parts. The simpler (Part A) requires subjects to use a pencil to connect circled numbers displayed on a page as quickly as they can. Part B displays numbers and letters; using a pencil, subjects must alternate between numbers and letters while maintaining the correct order in the number and letter strings. Time and accuracy are scored. This psychomotor test requires motor speed and steadiness as well as ability to order symbols presented in a visual array.

WECHSLER MEMORY SCALE (WMS) LOGICAL AND VISUAL MEMORY COMPONENTS (20)
Verbal Learning and Delayed Recall
We selected one of the prose passages from the WMS, read it to our subjects, asked them to repeat the passage verbatim after a 10 second delay, and immediately scored the number of memories produced. For the story, 22 memories was the maximum possible; if a person did not reach at least 15 memories on the first recollection, we presented the story again and required a second recall. This process was continued for a maximum of five trials if a person did not reach the criterion of 15 before then. Subjects were then asked to recall the prose passage 45 min later. Other testing occupied the interim. Three scores could be developed from this procedure: an immediate recall score (that is, number of memories at trial 1); a learning score (number of trials to reach criterion); and a delayed recall score (number of memories at 45 min).

Visual Learning and Recall
The Wechsler Memory Scale has three geometric stimulus cards, cards 1 and 2 containing one figure each, and card 3, two figures. Each card was presented to a subject for 10 seconds, and then the subject was required to reproduce each figure 10 seconds later on a piece of paper. The same principles of criterion learning and 45 min delayed recall were used as for the story above. Immediate, trials to criterion, and delayed recall scores were computed.

BROWN-PETerson TEST (21—22)
The Brown-Peterson distractor technique is a measure of short-term retention, and has been widely used as a test of short-term memory. Subjects were shown a card containing three consonants (a consonant trigram). There were two experimental manoeuvres which could, theoretically, either facilitate or interfere with short-term retention. To determine whether retention could be facilitated by rehearsal in our patients, they were required sometimes to recollect the trigram immediately after presentation, sometimes after a 2 second delay and other times after a 4 second delay. To determine our patients' sensitivity to "proactive interference" they were sometimes required to recollect the trigram after periods of 3, 6, 9 or 18 seconds of counting backwards by threes; other times no counting was required. There were 30 trigrams presented, randomised across the three rehearsal and five interference conditions. Subjects were allowed a rest period of 6 seconds between trials. The specific modification of the Brown-Peterson technique is very similar to the one employed by Butters and associates42 in their comparison of short-term memory in patients with early and late Huntington's Disease.

Results
Comparison of multiple sclerosis patients with non-patients
Table 1 compares neuropsychological performance of multiple sclerosis patients with that of non-patients. It will be seen that although multiple sclerosis patients were comparable in age and verbal intelligence to their nonpatient counterparts, they performed significantly worse on all other neuropsychological tests. Specifically, they were very significantly impaired on both Trail Making tests. It appears that the psychomotor speed demand of these tests was primarily responsible for the difficulties experienced by patients, since they had about
Table 1  Learning and memory in multiple sclerosis. Neuropsychological test results

<table>
<thead>
<tr>
<th>Tests</th>
<th>Groups*</th>
<th>Patients</th>
<th>Nonpatients</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>N = 43</td>
<td>N = 28</td>
<td></td>
</tr>
<tr>
<td>WAIS Vocabulary</td>
<td></td>
<td>11.9 ± 2.3</td>
<td>12.3 ± 2.7</td>
<td>1.52</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
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<td>ns</td>
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<tr>
<td>Trailmaking</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Part A (time-second)</td>
<td></td>
<td>51.4 ± 25.8</td>
<td>27.9 ± 9.3</td>
<td>21.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.0001</td>
</tr>
<tr>
<td>Trailmaking</td>
<td></td>
<td>105.8 ± 56.3</td>
<td>65.7 ± 22.8</td>
<td>12.6</td>
</tr>
<tr>
<td>Part B (time-second)</td>
<td></td>
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<td></td>
<td>0.001</td>
</tr>
<tr>
<td>WMS story</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trial 1 (# bits)</td>
<td></td>
<td>8.4 ± 3.4</td>
<td>12.6 ± 3.8</td>
<td>23.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.0001</td>
</tr>
<tr>
<td>Trials to criterion</td>
<td></td>
<td>3.2 ± 1.3</td>
<td>1.9 ± 0.6</td>
<td>26.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.0001</td>
</tr>
<tr>
<td>45 minute delay</td>
<td></td>
<td>13.5 ± 2.9</td>
<td>16.6 ± 3.0</td>
<td>19.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.0001</td>
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<tr>
<td>WMS Figures</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trial 1 (# bits)</td>
<td></td>
<td>13.6 ± 4.3</td>
<td>17.0 ± 2.8</td>
<td>13.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.001</td>
</tr>
<tr>
<td>Trials to criterion</td>
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<td>2.1 ± 1.3</td>
<td>1.0 ± 0.2</td>
<td>16.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.0001</td>
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<tr>
<td>45 minute delay</td>
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<td>15.0 ± 3.9</td>
<td>16.8 ± 2.8</td>
<td>4.21</td>
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<tr>
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<td></td>
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<tr>
<td>Brown-Peterson</td>
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</tr>
<tr>
<td>Total % correct</td>
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<td>59.3 ± 19.0</td>
<td>69.8 ± 21.1</td>
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<td></td>
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<td>0.04</td>
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<tr>
<td>No interference</td>
<td></td>
<td>98.6 ± 6.4</td>
<td>95.2 ± 8.9</td>
<td>4.19</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.05†</td>
</tr>
<tr>
<td>Interference</td>
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<td>49.6 ± 23.0</td>
<td>63.9 ± 25.4</td>
<td>6.06</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>0.02</td>
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<tr>
<td>No delay</td>
<td></td>
<td>45.3 ± 25.9</td>
<td>57.6 ± 29.7</td>
<td>3.37</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ns</td>
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<tr>
<td>4 second delay</td>
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<td>52.9 ± 26.3</td>
<td>62.1 ± 28.8</td>
<td>1.91</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ns</td>
</tr>
</tbody>
</table>

*m*mean ± SD  
†Multiple sclerosis patients better

Table 2  Learning and memory in multiple sclerosis. Predicting test performance with years of active disease, age, acute/quiescent status and education

<table>
<thead>
<tr>
<th>Dependent measures</th>
<th>Cumulative % accountable variance*</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Years of active disease</td>
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<tr>
<td>Trailmaking Part A</td>
<td>0.50</td>
</tr>
<tr>
<td>Trailmaking Part B</td>
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<tr>
<td>Brown-Peterson</td>
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<tr>
<td>Total % correct</td>
<td>0.28</td>
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<tr>
<td>No interference</td>
<td>0.15</td>
</tr>
<tr>
<td>9 seconds interference</td>
<td>0.01</td>
</tr>
<tr>
<td>WAIS vocabulary</td>
<td>0.08</td>
</tr>
</tbody>
</table>

*These are cumulative $R^2$ values and can be read as % variance in test accounted for by addition of each predictor. The $R^2$ under column 4 is identical to $R^2$ for the entire multiple regression, where $R = $ multiple correlation. Example: For Trails B, years of active disease accounts for 46% of variance in that test; age yields no added predictive power; education adds 1% prediction, and acute/quiescent status adds a final 1% for a total $R^2 = 0.48$. The unadjusted multiple R is $\sqrt{0.48}$, or 0.69 for this regression.

†p < 0.05, ‡p < 0.01, §p < 0.001.

the same level of difficulty with the cognitively easy Trail Making Part A as with the somewhat more cognitively complex Part B.

The results of the memory tests demonstrated that motor speed and dexterity were not, however, the only influence operating. With the exception of the requirement to reproduce a diagram in the visual section of the Wechsler Memory Scale, none of the other tests had any motor component whatever. Despite this, patients exhibited significant difficulty in short-term retention (Brown-Peterson Test), and, in particular, they appeared to be sensitive to the disrupting effect of enforced mathematical calculations which were introduced as interference. This
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Inference is supported by the higher level of statistical separation between performance of patients and nonpatients on the Brown-Peterson "with interference" score than with the "no interference" condition (table 1). Opportunity for rehearsal up to 4 seconds appears to be somewhat, but not dramatically helpful in improving patients' performance.

Turning to learning and delayed recall, it will be seen that patients showed worse immediate recall of both verbal and visual information; they also showed impaired learning in both modalities, since they required more trials to reach the pre-established criterion; finally, although all but two patients did in fact reach the criterion, and many, by virtue of having had more trials received more exposure to the material, patients nevertheless recalled fewer bits of information after 45 min than did their nonpatient counterparts.

Characteristics of multiple sclerosis patients in respect to neuropsychological functioning

In our efforts to understand potential sources of neuropsychological disadvantage in the patient group, we considered several factors which could theoretically influence performance. These included increasing age, educational disadvantage, length of disease, disease status (that is, flareup vs quiescent), and presence of ACTH or steroid medications. It will be appreciated that many of these putative causal factors are intercorrelated; for example, older patients are likely to have had the disease longer. Similarly, patients who have had the disease longer tend to have more frequent flareups and are more likely to receive steroids or ACTH. For this reason, we decided to explore the possible influence of these various factors on neuropsychological test performance in a multivariate fashion.

In the first set of analyses we performed step-wise multiple regressions to predict various neuropsychological test scores from number of years of active disease, age, acute/quiescent status, and education. Years of active disease was entered first.* Table 2 summarises these regression results. It should be noted that the figures presented are $R^2$ values, and therefore can be read as proportion of variance accounted by the variable in question. To derive the actual partial correlation for variables or the multiple correlation it is necessary to take the square root of the values presented. Hence, the multiple correlation for Trails A is the square root of 0.51, or approximately 0.7.

Turning first to the tests which load on psychomotor speed, it will be seen that the number of years of active disease accounted for most of the variability in Trails A and B. Age, education and active/quiescent status added almost nothing to the explanatory power of these equations. On the other hand, short-term memory (Peterson total) was significantly affected by all four factors—about 10% of the variance in Peterson errors was explained by each of years that the disease had been active, age, education, and whether an acute flareup had occurred. Further exploration of the Peterson results showed that under the simplest learning conditions (no interference) years of active disease provided the greatest explanatory power; as the complexity of the task increased (9 seconds of counting backwards interference) years of active disease, native intelligence (education) and acute/quiescent status all became important. At the most complex level (18 seconds interference) the equation became insignificant. This is explained by the fact that the 18 second condition was so difficult that even nonpatients had problems scoring well on it; indeed, we found no difference between patients and controls on the 18 second interference trial.

Equations which attempted to predict Wechsler memory scores from these four variables failed to reach significance. At the same time, vocabulary score was significantly predicted by education, flareup, and years of active disease.

When we constructed a new set of equations entering medicated/unmedicated status (and dropping out education in order to keep the number of predictor variables to a manageable number) we found that the presence of ACTH or steroids did not provide any additional (unique) explanatory power in the neuropsychological performance of our patients. We believe this to be a reflection of the fact that medication status is very highly correlated with flareup status ($\chi^2 = 10.1, p < 0.002$).

Discussion

The results of our study suggest that patients in early and middle phases of multiple sclerosis, beyond exhibiting some of the expected impairments in psychomotor functioning, also suffer decrements in the major domains of memory functioning. These include disturbances in short-term memory, learning, and delayed recall of learned material. It appears that verbal and nonverbal memory are affected equally. Efficiency of short-term retention seems to be particularly vulnerable to the effects of proactive interference, and the opportunity to rehearse for up to 4 seconds does not appear to be substantially helpful in correcting this difficulty with

*Years of active disease is defined as the number of years in which a patient had a minimum of 7 days of significant symptomatology. We constructed this variable rather than the more traditional "years since multiple sclerosis was diagnosed" to avoid artificially equating patients who might have had only a few attacks interspersed with long quiescent periods with patients whose disease was clearly far more active.
short-term retention. It appears further that the
defect in short-term storage of information is related
to the number of years that a patient's disease has
been active, and to the activity of the disease
(flareup vs quiescence) at the time of testing. Medication
with ACTH or steroids, on the other hand, does not appear to explain short-term retention
deficit; but this might be due to our difficulty in
disentangling the possible independent effects of
acute flareup and drugs prescribed for that flareup.

How do our findings relate to previous studies of
memory functioning in multiple sclerosis? As we
indicated in our introductory remarks, very few
neuropsychological studies of this disorder
performed systematic explorations of memory. Sur-
ridge performed ratings of memory on his 108
patients on the basis of a clinical psychiatric exami-
nation. He reported that approximately two-thirds
of his patients had some intellectual deterioration,
and that the typical difficulty was amnesia. The
companion study by Jambor examined most of
these same patients with several formal memory
tests. Jambor found that multiple sclerosis patients
had difficulties with a sentence learning task, requiring
more trials than controls to learn the Babcock Sentences. This result is comparable to our present
observation that patients with multiple sclerosis
required more trials to learn our prose passage.
Jambor also reported that his patients had difficulties
on a task involving delayed recall of pictures.
This appears similar to our observation that delayed
recall of figures is impaired in multiple sclerosis, but
the lack of detail in Jambor's report makes direct
comparison of our results problematic. In relating
these Surridge and Jambor data to our own, it is
important to recognise that their patients were, in
general, ill for substantially longer periods than
ours. For example, approximately half of their
patients were ill for 6 to 10 years, and another third
had the disease for 11 to 25 years. In contrast, 60%
of our patients were within the first five years of
their initial diagnosis, 30% were 6 to 10 years post
diagnosis, and only 10% were 11 to 15 years post
diagnosis. The fact that we found our sample to have
memory difficulties which were qualitatively similar
to those of Surridge and Jambor suggests that mem-
ory disturbance might begin evolving relatively early
in the natural history of multiple sclerosis.

The second systematic study of memory in multi-
ple sclerosis was that of Beatty and Gange whose
26 patients had a mean disease length of 9 years.
The task they employed, which involved free recall
of 24 words presented at 2 second intervals for four
trials, is not directly comparable to our procedure,
though it probably has most in common with our
story learning to criterion technique. These authors
found that their patients recalled significantly fewer
words over the four trials than did their controls,
and that although the patients benefited somewhat
from repeated testing, they did not gain at the same
rate as the nonpatients. Beatty and Gange's multiple
sclerosis patients were also deficient on a verbal
delayed recall task, which consisted of 25 multiple
choice questions about a short story presented 30
min previously. This finding is consistent with our
observation that patients had difficulty recalling a
prose paragraph after 45 min.

The third neuropsychological study to assess
learning and memory was that of Staples and Lin-
coln. Their approach to memory assessment was
more comprehensive than that of the other two
studies, involving as it did the full Wechsler Memory
Scale with the addition of verbal and visual
delayed recall requirements. Part of their study
involved comparing performance of multiple
sclerosis patients to published norms, while another
part compared 29 patients with multiple sclerosis to
a matched group of muscular dystrophy patients.
The results of the controlled investigation showed
that patients with multiple sclerosis were impaired in
immediate recall of Wechsler figures but not stories;
that they were impaired on delayed recall both of
verbal and nonverbal material; and that their verbal–verbal paired associate learning was also
deficient. Once again, we would note that the
patients in the Staples and Lincoln study were
described by the authors as having "severe long-
standing disability" with a mean duration of 15-9
years.

We consider that our findings both confirm and
extend the results of these several previous investig-
ations into the nature of memory disturbance in
multiple sclerosis. Our study is distinctive in two
respects. It is the first of which we are aware that has
focused on processes of short-term memory, which
are presumed to reflect the initial encoding of
information. Although the neuroanatomic under-
pinnings of memory are still imperfectly understood,
it is thought that the lesions of the dorsomedial
thalamic nuclei are especially likely to be associated
with deficits in learning, whereas medial temporal
lobe pathology interferes with ability to recall
information which was once learned. In the latter
instance, there appears to be a lateralisation of func-
tion such that recall of verbal material is influenced
by the left temporal lobe, while recall of nonverbal
material is related to right temporal function. If
these neuroanatomic models are correct, then our
findings would suggest that many of our patients
have foci of demyelination in the diencephalon
while others may have lesions bilaterally in the hip-
pocampus and related temporal structures. Unfor-
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Unfortunately, brain imaging (CT scanning) was not sufficiently sophisticated in our study to detect small lesions in these areas. With the advent of nuclear magnetic resonance (NMR) technology it will soon become feasible to validate these neuropsychological observations of memory disturbance with detailed information concerning the distribution, size, and activity of plaques of demyelination.

The second unique feature of our study has to do with the phase in the natural history of their disease in which our patients found themselves. The majority experienced active disease for less than 5 years, and there were only four patients whose disease had been active 11-15 years. Further, only two of our patients were chronically hospitalised, the rest living either independently or with family in the community. Despite their relative skew toward lack of gross physical or cognitive disability, our patients did manifest substantial memory difficulties.

In summary, patients in early and middle phases of multiple sclerosis often showed disturbances in short-term memory, learning, and recall of verbal and nonverbal information. Some of these memory disturbances were correlated with number of years of active disease, acute versus quiescent status of the disorder, age, and background educational level of the patient. ACTH and steroid medications appeared to play little or no role in the memory difficulties. We conclude that subtle, unrecognised memory disturbance is common even in early multiple sclerosis and recommend that future, more detailed investigations of memory and other neuropsychological functions be coupled with advanced brain imaging technology.

The authors are indebted to Dr PC Gautier-Smith, for his enthusiastic support of this study, and particularly, his willingness to identify patients who might be suitable as research subjects. Professors John Marshall, Newsom-Davis, and PK Thomas were also helpful in allowing us access to patients, as were Drs RW Ross Russell, JA Morgan-Hughes, PRudge, AN Gale, R Clifford Jones, AJ Lees, and E Byrne. Ms Marge Zeitsman assisted in scheduling some of our patients, and Ms Debi Taylor helped in the preparation of this manuscript.

This research was conducted while Dr I Grant was honorary research fellow at the Institute of Neurology, supported by the Foundations Fund for Research in Psychiatry, 1980-81.

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*J Neurol Neurosurg Psychiatry* 1984 47: 250-255
doi: 10.1136/jnnp.47.3.250

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