Effect of Methylprednisolone on CSF IgG Parameters, Myelin Basic Protein and Anti-Myelin Basic Protein in Multiple Sclerosis Exacerbations

Kenneth G. Warren, Ingrid Catz, Verona M. Jeffrey and Dorothy J. Carroll

ABSTRACT: Clinical exacerbations of multiple sclerosis (MS) are characterized by elevated levels of cerebrospinal fluid (CSF) myelin basic protein (MBP). The purposes of this study were to determine whether anti-MBP antibodies are present in increased titer in CSF of MS patients with exacerbations, and whether they can be suppressed by the administration of immunosuppressive dosages of methylprednisolone (MP). A solid phase radio-immunoassay (RIA) was used to detect free and total anti-MBP antibodies before and after acid hydrolysis of CSF. In MS exacerbations, the majority of elevated anti-MBP is in the free form. With the exception of subacute sclerosing panencephalitis (SSPE) and some cases of post infectious encephalomyelitis, anti-MBP antibodies are not present in either MS patients in remission or in non-MS controls. Anti-MBP levels remained elevated over a 10 day period when patients are managed by bed rest only or when treated with intravenous (IV) ACTH. IV administration of MP in "high" (160 mg/day) or "mega" (2 g/day) dosages produces a highly significant reduction of both MBP (p<0.01) and anti-MBP (p<0.001) levels. Total intrathecal IgG synthesis is also significantly suppressed by IV-MP but not by ACTH.

RESUME: Le traitement par la methylprednisolone d'une elevation des anticorps anti-proteine basale de la myeline dans les exacerbations de la sclerose en plaques (SEP) est caracterise par une diminution de la teneur en anticorps anti-MBP. Les anticorps anti-MBP libres et total dans le liquide cephalo-rachidien (LCR). Le but de la presente etude etait de determiner si les anticorps anti-MBP etaient en teneur plus elevees dans le LCR au cours des poussées de la maladie et de verifier si ces modifications peuvent etre supprimées par l'administration de doses immunosuppressives de methylprednisolone (MP). Le dosage des anticorps anti-MBP libres et totaux dans le LCR, après hydrolyse acide, s'est fait grâce à un radio-immunoassay (RIA). Lors des poussées de SEP la plus grande partie de l'élévation des anti-MBP est sous forme libre. Les anticorps anti-MBP ne sont pas présents chez les patients SEP en rémission ou chez les témoins non-SEP. Ils sont parfois présents dans la SSEP et chez quelques cas d'encéphalomyélite post-infectieuse. Les teneurs d'anti-MBP restent élevées sur une période de 10 jours si les patients sont traités par le repos au lit ou à l'aide d'ACTH intraveineux. L'administration intraveineuse (IV) de MP à des doses "élévées" (160 mg/jour) ou "mégas" (2 g/jour) produisit une réduction fortement significative des MBP (p < 0.001). La synthèse intrathécale des IgG totaux est également supprimée par le MP-IV, mais non par l'ACTH. Il semblerait donc que le MP en dose adéquate pourrait remplacer l'ACTH dans le traitement des poussées cliniques de SEP.
The 40 patients entered into this study had clinically definite MS and were experiencing exacerbations. They were randomly allocated to four treatment groups of 10 patients each: group A — non-treated control group; group B — ACTH 60 units/day intravenously for 10 days; group C — “high” dose methylprednisolone (MP) 160 mg/day intravenously for 10 days and group D — “mega” dose MP 2 g/day intravenously for 10 days. Matched CSF and serum samples were obtained simultaneously from all patients before treatment and within 12 hours of terminating the medication. The untreated control group also had samples obtained 10 days apart.

Total protein (TP), albumin (alb), IgG, MBP as well as free and total anti-MBP levels were measured in CSF. Alb and IgG levels were also determined in serum. The degree of breakdown of the BBB was estimated by the CSF/serum alb ratio. Intrathecal IgG synthesis was measured by CSF IgG/alb ratio, an IgG Index and daily rate of CNS IgG synthesis. Cerebrospinal fluid MBP, an indicator of disease activity, was determined by radioimmunoassay (RIA).

Free and total CSF anti-MBP levels were determined before and after acid hydrolysis by a solid phase RIA. CSF samples were acidified for 1 hour (pH = 3) in order to dissociate possible preformed immune complexes. These samples were neutralized (pH = 7) prior to performing the assay. Final IgG concentration was adjusted to 0.010 g/l in all CSF samples. Aliquots of 100 µl CSF before and after acid hydrolysis were incubated for 2 hours at room temperature (RT) in immulon plates coated with human-MBP (1 µg/well). After 5 washes, goat anti-human Ig was added and incubated for 1 hour at RT. After another 5 washes radio-labelled (125) Staph A Protein (50,000 counts/well) was added and incubation continued for 2 more hours at RT. Finally, after 5 more washes the wells were counted individually and results were expressed as % bound total radioactivity (TC). Blanks were performed with each sample to determine nonspecific adherence to uncoated immulon plates. The nonspecific binding (≤ 1% TC) was subtracted from the respective sample counts. When CSF samples with initially high IgG and anti-MBP values were serially diluted, the anti-MBP levels were parallel to the IgG concentrations. This assay was also validated by absorbing CSF anti-MBP with MBP prior to anti-MBP assay. Absorption to MBP resulted in complete elimination of anti-MBP from samples that initially had high anti-MBP titers. Two known positive and two negative controls were included with each run. Within assay variability (CV) for 25 sets of duplicates was 3.86 while between assay variability (CV) for one set of quadruplicates over 10 different runs was 7.75.

Intragroup mean and standard deviations (SD) were calculated for each parameter before and after treatment. The pre and post treatment values were statistically compared by Student’s t test.

**RESULTS**

“Normal” levels of free and total CSF anti-MBP were determined in a group of 88 control patients consisting of 25 with psychoneurosis, 32 with degenerative disc disease and 31 with various neurological diseases exclusive of MS. CSF free anti-MBP was 0.5±0.3 while total anti-MBP was 2.3±0.5. In this series, the only neurological diseases associated with relatively high titers of CSF anti-MBP in the bound form only were subacute sclerosing panencephalitis and 2 of 8 cases of post infectious encephalomyelitis.

The results of the 4 treatment methods are summarized in Table 1.

Group A — non-treated control group: This group showed no change in any of the measured parameters over the 10 day period. CSF TP and alb as well as the degree of breakdown of the BBB remained approximately the same. Total intrathecal IgG synthesis as measured by three different parameters remained high. CSF-MBP also remained elevated indicating continuation of disease activity. Both free and total CSF anti-MBP levels showed no tendency to drop towards normal values.

Group B — ACTH treatment: With one exception, results of this group were similar to group A. CSF-MBP showed a modest but significant drop from 21.8±2.2 to 16.0±3.2 µg/L (p<0.05). The BBB, intrathecal IgG synthesis and CSF anti-MBP showed no significant change.

Group C — “high” dose MP and Group D — “mega” dose MP: CSF TP and alb remained unchanged. The degree of breakdown of the BBB (Figure 1) also remained elevated in these 2 groups. Absolute CSF IgG was significantly lowered from 0.97±0.013 to 0.046±0.002 g/L (p<0.01) by “mega” dose MP; total intrathecal IgG synthesis was also more significantly suppressed by “mega” dose treatment: CSF IgG/alb ratio was significantly lowered from 0.58±0.06 to 0.23±0.07 (p<0.01) by
**DISCUSSION**

Although intrathecal IgG synthesis is significantly increased in MS patients with exacerbations, it was not the purpose of this study to determine how much of the total CSF IgG was

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### Table 1: Effects ACTH and Methylprednisolone on CSF parameters (mean ±2SD) of MS patients with exacerbations

<table>
<thead>
<tr>
<th></th>
<th>No Treatment (A)</th>
<th>ACTH: 60 u/d (B)</th>
<th>MP: 160 mg/d (C)</th>
<th>MP: 2 g/d (D)</th>
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<tbody>
<tr>
<td></td>
<td>Pre</td>
<td>Post</td>
<td>Pre</td>
<td>Post</td>
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<tr>
<td>CSF Total Protein</td>
<td>0.64±0.30</td>
<td>0.60±0.25</td>
<td>0.54±0.20</td>
<td>0.46±0.14</td>
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<tr>
<td></td>
<td>0.35±0.10</td>
<td>0.36±0.10</td>
<td>0.36±0.10</td>
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<tr>
<td>CSF Albumin</td>
<td>0.29±0.086</td>
<td>0.29±0.086</td>
<td>0.28±0.048</td>
<td>0.28±0.052</td>
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<tr>
<td>0.164±0.048</td>
<td>0.029±0.029</td>
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<tr>
<td>1000 (CSF/Serum Alb)</td>
<td>7.9±2.1</td>
<td>8.1±2.0</td>
<td>9.2±1.3</td>
<td>8.5±1.5</td>
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<tr>
<td>3.7±1.8</td>
<td>0.106±0.022</td>
<td>0.120±0.022</td>
<td>0.086±0.066</td>
<td>0.069±0.013</td>
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<tr>
<td>CSF IgG</td>
<td>0.106±0.022</td>
<td>0.120±0.022</td>
<td>0.086±0.066</td>
<td>0.069±0.013</td>
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<tr>
<td>0.024±0.016</td>
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<tr>
<td>IgG Index</td>
<td>0.42±0.05</td>
<td>0.44±0.11</td>
<td>0.31±0.10</td>
<td>0.29±0.08</td>
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<tr>
<td>0.14±0.06</td>
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<td>0.14±0.06</td>
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<tr>
<td>MBP</td>
<td>1.68±0.12</td>
<td>1.64±0.16</td>
<td>1.10±0.14</td>
<td>1.10±0.22</td>
</tr>
<tr>
<td>0.5±0.1</td>
<td>1.64±0.16</td>
<td>1.10±0.14</td>
<td>1.10±0.22</td>
<td>1.10±0.22</td>
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<tr>
<td>CNS IgG Synthesis</td>
<td>32.0±6.0</td>
<td>36.0±8.0</td>
<td>17.2±7.2</td>
<td>15.8±6.2</td>
</tr>
<tr>
<td>0.0±4.0</td>
<td>36.0±8.0</td>
<td>17.2±7.2</td>
<td>15.8±6.2</td>
<td>16.0±2.8</td>
</tr>
<tr>
<td>MBP</td>
<td>7.4±2.6</td>
<td>7.0±2.8</td>
<td>21.8±2.2</td>
<td>16.0±3.2</td>
</tr>
<tr>
<td>0.0±2.0</td>
<td>7.0±2.8</td>
<td>21.8±2.2</td>
<td>16.0±3.2</td>
<td>16.0±3.2</td>
</tr>
<tr>
<td>Free Anti-MBP</td>
<td>20.0±2.0</td>
<td>20.5±2.0</td>
<td>19.0±2.5</td>
<td>20.5±4.0</td>
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<tr>
<td>0.4±0.2</td>
<td>20.5±2.0</td>
<td>19.0±2.5</td>
<td>20.5±4.0</td>
<td>20.5±4.0</td>
</tr>
<tr>
<td>Total Anti-MBP</td>
<td>22.5±2.5</td>
<td>23.6±3.5</td>
<td>22.5±2.5</td>
<td>22.5±4.0</td>
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<tr>
<td></td>
<td>22.5±2.5</td>
<td>23.6±3.5</td>
<td>22.5±2.5</td>
<td>22.5±4.0</td>
</tr>
</tbody>
</table>

Pre = data prior to treatment (day 0)  Post = data after 10 days of treatment.

Student’s t test p value:  = p<0.05  = p<0.01  = p<0.001

Group A (non-treated): No significant change occurred in any parameter.

Group B (ACTH 60 u/day): A modest but significant (*) drop occurred in CSF MBP levels only.

Group C (Methylprednisolone 160 mg/day): Significant reductions occurred in IgG Index (**), daily CNS IgG synthesis (**), CSF-MBP (**), and both free and total anti-MBP (**).

Group D (Methylprednisolone 2 g/day): Significant reductions occurred in CSF IgG (**), CSF IgG/albumin ratio (**), IgG Index and daily IgG synthesis (**), CSF-MBP (**), and free and total anti-MBP levels (**).

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antibody with specificity for MBP. Nevertheless, CSF anti-MBP antibodies were detected in these patients by a solid phase RIA using human MBP (whole molecule). Recently, Panitch et al., using a RIA with guinea pig MBP, reported low anti-MBP levels in MS patients with exacerbations. While other authors have found antibodies against MBP in MS CSF, there have also been reported negative results. In the present report, high CSF anti-MBP levels were found in MS patients with clinical exacerbations associated with elevated CSF-MBP levels. With the exception of a patient with SSPE and 2 of 8 patients with post infectious encephalomyelitis, a large control group of psychiatric and other neurological diseases had no detectable anti-MBP in the CSF. The fact that anti-MBP was somewhat higher after acid hydrolysis may be explained by the presence of immune complexes in CSF. The data illustrate high free to total anti-MBP ratios, therefore an exacerbation of MS may be dependent upon a pulse of anti-MBP molecules being produced intrathecally.

When MS exacerbations were managed for 10 days by bed rest only, all CSF parameters remained unchanged and abnormally elevated. The BBB remained impaired permitting leakage of serum proteins into the intrathecal compartment. CNS IgG synthesis continued at the same rate, and disease activity as indicated by elevated CSF-MBP was maintained. Anti-MBP antibodies persisted at high levels.

It has previously been suggested that short term, high dosage use of intramuscular ACTH hastened improvement of symptoms and signs of MS patients with exacerbations and that de novo CNS IgG synthesis was reduced by ACTH therapy. In the present report, the IV administration of 60 units ACTH/day for 10 days had a negligible effect on all CSF parameters. As noted in the untreated group, the BBB remained impaired and

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**Figure 3** — Intrathecal IgG synthesis as indicated by daily CNS IgG synthesis (mean ±2SD) is illustrated for the four treatment groups. A significant reduction occurred in both methylprednisolone groups (C and D).

**Figure 4** — Disease activity as indicated by CSF-MBP (mean ±2SD) levels is illustrated for the four treatment groups. A modest but significant reduction was produced by ACTH (B) and a more significant reduction was produced by methylprednisolone therapies (C and D).

**Figure 5** — Free and total CSF anti-MBP levels (mean ±2SD) are illustrated for the four treatment groups.

*Hatched area* = free anti-MBP (prior to acid hydrolysis).

*Total area* = total anti-MBP (post acid hydrolysis).

- **Group A** — no change.
- **Group B** — no change.
- **Group C and D** — a highly significant reduction in both free and total anti-MBP after 10 days of IV methylprednisolone therapy.
intrathecal IgG synthesis was maintained. While CSF anti-
MBP levels persisted there was a modest reduction of CSF-MBP.
Uncontrolled clinical observations have suggested that IV-MP
may be of clinical value for MS patients with active disease.23-27
A dose of 1g MP per day administered to MS patients with
exacerbations reduced contrast enhancing lesions detected by
computed tomography.28 In the present study the BBB as mea-
sured by the CSF/serum alb ratio remained unchanged. It has
also been suggested that a large dose of 1g MP per day may
result in suppression of CNS IgG synthesis.29 In this report two
dosage regimes of MP were studied: a “high” dose (160 mg/day)
as well as a “mega” dose (2 g/day). In contrast to the untreated
or ACTH treated groups the MP therapies produced marked
changes in many of the CSF parameters. Although the “mega”
dosage produced a more significant drop, intrathecal IgG syn-
thesis was significantly reduced by both MP regimes. Both MP
therapies similarly reduced disease activity as measured by
CSF-MBP. Free and total CSF anti-MBP levels were also
significantly reduced by MP in either dosage. The effects of the
“high” dose administration was nearly equivalent to that pro-
duced by the “mega” dose. Adverse clinical side effects including
acute psychosis and attempted suicide, hair loss, steroid with-
drawal skin rashes, catatric formation and aseptic necrosis of
the femoral head were unacceptably high in the group of patients
who received “ mega” dose therapy in this study. These side
effects did not occur in patients who received the lower dosage
(160 mg/day). According to the results observed in this study, it
may not therefore be necessary to give “ mega” dosages of MP
to MS patients. However, biological effects not measured in
this study may only occur with “ mega” dosage therapy.

In conclusion, the results of this study suggest that IV-MP is
of greater value than ACTH in reducing certain CSF parameters.
Unpublished longitudinal case studies of MP treated patients in
our Multiple Sclerosis Research Clinic have shown that intrathecal
IgG synthesis, CSF MBP and anti-MBP levels remained sup­
pressed and the disease remained clinically inactive for periods
up to 6 to 12 months. Conversely, other cases with more aggres-
sive disease, responded negligibly to MP therapy.

Since some of the intrathecally produced IgG may be protective,
suppression of total IgG by MP may be detrimental. However,
suppression of potentially destructive IgG components such as
anti-MBP would convert an active disease process into an
inactive one and the protective antibodies may no longer be
required. Because CSF anti-MBP levels are increased in MS
patients with active disease but not in patients in remission
anti-MBP may be involved in the pathogenesis of demyelination.
Doubleblind controlled clinical trials of MP therapy with mon-
toring of the role of anti-MBP are required.

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