**CONCISE REPORT**

**Methotrexate treatment in juvenile idiopathic arthritis: when is the right time to stop?**

D Foell, M Frosch, A Schulze zur Wiesch, T Vogl, C Sorg, J Roth

**Objectives:** To investigate whether prolonged methotrexate (MTX) treatment after induction of remission influences the subsequent duration of remission in patients with juvenile idiopathic arthritis (JIA), and to analyse the usefulness of myeloid related proteins 8 and 14 (MRP8/MRP14) as predictive markers for the stability of remission at the time when MTX is withdrawn.

**Methods:** Twenty-five patients with oligoarticular and polyarticular JIA who received MTX to induce remission were followed up. MTX treatment was stopped after a mean of 3.8 months (group 1) or 12.6 months (group 2) after remission was documented. Differences in the number of relapses between these groups were looked for. Additionally, MRP8/MRP14 were analysed by ELISA in 22 patients.

**Results:** No difference was found in the number of relapses between patients with prolonged or early discontinued MTX treatment. Patients who were in stable remission had significantly lower MRP levels when MTX was discontinued than patients with relapses. With a cut off point for MRP8/MRP14 at 250 ng/ml, sensitivity and specificity were 100% and 70%, respectively.

**Conclusion:** Longer duration of MTX treatment after induction of remission does not generally improve the status of remission in patients with JIA. Residual synovial inflammation seems to influence the rate of relapses after discontinuation of MTX treatment. MRP8/MRP14 indicate residual activity even in the absence of other laboratory or clinical signs of continuing inflammation. Normal serum concentrations of MRP8/MRP14 in clinical inactive arthritis may help to identify patients in whom MTX can be safely withdrawn after remission is achieved.

**Patients and Methods**

**Patients**

We performed a prospective study including 25 children who fulfilled the criteria for JIA with an oligoarticular or polyarticular course of the disease. At study entry all patients had clinical symptoms of active arthritis, as defined previously. All patients received MTX (10 mg/m² body surface area) to induce remission. MTX treatment was stopped earlier than six months (group 1, 15 patients) or later than six months (group 2, 10 patients) after remission according to American College of Rheumatology (ACR) criteria had been documented. Patients were followed up over 12–36 months at intervals of 12 weeks after MTX was stopped. Table 1 summarises the characteristics of patients.

**Documentation of clinical parameters**

Disease activity was documented by the doctor's global assessment and patient's/parental assessment of disease activity, determination of functional ability, and the number of active joints (joint swelling or limitation of movement, with either pain on movement or tenderness). Patients were categorised as having active disease, or were considered to be in remission based on the ACR criteria for at least three consecutive months, including duration of morning stiffness not exceeding 15 minutes, no fatigue, no active arthritis, and an erythrocyte sedimentation rate (ESR) <20 mm/1st h.

**Abbreviations:** ACR, American College of Rheumatology; CRP, C reactive protein; ELISA, enzyme linked immunosorbent assay; ESR, erythrocyte sedimentation rate; JIA, juvenile idiopathic arthritis; MRP, myeloid related protein; MTX, methotrexate
Laboratory examinations

Serum concentrations of MRP8/MRP14 were determined by a sandwich enzyme linked immunosorbent assay (ELISA) following the system established in our laboratory. 11 MRP8/MRP14 serum levels of 22 patients were analysed in active disease before starting MTX, and also in remission at the time when treatment with MTX was stopped. In the remaining three patients no serum was obtained at this time. ESR (Westergren method) and C reactive protein (CRP; remaining three patients no serum was obtained at this time when treatment with MTX was stopped. In the disease before starting MTX, and also in remission at the duration of remission in groups of patients. Analyses were performed to study the rate of relapses and for tested variables. In addition, survival and log rank sensitivity, specificity, and likelihood ratio over a broad range whole follow up period showed no significant difference for 1, compared with 60% in group 2. A log rank analysis for the year after stopping MTX, the flare up rate was 33% in group (60%) relapsed in group 2 (late withdrawal). Within the first relapse in group 1 (early withdrawal), while six patients discontinued MTX treatment. Seven patients (47%) had a relapse of JIA within two years after treatment with MTX was stopped. There was no difference between patients with prolonged or early relapses

**Table 1** Characteristics of 25 patients with JIA

<table>
<thead>
<tr>
<th>Discontinuation of MTX</th>
<th>Groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Early</td>
</tr>
<tr>
<td>Number of patients</td>
<td>15</td>
</tr>
<tr>
<td>Diagnosis (oligoarticular/polyarticular JIA)</td>
<td>7/8</td>
</tr>
<tr>
<td>Sex (female/male)</td>
<td>13/2</td>
</tr>
<tr>
<td>Age (years), mean (range)</td>
<td>6.6 (3–12)</td>
</tr>
<tr>
<td>MTX (mg/m²), mean (range)</td>
<td>9.9 (5–15)</td>
</tr>
<tr>
<td>MRP8/MRP14 serum levels</td>
<td>3.8 (0–6)</td>
</tr>
<tr>
<td>Duration of remission (months), mean (range)</td>
<td>14.9 (3–36)</td>
</tr>
<tr>
<td>MRP8/MRP14 active (ng/ml), mean (SEM)</td>
<td>2970 (1070)</td>
</tr>
<tr>
<td>CRP active (mg/l), mean (SEM)</td>
<td>1 (1)</td>
</tr>
<tr>
<td>CRP active (mg/ml), mean (SEM)</td>
<td>26.5 (4.5)</td>
</tr>
<tr>
<td>CRP active (mg/l), mean (SEM)</td>
<td>9.3 (1.1)</td>
</tr>
<tr>
<td>Active joints (maximally active), mean (range)</td>
<td>3.1 (1–8)</td>
</tr>
</tbody>
</table>

*Continuation of treatment with MTX after diagnosis of remission was documented.

**RESULTS**

Relapses after discontinuation of MTX treatment

Thirteen (52%) patients had a relapse of JIA within two years after treatment with MTX was stopped. There was no difference between patients with oligoarticular or polyarticular JIA.

Influence of the time of MTX withdrawal on the rate of relapses

We found no difference in the duration of remission and the number of relapses between patients with prolonged or early discontinued MTX treatment. Seven patients (47%) had a relapse in group 1 (early withdrawal), while six patients (60%) relapsed in group 2 (late withdrawal). Within the first year after stopping MTX, the flare up rate was 33% in group 1, compared with 60% in group 2. A log rank analysis for the whole follow up period showed no significant difference for

**Predictive value of MRP levels for the risk of relapse**

Patients who were in stable remission for at least one year after MTX was stopped had significantly lower MRP levels when MTX was discontinued than patients who had a relapse within one year (mean (SEM) 210 (30) ng/ml v 590 (90) ng/ml; **p**, 0.001). (B) MRP8/MRP14 concentrations were analysed in serum from patients in remission, obtained at the time when MTX was withdrawn. Ticks in the graphs for both groups indicate censored data for patients with subsequent follow up of <36 months. A log rank analysis confirmed that the difference between both groups was not significant (p = 0.35).

The duration of remission between the groups (0.87; p = 0.35). Figure 1 shows the survival analysis.

**Figure 1** Duration of remission. A survival analysis showed that patients after early discontinuation of MTX (group 1) had neither earlier flare ups nor more relapses than patients with late discontinuation (group 2). Point 0 indicates the time when MTX was withdrawn. Ticks in the graphs for both groups indicate censored data for patients with the duration of remission between the groups (0.87; p = 0.35).

**Figure 2** Serum levels of MRP8/MRP14. (A) MRP8/MRP14 concentrations were analysed in 22 patients with oligoarticular and polyarticular JIA before and after the start of MTX treatment. There was a significant difference between serum concentrations in active disease before starting treatment with MTX and in inactive disease after successful MTX treatment were 490 (80) ng/ml in group 1 and 2920 (970) ng/ml in group 2. Mean MRP8/MRP14 serum levels in inactive disease after successful MTX treatment were 490 (80) ng/ml in group 1 and 420 (80) ng/ml in group 2.

**Figure 2** Serum levels of MRP8/MRP14. (A) MRP8/MRP14 concentrations were analysed in 22 patients with oligoarticular and polyarticular JIA before and after the start of MTX treatment. There was a significant difference between serum concentrations in active disease before starting treatment with MTX and in inactive disease after successful treatment, respectively. Box plots show median, mean (bold line), 25th and 75th centiles. Error bars indicate 5th and 95th centiles (**p**, 0.001). (B) MRP8/MRP14 concentrations were analysed in serum from patients in remission, obtained at the time when MTX treatment was stopped. Two groups of patients were compared according to their outcome within one year after withdrawal of MTX. Serum levels were significantly higher in patients who had a relapse within the following year than in patients who stayed in stable remission. Data points show individual MRP8/MRP14 serum concentrations. The dashed line indicates a cut off point at 250 ng/ml with a sensitivity to detect risk for a relapse of 100%, while specificity was 70% (**p** = 0.01).
Our data confirm the problem of a high relapse rate after withdrawal of MTX. Taken together, 13 of our 25 patients (52%) had a relapse when MTX treatment was discontinued after clinical remission was achieved. A relapse may occur owing to new developing synovial inflammation. On the other hand, the risk of relapse may be influenced by residual synovial inflammation, even in the absence of clinical signs, when MTX is discontinued. So far, a reliable marker for such non-apparent disease activity in JIA has not been found in a prospective follow up. Our results indicate that prolonged treatment with MTX does not improve the status of remission in general. Thus, prolonged treatment with MTX for all patients, as proposed by some authors, does not generally seem to be appropriate. It is more important to identify individual patients at special risk for relapse in whom withdrawal of MTX might be harmful.

MRP8/MP14 are released by activated phagocytes upon contact with endothelial cells in inflamed tissue. Their serum concentrations correlate well with local inflammation. In our study MRP8/MP14 serum concentrations decreased significantly in response to effective treatment with MTX. However, individual differences in MRP8/MP14 serum levels in inactive JIA point to the fact that a group of patients who are considered to be in remission according to clinical and routinely used laboratory criteria may have undetected local disease activity. These patients would be at special risk for a clinically apparent relapse of JIA as is frequently seen after withdrawal of MTX treatment.

Comparing a subgroup of patients who relapsed with those who did not, we found significantly lower serum levels of MRP8/MP14 for the latter group. Thus, raised MRP8/MP14 serum levels point to patients with clinically undetectable local synovial inflammation for whom prolonged treatment with MTX will be beneficial. Taking into account the high pre-test probability of 50% for a relapse after discontinuation of MTX treatment, a likelihood ratio of 4.6 results in a substantial shift of the post-test probability to about 85% in the presence of high MRP8/MP14 levels.

Although our study group is relatively small these data demonstrate that MRP8/MP14 are markers for clinically non-apparent disease activity. Further prospective studies have to prove that MRP8/MP14 serum levels may be used to guide anti-inflammatory treatment in JIA. So far, the impact of the time point of MTX discontinuation on the risk for relapses has not been studied systematically. We cannot exclude the possibility that reasons related to patient characteristics influenced the decision for stopping MTX earlier or later, and that this might have influenced the subsequent occurrence of flares. Therefore, a controlled randomised study will be necessary to analyse if treatment with MTX can be safely stopped once individual patients have reached a status of remission without signs of residual synovial inflammation.

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REFERENCES
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