Efficacy and safety of methotrexate in articular and cutaneous manifestations of systemic lupus erythematosus

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Abstract

Aim: A prospective open-label study comparing the efficacy and safety of methotrexate (MTX) and chloroquine (CQ) in articular and cutaneous manifestations of systemic lupus erythematosus (SLE).

Methods: Consecutive SLE patients were randomly assigned to either 10 mg MTX weekly or 150 mg CQ daily during 24 weeks. Outcome measures were: numbers of swollen and tender joints, duration of morning stiffness, visual analog scale (VAS) for articular pain, physician global assessment index, patient global assessment index, SLE Disease Activity Index (SLEDAI), disappearance of skin rash and erythrocyte sedimentation rate (ESR).

Results: Forty-one patients consented to participate, 15 were allocated in the MTX group and 26 in the CQ group. Two patients on MTX dropped out due to side-effects and two in the CQ group, one due to side-effects and one due to inefficacy. Baseline demographic, clinical and laboratory parameters of the two groups were nearly identical. In both groups the clinical and laboratory parameters improved significantly over 24 weeks, except the ESR in the MTX group. The results of the outcome measures at the end of the trial did not differ significantly between the two groups, except morning stiffness ($P < 0.05$ in favor of the MTX group) and ESR ($P < 0.01$ in favor of the CQ group). Rise of serum alanine aminotransferase was observed in two cases in the MTX group and in none in the CQ group.

Conclusion: Low-dose MTX appears to be as effective as CQ in patients with articular and cutaneous manifestations of SLE, having an acceptable toxicity profile. Results of this prospective study need to be confirmed in a larger study.

Key words: articular, chloroquine, cutaneous, methotrexate, SLE.

INTRODUCTION

Arthralgia and arthritis occur in approximately 90% and skin and mucous membranes are involved in 80% of cases of systemic lupus erythematosus (SLE). Conventionally, antimalarials are used to control these articular and mucocutaneous manifestations in non-organ threatening SLE. In a recent review of 95 articles on antimalarial treatment in SLE, high levels of evidence were found that both hydroxychloroquine (HCQ) and chloroquine (CQ) prevent lupus flares and increase long-term survival of patients with SLE; moderate evidence was found of protection against irreversible organ damage, thrombosis and bone mass.
loss. Toxicity related to antimalarials is infrequent, mild and usually reversible, with HCQ having a safer profile than CQ, although comparative data come only from one observational study.

In a retrospective study assessing the reasons for discontinuation of antimalarial drugs in SLE patients, 42% discontinued HCQ due to disease remission, 8% due to inefficacy and 29% stopped because of adverse effects, particularly severe anorexia, nausea, headache, dizziness, deafness, visual disturbance and myopathy. In a substantial proportion of cases, antimalarials induce and maintain only partial remission, necessitating addition of low to high doses of corticosteroids and sometimes cytotoxic drugs like azathioprine or cyclophosphamide, and antimalarials are especially inadequate in managing severe lupus, especially lupus nephritis.

The high efficacy/toxicity trade-off and high long-term drug survival rates of methotrexate (MTX) have been documented in rheumatoid arthritis patients in several clinical trials and meta-analyses. Despite existence of some evidence as early as 1965, the interest in the effects of MTX in SLE has grown only in recent years. In 1965, Miescher and Riethmüller used MTX at a dose of 50 mg per week intravenously in 10 SLE patients with arthralgia, skin rash and vascular purpura. They noted a rapid response: the manifestations of SLE diminished within 1 or 2 weeks. Subsequently, MTX was proven to be effective in controlling articular and cutaneous manifestations of SLE in several clinical trials. A few of these trials were retrospective. Other trials were uncontrolled and included either steroid-resistant patients or patients dependent on high doses of steroids.

To our knowledge only two prospective randomized placebo-controlled trials on the effect of MTX in SLE have been published. The first by Carneiro and Sato reported the efficacy of 15–20 mg MTX in 41 patients controlling cutaneous and articular activity of SLE and corticosteroid dose reduction. Side-effects were frequent (gastrointestinal complaints and hepatic enzyme elevations) but only two out of 18 receiving MTX had to stop due to toxicity. A recent trial showed in a 12-month study a steroid-sparing effect of MTX 7.5 increasing to 20 mg per week plus folic acid in 86 randomized patients with moderately active rather than severe lupus. To our knowledge the efficacy of MTX in non-severe SLE has been reported in only two double-blind placebo-controlled trials. Considering the lack of tolerability or efficacy of antimalarials in subsets of patients and the long-term efficacy of MTX in steroid-resistant SLE and its relative safety, despite sometimes considerable side effects, it has become imperative to investigate its status as an alternative for antimalarials in non-organ-threatening articular and cutaneous SLE.

The present study was undertaken to compare the efficacy and toxicity of MTX with those of CQ in muco-cutaneous and articular SLE. The rationale is that in some cases antimalarials have insufficient effect and side-effects and a second affordable effective drug is necessary in developing countries such as Bangladesh. In this study CQ 150 mg base tablet daily was chosen as it is the only antimalarial drug available for SLE in Bangladesh due to its low price.

We chose to keep a fixed weekly dose of 10 mg MTX for the whole duration of the study, in order to assess the outcome at a fixed and low dose. The weight of most of the patients was in the range of 35–45 kg, so the dosage per kg is comparable with that of higher dosages as used in Western countries, where people tend to have higher weight. We decided to study the efficacy at a stable and low MTX dose as many of the patients in daily Bangladeshi practice are using medical treatment without regular clinical follow-ups, due to often long distances to the hospitals and lack of funds to do blood tests. This means that changing dosages, in these often illiterate patients, would be difficult to control. Obviously, during our study all participating patients came for follow-up and laboratory tests at the allocated times.

MATERIALS AND METHODS

This prospective open-label randomized clinical trial was conducted at the lupus clinic of a tertiary care center in Dhaka, the capital of Bangladesh. The study period was June 2001 to November 2002 and included 6-month follow-up. Patients fulfilling American College of Rheumatology (ACR) criteria of SLE and suffering from arthralgia, or arthritis and active skin lesions, were selected for this study. A total of five patients refused to participate after explanation. Exclusion criteria were: involvement of any other systems, pregnancy, lactation, any form of eye problems, history of taking antimalarials within the last 4 months or corticosteroids equivalent to > 20 mg of prednisolone per day, raised serum alanine aminotransferase (ALT), and raised serum creatinine.

All patients gave verbal consent. No written consent was attempted because participants were mostly illiterate. After obtaining informed verbal consent, the
subjects were evaluated clinically and the following laboratory tests were done: routine urinalysis, complete blood counts (CBC), erythrocyte sedimentation rate (ESR), antinuclear antibodies (ANA), anti-double-stranded DNA (anti-dsDNA), serum ALT and creatinine. Serology-like anti-Sjögren syndrome A and B and antiphospholipid antibodies were not done due to financial constraints.

The patients were randomly allocated to either MTX or CQ groups. Randomization was performed following a random number table without considering their presentation. We followed the vertical series of odd and even numbers.

Methotrexate was given at a dose of 10 mg/week and CQ 150 mg per day to the respective groups throughout the study period of 24 weeks. The patients were allowed to continue corticosteroids in a fixed dose that they were taking for at least 2 months before the start of the study and at doses not exceeding 10 mg/day. Increasing the dose was not permitted, also not in any other route like intramuscular, intravenous or intra-articular.

Measures

As this study was restricted to cases with skin and joint lesions, we used the ACR core set of outcome measures which are often used in studies in (rheumatoid) arthritis patients. Outcome measures were: number of swollen and tender joints; duration of morning stiffness; visual analog scale (VAS) for articular pain, physician’s global assessment index, patient’s global assessment index and ESR. As a multi-item disease activity measure we calculated the SLE Disease Activity Index (SLEDAI). For the skin we used the disappearances of skin rash as an outcome measure. Skin lesions of any type such as subacute cutaneous lupus erythematosus (SCLE), chronic discoid lupus erythematosus (CDLE), and butterfly rash were scored as present or absent.

Side-effects were recorded at each visit. CBC, serum ALT and creatinine were measured every 2 weeks during the first month and monthly thereafter as follow-up tests. Ophthalmic evaluation was performed at the end of 6 months.

Statistics

Data were entered into SPSS 10.1 (SPSS Inc., Chicago, IL, US). In case of continuous variables, significance of difference between groups was assessed by Student’s t-test in cases of normally distributed, and Mann–Whitney U-test in non-normally distributed observations. Within-group differences between pre- and post-treatment measures were assessed by Wilcoxon signed rank test. The differences between pre- and post-treatment values of discrete variables were assessed by Fisher’s exact test. The final analysis was not done with intention to treat, and only done in the 37 out of 41 who completed the 24 weeks period.

Ethics

The study was approved by the Ethics Committee of the Bangabandhu Sheikh Mujib Medical University Shahbagh, Dhaka, Bangladesh. The study was performed following the Declaration of Helsinki principles and informed (oral) consent was obtained from all participants before enrolment. As most patients were illiterate, we explained the method orally to the patients and their families and gave extensive possibility for them to ask any questions.

RESULTS

Forty-one subjects consented to participate in the trial. According to the random order 15 were allocated to the MTX and 26 to the CQ group. Two patients in the MTX group were excluded from therapy, one due to central nervous system involvement, manifested by convulsions, and another due to hepatitis. Two patients in the CQ group discontinued therapy, one due to lack of efficacy and one due to psychosis (the psychosis improved after discontinuation of CQ).

Out of 37 completers, 36 were female. Baseline demographic, clinical and laboratory parameters were nearly identical in the two groups (Table 1). All patients were positive for ANA and at the start of the study all had anti-dsDNA antibodies. After 24 weeks in the CQ group in 8/24 cases dsDNA was still positive and in the MTX group 4/15 still had increased dsDNA. Two patients in the MTX group and four in the CQ group were on a stable dosage of prednisolone, with a maximum of 10 mg. As our patients were poor, they could not afford using sunscreen and/or topical steroids.

At baseline the groups did not differ regarding age, sex, illness duration, duration of morning stiffness, joint swelling and pain and VAS pain, patient and physician global assessment (Table 1). The rather high joint tenderness and duration of morning stiffness in both groups may be ascribed to the humid climate, or to sometimes exaggerated estimation by these illiterate patients.
In the CQ group, all outcome measures improved significantly during the study period. In the MTX group, over 24 weeks, SLEDAI and all clinical and laboratory parameters except ESR improved significantly compared to baseline values (Table 2).

The findings in the outcome measures at the end of the trial did not differ significantly between groups, except morning stiffness ($P < 0.05$ in favor of MTX group) and ESR ($P < 0.01$ in favor of CQ group) (Table 2). Improvement of skin rash (near disappearance) was significant in both groups but differences between groups were non-significant (Table 3).

**Side effects**

Anorexia and nausea were common in both groups. Seven patients in the MTX group and four in the CQ group noticed anorexia and nausea. In most cases anorexia and nausea were mild and all subjects were able to continue the drugs and complete the trial. Rise of serum ALT was observed in two cases in the MTX group. Viral markers were found negative in these

### Table 1 Baseline characteristics of methotrexate and chloroquine groups

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Methotrexate ($n = 13$) (mean ± SD)</th>
<th>Chloroquine ($n = 24$) (mean ± SD)</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>24.0 ± 4.5</td>
<td>24.9 ± 7.0</td>
<td>0.299</td>
</tr>
<tr>
<td>Sex (female/male)</td>
<td>13/0</td>
<td>23/1</td>
<td></td>
</tr>
<tr>
<td>Duration of illness (months)</td>
<td>15.4 ± 12.1</td>
<td>12.2 ± 9.5</td>
<td>0.216</td>
</tr>
<tr>
<td>Number of swollen joint</td>
<td>7.8 ± 9.7</td>
<td>2.7 ± 4.6</td>
<td>0.499</td>
</tr>
<tr>
<td>Joint swelling index</td>
<td>11.7 ± 19.4</td>
<td>3.4 ± 5.7</td>
<td>0.475</td>
</tr>
<tr>
<td>Number of tender joint</td>
<td>20.1 ± 10.0</td>
<td>15.2 ± 11.2</td>
<td>0.311</td>
</tr>
<tr>
<td>Joint tenderness index</td>
<td>35.7 ± 21.7</td>
<td>23.0 ± 17.4</td>
<td>0.233</td>
</tr>
<tr>
<td>Morning stiffness (min)</td>
<td>45.0 ± 27.0</td>
<td>29.4 ± 29.0</td>
<td>0.386</td>
</tr>
<tr>
<td>VAS for pain</td>
<td>5.4 ± 2.3</td>
<td>4.5 ± 2.6</td>
<td>0.251</td>
</tr>
<tr>
<td>Physician’s global assessment index</td>
<td>3.4 ± 0.7</td>
<td>3.3 ± 1.0</td>
<td>0.221</td>
</tr>
<tr>
<td>Patient’s global assessment index</td>
<td>3.5 ± 0.7</td>
<td>3.3 ± 0.8</td>
<td>0.226</td>
</tr>
<tr>
<td>Skin rash</td>
<td>6</td>
<td>19</td>
<td>0.499</td>
</tr>
<tr>
<td>ESR (mm/1st h)</td>
<td>73.5 ± 38.7</td>
<td>56.9 ± 30.4</td>
<td>0.455</td>
</tr>
<tr>
<td>ALT (U/L)</td>
<td>35.0 ± 14.0</td>
<td>28.2 ± 10.8</td>
<td>0.212</td>
</tr>
<tr>
<td>Platelet count (per mm$^3$)</td>
<td>240,308 ± 124,751</td>
<td>288,542 ± 77,871</td>
<td>0.412</td>
</tr>
<tr>
<td>Total white blood cell count (cm$^3$)</td>
<td>7,415 ± 2,105</td>
<td>7,242 ± 2,443</td>
<td>0.122</td>
</tr>
<tr>
<td>Hemoglobin (g/dL)</td>
<td>10.3 ± 1.6</td>
<td>10.8 ± 1.5</td>
<td>0.102</td>
</tr>
</tbody>
</table>

ALT, alanine aminotransferase; ESR, erythrocyte sedimentation rate; VAS, visual analog scale.

### Table 2 Changes in the outcome measures in methotrexate ($n = 13$) and chloroquine ($n = 24$) groups

<table>
<thead>
<tr>
<th>Variables</th>
<th>Methotrexate group</th>
<th>Chloroquine group</th>
<th>Inter-group $P$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 week</td>
<td>24 weeks</td>
<td>0 week</td>
</tr>
<tr>
<td></td>
<td>(mean ± SD)</td>
<td>(mean ± SD)</td>
<td>(mean ± SD)</td>
</tr>
<tr>
<td>Number of swollen joint</td>
<td>7.77 ± 9.68</td>
<td>0.77 ± 1.74</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Joint swelling index</td>
<td>11.7 ± 19.4</td>
<td>1.4 ± 3.1</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Number of tender joint</td>
<td>20.1 ± 10.0</td>
<td>3.3 ± 5.3</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Joint tenderness index</td>
<td>35.7 ± 21.7</td>
<td>4.5 ± 9.1</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Morning stiffness (minute)</td>
<td>45.0 ± 27.0</td>
<td>7.7 ± 14.8</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>VAS for pain</td>
<td>5.4 ± 2.3</td>
<td>1.4 ± 2.1</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Physician’s global assessment index</td>
<td>3.4 ± 0.7</td>
<td>1.5 ± 1.1</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Patient assessment index</td>
<td>3.5 ± 0.7</td>
<td>1.6 ± 1.2</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>ESR (mm at 1st h)</td>
<td>73.5 ± 38.7</td>
<td>57.3 ± 29.1</td>
<td>NS</td>
</tr>
<tr>
<td>SLEDAI</td>
<td>12.5 ± 1.2</td>
<td>2.8 ± 2.4</td>
<td>&lt; 0.01</td>
</tr>
</tbody>
</table>

*Wilcoxon signed-rank test. ESR, erythrocyte sedimentation rate; SLEDAI, Systemic Lupus Erythematosus Disease Activity Index; VAS, visual analog scale.*
cases and the liver functions became normal after dis-
continuation of MTX. The number of adverse events
was significantly higher in the MTX group compared
to the CQ group (Table 4). The ophthalmic evaluation
at the end of 6 months did not reveal any abnormali-
ties in either group of patients. No flares of SLE or
end organ damage were observed during the study.

DISCUSSION

The role of antimalarials in the treatment of cutaneous and articular manifestations of SLE has been well
established. In some retrospective and uncontrolled studies, MTX has been reported to be effective in
controlling steroid-dependent or steroid-resistant articular and cutaneous manifestations of SLE. A few of
these trials were retrospective. Other trials were uncontrolled and included either steroid-
resistant patients or patients dependent on high doses of steroid.

Only two prospective randomized placebo-con-
trolled trials on the efficacy of MTX in SLE have been
published, both showing good effect of MTX on
controlling cutaneous and articular activity of SLE
and corticosteroid dose reduction despite frequent
side-effects (gastrointestinal complaints and hepatic
enzyme elevations). In both studies higher MTX dos-
ages were used than in ours. A recent trial was done
in a more severe patient group and half the patients
had renal and two-thirds cardiovascular, hematologic
and other organ involvements. In either arms of the
study two-thirds or virtually all (placebo arm) of the
patients received HCQ so the results of this study can-
not be compared with ours.

None of the studies addressed the issue of safety and
efficacy of MTX compared to those of antimalarials.
These studies opened our eyes to the use of MTX in
articular and cutaneous SLE, but they did not
answer the question if MTX could be used as an alter-
native for antimalarials.

Our prospective, controlled study was the first one
designed to compare the efficacy and safety of MTX
with CQ in patients with articular and cutaneous lupus not resistant to or dependent on high-dose cor-
ticosteroids. As with the previous studies, we dem-
strated the efficacy of MTX in controlling articular and
mucocutaneous manifestations of SLE. There was a
significant improvement in outcome measures in both
MTX and CQ groups and the two groups did not sig-
nificantly differ at the end of the 24-week trial period
in most outcome measures. This indicates that MTX is
as effective as CQ in articular and mucocutaneous
SLE. However, two observations in our limited trial
need re-investigation. The fall in ESR was not signifi-
cant in the MTX group. This may reflect the well-
known fact that ESR does not correlate with disease
activity in SLE and is mainly an indication of, for
example, infection. Alternatively, antimalarials might
be more effective in controlling some aspects of SLE
than MTX. The adverse reactions, although not severe
enough to warrant withdrawal of the drug, were more
frequent in the MTX group. In rheumatoid arthritis,
MTX is more potent than antimalarials, but in many
cases higher dosages of MTX are needed to get an opti-
mal effect. If we had used a higher dose of MTX, the
result might have been that MTX would appear to be
more effective than CQ. On the other hand, the
patients had a very low weight and the dosage per kg
used in our study is comparable with about 20 mg/
week in a Western population. In any case, further
studies are needed.

We feel that there is a role for MTX in the treatment of
patients with articular and mucocutaneous SLE. In
daily practice a failure of antimalarial drugs often
results in long-term treatment with higher doses of corticosteroids. Our findings and those of others indicate that MTX can be used as a steroid-sparing agent. Cyclophosphamide and azathioprine are also effective and have steroid-sparing potential, but there is concern about the risks of carcinogenicity or irreversible steril-
ity. To date no evidence of carcinogenicity has

### Table 3 Number of subjects with skin rash before and after treatment

<table>
<thead>
<tr>
<th>Group</th>
<th>Total At 0 week</th>
<th>At 24 weeks</th>
<th>P-value</th>
<th>Inter-group P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methotrexate</td>
<td>13</td>
<td>06</td>
<td>0</td>
<td>&lt; 0.001 NS</td>
</tr>
<tr>
<td>Chloroquine</td>
<td>24</td>
<td>19</td>
<td>03</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

*NS, not significant.

### Table 4 Adverse effects of methotrexate and chloroquine after 24 weeks

<table>
<thead>
<tr>
<th>Adverse effects</th>
<th>Methotrexate (n = 13) (%)</th>
<th>Chloroquine (n = 24) (%)</th>
<th>P*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anorexia and nausea</td>
<td>7 (53.8)</td>
<td>4 (16.7)</td>
<td>&lt; 0.01</td>
</tr>
<tr>
<td>Raised ALT</td>
<td>2 (15.4)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>9 (69.2)</td>
<td>4 (16.7)</td>
<td></td>
</tr>
</tbody>
</table>

*Fisher’s exact test. ALT, alanine aminotransferase.
been reported with MTX. So MTX may be preferable to these agents.

We permitted enrolment of patients on stable dosage of maximum 10 mg prednisolone during the last 2 months before enrolment and no addition or change of dose during the trial period was allowed. Only two patients in the MTX group and four in CQ group were using corticosteroids, so no subgroup analysis was done.

Our study had several limitations and flaws that need to be specifically addressed in designing future studies on efficacy of MTX in SLE. The first limitation is that the study was designed as an equivalence trial, trying to demonstrate that MTX is equivalent to CQ. Although generally no differences were found between the efficacy of MTX and CQ, the small sample size in this study may have masked some true differences between the groups. To confidently conclude that there is absolutely no difference between the two treatments, one would need a very large sample size in order to achieve enough power to identify even small differences on all outcome measures. Therefore, the conclusion that MTX is equivalent to CQ needs to be interpreted with some caution. Although we realize that the sample size in this study was quite small, as it is in most studies on this issue, we had to make do with the available number of SLE cases fulfilling the criteria. SLE is a rare disease and studies with larger samples need multi-center collaboration.

Another flaw of our study is that there was no blinding to treatment and this may have biased the outcomes. Owing to lack of financial and logistic support, we could not make a double-blind design. But we do not feel that this has influenced the results, as the majority of the patients were illiterate, so in that way they were more or less blinded. Moreover, this study was done without any funding from industry or government.

We were compelled to use in CQ instead of HCQ, the antimalarial more commonly used in SLE. HCQ is more expensive and not available in Bangladesh and many parts of the world. Moreover, it has been suggested that CQ might be more effective although more toxic than HCQ. Both HCQ and CQ have comparable immunomodulatory and anti-inflammatory effects and are effective in SLE. Doses used for both MTX and CQ may appear subtherapeutic. Although we considered applying higher or flexible dosages like the treatment schemes applied in rheumatoid arthritis used in Western countries, we decided to choose in this study for low and fixed dosages in these patients with a weight of generally 35–45 kg. Additionally, we did not give folic acid as a supplement, which considering the current literature in RA would have been preferable. Despite this, we observed few side-effects in the MTX-treated patients.

The assessment of the skin lesions was rather crude – either presence or absence of skin lesions. We did not allow changes in corticosteroid dosages, so possible changes can be ascribed only to MTX or CQ treatments.

A follow-up period of 6 months, as it was applied in most previous studies with MTX in SLE, does not establish sustained efficacy, so in the future longer follow-up is needed.

CONCLUSIONS

As a result of our study we conclude that both drugs are useful in SLE patients with skin and articular complaints and as both drugs are cheap and affordable in our country, we advise its use for the benefit of our patients.

Long-term multi-center double-blind collaborative studies, including large samples, will further clarify the relative safety, efficacy and cost-effectiveness of MTX in the treatment of articular and cutaneous lupus as an alternative to the antimalarials.

ACKNOWLEDGMENTS

We would like to thank all the patients who participated for voluntarily giving up their time to be involved in this study.

CONFLICTS OF INTEREST

The authors have no conflicts of interest.

REFERENCES


