Macrocytosis in multiple sclerosis. A study in 82 de novo Arab patients

A S Najim Al-Din, M Khojali, H Habbosh, S Farah, A R Idris, F Al-Muhtasib

Abstract

Macrocytosis, without anaemia, was common in 82 de novo multiple sclerosis patients compared with a similar number of age and sex matched controls. This was an early phenomenon in the course of the disease and was not influenced by the age of the patients nor the duration of the disease. None of the patients proved to have pernicious anaemia, yet the similarity in the geographical and sex distribution as well as the similarity in HLA associations of multiple sclerosis and pernicious anaemia may indicate that both diseases are under similar genetic influence.

Mild macrocytosis has been reported in cases of multiple sclerosis (MS) without anaemia. This has raised the question of whether an abnormality of vitamin B₁₂ absorption, transport or metabolism was associated with MS. More recently MS patients with unusual vitamin B₁₂ deficiencies were described. To verify this observation we retrospectively studied the haemoglobin (Hb) and the red cell mean corpuscular volume (MCV) in our group of Arab MS patients.

Patients and methods

The study population comprised 82 clinically definite MS Arab patients who fulfilled the Poser et al criteria. All of them presented within four years of the onset of the first symptom and were not diagnosed before or had received specific medications before presentation to us. Their Hb and MCV values were recorded before the initiation of any form of specific therapy. Each MS patient was matched for sex and age with a neurological patient who attended a neurology outpatient clinic during the same period (1985-89). Groups of patients who had conditions which led to abnormal Hb and/or MCV (epileptics on anticonvulsants, dementia, depression, neuropathies and subacute combined degeneration of the cord) were excluded. Among the 82 controls 32 had tension headache, 27 migraine, 15 spondylotic lumbar or cervical spinal diseases, six benign intracranial hypertension and two had peripheral vestibular disease. None of the MS patients nor the controls were on drugs that were likely to alter either measurement. All blood counts were measured using Coulter Blood Counter Model S-Plus (Coulter Electronics, Luton, UK). All data were entered into the vax-11/8810 computer and analysis was carried using SPSS package.

Results

The age, sex distribution and haematological data of patients and controls are shown in table 1. Fifty one (62%) of the patients had a duration of disease of one to 12 months before haematological evaluation and only five (6%) had a duration of three to four years before these tests were performed. The mean duration of disease was 15-9 months (range 1-48 months) among all cases. The two groups were similar in age, sex distribution and mean haemoglobin levels. In both groups the mean (SD) MCV was found to be within normal limits, the normal reference range for MCV in our laboratory is 84 (7) fl. The MS patients exhibited an elevated MCV [87-5 (6-4) fl] when compared with controls [84-1 (5-6) fl] and the difference was found to be highly significant (p < 0-0001). This difference was also observed when comparing MS female and male patients with their respective sexes among the controls and it was found to be highly significant, p < 0-009, p < 0-003, respectively (table 2). There was no influence of age on the level of MCV among both patients and controls. Likewise the duration of the disease before haematological tests were performed had no influence on the MCV.

Only four females from both groups had an MCV lower than the normal limits for our laboratory (77 fl). Four (5%) of the controls had an MCV higher than the upper limits for normal (91 fl), this ranged from 91-6-95-7 fl. Abnormally high MCV was observed in 22 (27%) of the patients with a range of 91-6-100 fl. Sixty three (77%) of the patients compared with 40 (49%) of the controls had an MCV higher than the normal mean. Even when the 22 patients with abnormally high MCV and the four with the abnormally low MCV were excluded, still 73% of the remaining 56 MS patients with normal MCV had a measurement which was higher than the normal mean.

Only two MS patients, one with an MCV of 94-8 and the other 100 fl had a low haemoglobin (10-8 and 10-9 gm/dl respectively). In neither was the B₁₂ metabolism studied on first presentation.

The serum B₁₂ and folate were assayed in 14 of the patients reported above, who were not on azathioprine, in 1989 and all the values were within normal limits.
Table 1  Sex, mean (SD) age, haemoglobin and MCV among (B12) multiple sclerosis and controls and neurological control

<table>
<thead>
<tr>
<th></th>
<th>MS (n = 82)</th>
<th>Control (n = 82)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) age (years)</td>
<td>31.7 (8.7)</td>
<td>31.7 (8.7)</td>
<td>NS</td>
</tr>
<tr>
<td>Sex ratio M:F</td>
<td>30:52</td>
<td>30:52</td>
<td></td>
</tr>
<tr>
<td>Mean (SD) +g/dl</td>
<td>13.8 (1.6)</td>
<td>13.5 (1.5)</td>
<td>NS</td>
</tr>
<tr>
<td>Mean (SD) MCV</td>
<td>87.5 (6.4)</td>
<td>84.1 (5.6)</td>
<td></td>
</tr>
<tr>
<td>Mean (SD) duration months</td>
<td>15.9 (13.6)</td>
<td></td>
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</tr>
</tbody>
</table>

*Normal laboratory value = 84 (7-0) fl.

Table 2  Mean (SD) age, haemoglobin and MCV by sex among MS patients and controls

<table>
<thead>
<tr>
<th></th>
<th>Females</th>
<th></th>
<th>Males</th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>MS n = 52</td>
<td>Control n = 52</td>
<td>MS n = 30</td>
<td>Control n = 30</td>
</tr>
<tr>
<td>Mean (SD) age (years)</td>
<td>30.1 (8.5)</td>
<td>30.1 (8.5)</td>
<td>34.5 (8.5)</td>
<td>34.5 (8.5)</td>
</tr>
<tr>
<td>Mean (SD) +g/dl</td>
<td>13.0 (1.2)</td>
<td>12.8 (1.3)</td>
<td>15.0 (1.5)</td>
<td>14.7 (1.1)</td>
</tr>
<tr>
<td>Mean (SD) MCV</td>
<td>86.0 (6.8)*</td>
<td>82.7 (5.7)</td>
<td>90.1 (4.6)*</td>
<td>86.4 (4.3)</td>
</tr>
<tr>
<td>Mean (SD) duration month</td>
<td>14.4 (13.0)</td>
<td></td>
<td>18.5 (14.4)</td>
<td></td>
</tr>
</tbody>
</table>

*Significant difference p = 0.009 t = 2.67.
†Significant difference p = 0.003 t = 3.07.
‡Difference not significant when comparing females and males.

Discussion

The observation by Reynolds of unusual vitamin B12 deficiencies in 10 patients might have implications on the pathoetiology of MS.5 The report by his group of macrocytosis without anaemia in 27 MS patients gives further support for this observation.2

We studied this phenomenon in 82 de novo patients with relatively short duration of the disease to overcome any effect of drug or dietary influences. Our data supports the study by Reynolds4 that a tendency for macrocytosis without anaemia is noted in MS. The majority of our MS patients were thus found to have an MCV value higher than the normal mean early in the course of their disease before the start of any specific therapies. More significant is the observation that 27% of our patients had abnormally high MCV. Yet anaemia was exceptional as it was noted in only two patients with a high MCV. The macrocytosis was not a function of the duration of the disease and was frequently discovered in patients who were studied within a week of presentation with the first symptom. Although we studied the serum B12 and folate in a limited number of patients (several years after the onset) these were normal.

The significance of our observation and previous reports is subject to debate. Although none of our patients proved to have pernicious anaemia (PA) it is interesting that both MS and PA are most common in whites of northern European ancestry and both affect females more than males.2 Both diseases are of unknown aetiology where both genetic susceptibility and autoimmune reaction are postulated. Furthermore, PA is reported to have a geographical distribution similar to MS.5 Interestingly the HLA studies in PA have shown an association with HLA-A3,7 HLA-B7,8 HLA-DRW29 and DR211 similar to MS.

Whether metabolic abnormalities of vitamin B12 play a role in the aetiology of MS is not yet clear. The phenomenon observed probably reflects a genetic overlap of two different diseases. It is possible that both of these diseases require a common genetic susceptibility but are manifest under the influence of different environmental factors.

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