Pregnancy and multiple sclerosis: a longitudinal study of 125 remittent patients

E Roullet, M-H Verdier-Taillefer, P Amarenco, G Gharbi, A Alperovitch, R Marteau

Abstract

The relationship between pregnancy and multiple sclerosis (MS) was assessed in a clinic-based, prospectively followed, population of 125 patients with a remittent onset of MS who had been followed for a mean (SD) of 10.3 (0.1) years. Thirty three women had a total of 49 pregnancies of which 32 had been full term and 17 terminated. There was a three-fold increase in the relapse rate per year during the first three months following delivery, compared with the baseline period of the same patients [1.62 (0.38) vs 0.51 (0.08) p = 0.05]. During pregnancy itself, the relapse-rate was not different from baseline. The overall relapse rate of the pregnancy group was lower than that of a control group without pregnancies after MS onset, but similar to that of patients who had children after MS onset, but no pregnancy during follow up. Pregnancy did not lead to increased disability. These results confirm that post partum increase in relapse rate is the main event related to pregnancy in MS and underline the difficulties of undertaking prospective studies in this field.

Methods

PATIENTS AND COLLECTION OF DATA

Since 1967, MS patients in our clinic have been evaluated in a standardized way. At the first visit, age at MS onset, number of children and date of pregnancies are collected retrospectively. Clinical assessment is made by a group of 3 neurologists who use uniform criteria for diagnosis, events, and disability. Diagnosis was based on Schumacher's criteria before 1983 and on Poser's thereafter. Events and disability are recorded prospectively at each visit. All patients are examined at least once a year and more frequently if worse; they are instructed to phone at each worsening of symptoms, and are then examined if judged necessary. The occurrence of new symptoms or the noticeable aggravation of old symptoms, lasting more than 24 hours, with or without objective changes on clinical examination are considered as a relapse, whereas isolated paroxysmal symptoms such as Lhermitte's sign or trigeminal neuralgia are not. Progression is the progressive worsening of symptoms and signs over a period of six months or longer. The overall evolution of MS is thus of 4 subtypes: remittent, remittent-progressive, progressive from onset or unclassified. When a patient has entered a progressive phase, he remains classified as progressive, even if relapses occur later; relapses are not scored during the progressive phase because even prospective data are judged unreliable in this setting. Permanent disability is assessed through Kurtzke's Expanded Disability Status Scale (EDSS). From 1967–84, disability had been assessed through our own disability scale and was then converted to EDSS. In female patients, the occurrence and outcome of all pregnancies were recorded, allowing precise location of the distribution of relapses during pregnancy and postpartum. The end of the study was 1 September 1989 or the last visit before that date. Disability and evolutive type were assessed and the total number of relapses
Pregnancy and multiple sclerosis: a longitudinal study of 125 remitted patients

Table 1: Comparison of initial data in 3 groups of remitted MS patients

<table>
<thead>
<tr>
<th>Women with pregnancies during follow up (Pregnancy group)</th>
<th>Women with pregnancies after MS onset (Control group 1)</th>
<th>Women without pregnancies after MS onset (Control group 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n = 33</td>
<td>n = 17</td>
<td>n = 75</td>
</tr>
<tr>
<td>Age (years) mean (SEM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>38.5 (1.5)</td>
<td>43.1 (2.1)</td>
<td>39.4 (1.2)</td>
</tr>
<tr>
<td>Age at onset (years) mean (SEM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>23.9 (0.7)</td>
<td>23.9 (1.2)</td>
<td>26.7 (0.7)</td>
</tr>
<tr>
<td>Duration of MS at entry (years) mean (SEM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.6 (0.5)</td>
<td>7.3 (1.1)</td>
<td>2.8 (0.4)</td>
</tr>
<tr>
<td>Duration of relapsed follow up (years) mean (SEM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10.5 (1.1)</td>
<td>10.0 (1.7)</td>
<td>8.9 (0.8)</td>
</tr>
<tr>
<td>EDSS at entry mean (SEM)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.15 (0.22)</td>
<td>3.44 (0.50)</td>
<td>2.51 (0.17)</td>
</tr>
<tr>
<td>p</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ns</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

RESULTS

PREGNANCY GROUP

A total of 49 pregnancies were observed during follow up. Twenty one women had one pregnancy, 8 had two and 4 had three pregnancies. Thirty two pregnancies were full-term pregnancies and 17 spontaneous or voluntary abortions; seven of these were the only pregnancy in 6 women. Deliveries of full-term pregnancies were uneventful, except in the only twin pregnancy in which placenta praevia caused the death of one twin. One other child had multiple malformations. The deliveries and children of the three patients treated by azathioprine during full-term pregnancies were normal. Premature termination of pregnancy before 3 months was caused by spontaneous abortion in 4 cases and was a medical decision in 13 cases. MS was the main or the only factor of the decision in 11 of these latter cases: 5 of them were experiencing a relapse, 3 were receiving azathioprine, and although MS was not progressive, therapeutic abortion was undertaken in 3 other cases. Other causes of premature termination of pregnancy were ectopic and molar pregnancies (one each). The average follow up after termination of the last pregnancy was 100-3 months (range: 6 to 264).

The crude relapse-rate during follow up in the pregnancy group as a whole was 0.59 (SEM 0.07). One hundred and eighty seven relapses occurred during follow up in this group; 25 of them occurred during pregnancy and 19 during postpartum periods; only one woman in the pregnancy group had no relapse during follow up. Thirteen patients had all relapses outside the pregnancy and postpartum periods. Nineteen relapses occurred during the 32 full-term pregnancies, and 17 during the corresponding postpartum periods; thus 53% of full-term pregnancies were followed by a relapse. Six relapses occurred during prematurely terminated pregnancies and 2 during the corresponding postpartum periods.

The figure shows relapse-rates for different periods of follow up for full-term pregnancies. Relapse-rate was 0.51 (SEM 0.08) during baseline; during the 3 trimesters of pregnancy, relapse-rates were quite similar [0.61 SEM 0.27 to 0.84 SEM 0.32] and not dif-

Figure: Relapse rate in women with pregnancies during follow up.
frent from baseline. During the first 3 months following delivery of full-term pregnancies, there was a sharp increase of relapse-rate (1.62 SEM 0.38) significantly different from baseline (p = 0.05). During the following 3 months, relapse-rate was again not different from baseline (0.46 SEM 0.26 NS). During prematurely-terminated pregnancies, relapse-rate was excessively high (2.10 SEM 0.70) and significantly different from the baseline value (p = 0.05).

The severity of relapses was variable. Relapses occurring during pregnancy were generally mild or moderate (24/25) and left no or minimal residual disability (only 4 patients had an increase in EDSS of 1 point or more). One patient had a severe multifocal relapse (EDSS: 6) during the first trimester of pregnancy, which cleared completely without steroids. During the postpartum period, relapses tended to be more severe and 8 (out of 19) resulted in an increment of EDSS of 1 point or more. One patient had onset of severe bilateral optic neuritis and complete paraplegia 24 hours after delivery, with slow improvement and marked residual disability (EDSS: 4-0); five years later she had another pregnancy, and again experienced early relapse (2 weeks after delivery), which cleared without further sequelae.

Eight patients entered the progressive phase of MS during follow up, an average of 91.1 months after the last pregnancy (range 67 to 127). During follow up, EDSS was unchanged in 13 patients; it deteriorated by one point in 5 patients, by 2 in 7 and by 3 or more points in eight patients.

**Discussion**

We included in this study all MS patients registered at our clinic during a 22 year period; 33% of them were seen within 2 years of onset; the age and disability distribution (data not shown) of this clinic-based population are similar to those of geographically-based populations,12 and we believe that the prospective follow up of this population gives valuable information on the natural history of MS in relation to pregnancy.

When remittent MS patients were used as their own controls, we could confirm the stepwise increase of relapse-rate during postpartum already shown in all earlier retrospective studies and in the only prospective study published to date.14 In that latter study, 6 out of 8 (75%) patients had a relapse during the first 7 weeks of postpartum; the corresponding figure in our study was 43%. The postpartum increase in relapse-rate is apparent during the first 3 months after delivery, but during the following 3 months relapse-rate turns back to its baseline value; the transient nature of this phenomenon was evident in some but not all retrospective studies.

Pregnancy itself was analysed in three retrospective studies8,10,13 which showed a reduction of the frequency of relapse during pregnancy either in the first and the second10 or the third trimester.8,13 Calculation of relapse-rate in one of these studies8 was hampered by the arbitrary attribution of undated relapses to the postpartum or pregnancy periods. Although relapse-rates were slightly higher during pregnancy than during baseline in our study, we found no significant differences between any trimester of full-term pregnancy and the baseline non-pregnancy-associated period. This difference with earlier studies may be attributed to the prospective collection of data in our study. This result contrasts with the significant, three-fold, increase of relapse-rate during the postpartum period, and indicates that the magnitude of any difference between pregnancy itself and baseline on this particular point is probably small. In that respect, MS could differ from other putative autoimmune disease in which remission is thought to result from immunosuppression induced by pregnancy.19

Despite the increase of relapse-rate observed during the first 3 months following delivery, the overall relapse-rate during follow up in the pregnancy—group was not significantly different from that of the control groups without pregnancy during follow up. Interestingly, the relapse-rate of patients who had children after MS onset, either during follow up or before that period (control group 1), was lower than the relapse-rate of the patients who had no pregnancy after MS

| Table 2 Comparison of evolutive data between 3 groups of remittent MS patients |
|-----------------------------------|-----------------------------------|-----------------------------------|
|                                   | Women with pregnancies during follow up (Pregnancy group) | Women with pregnancies after MS onset (Control group 1) | Women without pregnancies after MS onset (Control group 2) |
| Relapse RATE/year<sup>4</sup> mean (SEM)<sup>2</sup> | 0.64 (0.13) | 0.55 (0.20) | 0.86 (0.09) | 0.07 |
| Variation of EDSS mean (SEM)** | 1.57 (0.35) | 0.82 (0.50) | 1.32 (0.24) | ns |
| Transition to progressive form (per cent) | 24% | 23% | 17% | ns |

<sup>4</sup> Relapse rate/year: per each woman, ratio of number of relapses over duration of remittent phase (in years).
<sup>2</sup> Adjusted on age at MS onset and on duration of disease at entry in the study.
<sup>**</sup> Adjusted on age at MS onset and on duration of the disease at the end of follow up.
Pregnancy and multiple sclerosis: a longitudinal study of 125 remitted patients

onset. This result could be interpreted as indicating a protective effect due to pregnancy. Although we did not collect data on the reasons that led patients to choose to start a pregnancy, an alternative and more probable explanation would be that only patients with unfrequent relapses chose to have children, thus explaining the apparent differences in relapse-rate between groups. Whether pregnancies can reduce relapse-rate in some MS patients should be assessed in prospective studies in which: 1) patients would receive baseline information before deciding to start a pregnancy; and 2) the determinants of their choice would be collected before onset of pregnancy.

A high relapse rate was observed during prematurely-terminated pregnancies, but no firm conclusion can be drawn from this result. The number of events is small, and in fact termination of pregnancy was decided because of the relapse in some of these patients.

Three retrospective studies evaluated long-term disability in relation to pregnancy: no association was found between the number of children or timing of pregnancies and disability. During the prospective follow-up of our remittent patients, increasing disability was not related to pregnancies, even when prognostic variables such as age at onset and duration of MS were taken into account. In the other study in which these variables were assessed, similar results were found. Few remittent patients become progressive or experience a significant increase in disability during the first years of MS, however.

Obtaining significant prospective data on disability in relation to pregnancy clearly needs very long follow-up periods, which may prove unrealistic.

Our study did not address the mechanisms involved in the postpartum increase of relapse-rate which is the most consistent finding, common to all retrospective and prospective studies. Postpartum period, stressful events and viral infections are the only clear triggers of exacerbations in MS. The non specific activation of the immune system which occurs during viral infection through production of cytokines such as gamma-interferon has been advocated as a pathogenetic mechanism in infection-induced relapses. Although alterations of immune parameters have been described during normal pregnancy their magnitude is low and their clinical significance outside the maintenance of pregnancy itself may be poor.

Hormonal factors affect the course of experimental allergic encephalomyelitis, the animal model of MS. Whether the increase in relapse-rate is related to the drop of female hormones after parturition, to the secretion of neurotransmitters and hormones such as prolactin, or to more subtle interferences between hormonal and immune networks remains to be determined. Future studies on pregnancy and MS might concentrate on the postpartum period.