Efficacy of Leflunomide 100 mg Weekly Compared to Low Dose Methotrexate in Patients With Active Rheumatoid Arthritis. Double Blind, Randomized Clinical Trial

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OBJECTIVE: To determine the clinical efficacy and safety of Leflunomide (LFN) 100 mg/week compared to low dose Methotrexate (MTX) 10 mg/week in a double-blind, randomized, controlled trial with 52 weeks of follow up in Rheumatoid Arthritis (RA) patients.

PATIENTS AND METHODS: Patients who met ARC1987 criteria for RA were included. All patients had medical records, including laboratory tests and hand X-rays. Clinical evaluations for improvement and ACR and EULAR response criteria were performed. Statistical analysis for independent’s samples between both groups defined a P value of ≤.05. Safety was evaluated by comparing the proportion of adverse events (AE) registered.

RESULTS: Of the 90 patients screened, five were withdrawn and the remaining 85 patients were randomized: 43 LFN and 42 MTX. Sixty-three patients completed the study, 72% in the LFN group and 74.4% in the MTX group. ACR20 improvement criteria were achieved by LFN group in 90.3%, and in MTX 78.1% (P=.14) at week 52. EULAR improvement criteria applied at the end point showed a DAS28 score for the LFN group of 3.45, and for the MTX group was 3.67 (P=.43). Total withdrawals including loss during follow up, AE and lack of efficacy for each group was 12 patients in the LFN group, and 10 patients in the MTX group. Regarding safety, no serious AE of a life threatening nature were reported.

CONCLUSIONS: These outcomes confirm that LFN 100 mg/week offers an adequate and sustained improvement effect on the clinical manifestations of RA, similar to low dose treatment with MTX 10 mg/every week after 52 weeks of follow up; it may be a good therapeutical option alone or in combination with other anti-rheumatic drugs.

Eficacia de leflunomida 100 mg semanales comparado con dosis bajas de metotrexate en pacientes con artritis reumatoide activa. Estudio clínico doble ciego aleatorizado


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Introduction

Leflunomide (LFN) is a non-biological disease-modifying antirheumatic drug (DMARD), an inhibitor of purine synthesis, which is indicated for the treatment of rheumatoid arthritis (RA). There are several published clinical studies that have demonstrated its benefit and safety, being considered equivalent to treatment with sulfasalazine (SFA) or methotrexate (MTX).1-3

One problem that prevails in the treatment of RA is the compliance (respecting prescription doses) and adherence (to maintain the treatment for a long period of time) to DMARD treatment, a difficult situation to achieve due to multiple factors such as are polypharmacy, adverse drug events and the high cost of treatment, especially in those patients who lack social security coverage, all of which makes it difficult to obtain good long-term clinical outcomes in routine clinical practice.

Seeking treatment alternatives that benefit the compliance and adherence of the treatments, as well as maintains the effectiveness of antirheumatic treatment, we developed an open descriptive study using LFN weekly doses of 100 mg in active RA patients followed up to 6 months, who achieved clinical improvement according to criteria of the American College of Rheumatology (ACR), with no evidence of serious adverse events.4

In the present study, we describe the efficacy and safety results of patients treated in a randomized, comparative, double-blind trial of LFN given at a weekly dose of 100 mg, compared with a fixed low dose of MTX 10 mg/week with 52 weeks of follow up.

Patients and Methods

Patient Population

Patients included in the study were adults who met the ACR1987 criteria for classification as active RA. Patients were enrolled from June 2004 to December 2007 from the outpatient clinic of RA. Active RA was defined for those patients who had at least 6 or more swollen (SJ) and painful (PJ) joints, morning stiffness greater than 30 min and erythrocyte sedimentation rate (ESR) of 20 mm/h or greater. Previous treatment with DMARDs should have been suspended at least one month prior to enrollment, and more than 3 months prior for LFN or MTX. Newly diagnosed patients without DMARD treatment were also included.

The use of prednisone or its equivalent was allowed with a regular dose not exceeding 10 mg daily for the shortest possible time. Patients were excluded if a history of high alcohol consumption was present and pregnancy or a possibility thereof. Baseline laboratory studies requested for inclusion were: normal count of white blood cells, hemoglobin concentration greater than 12 g/dl, albumin levels ≥ 3.5 g/dl, normal liver function tests and if female, negative pregnancy test.

Study Protocol

A randomized controlled trial with a 52-week follow up started in 2004 after approval by the local Committee for Research and Ethics with the registration number 11331-1200-2098-UExe 322-2003 at the State and Municipalities Social Security Institute of Mexico (ISSEMyM), conducted under the guidelines of the International Declaration of Helsinki. The process of informed consent was required for all patients and in addition, females and patients of reproductive age were required to show confirmation of not being pregnant and the use of effective birth control during the development of protocol or until the doctor indicated.

All patients underwent complete medical history, physical examination, laboratory tests and radiographs of hands and feet, the latter for purposes of diagnostic classification. Clinimetric determinations were recorded, which included: 28 joint count (tender and swollen), patient (PGA) and physician (MDGA) global assessment on a visual analog scale (VAS 0–100 mm), patient pain score (VAS 0–100 mm), a validated functional physical limitation questionnaire for Spanish-speaking patients (HAQ-Di in Spanish);5 baseline laboratory studies required were erythrocyte sedimentation rate (ESR, Westergren), C-reactive protein, blood count and liver function tests. Clinical and laboratory tests were performed at the start of patient enrollment and monthly for 3 months, followed by visits every 2 months to complete the 52-week follow up. The X-ray studies of hands and feet were made only at the beginning of the protocol. All laboratory tests and imaging were performed at Toluca’s ISSEMyM Medical Center, under standardized techniques validated according to international protocols of good clinical laboratory practice.

The primary study objective was to evaluate the clinical improvement of the disease according to ACR improvement criteria, with 20% improvement in swollen and tender joints and at least one of the following to determine ACR improvement: pain, global assessment of disease by the patient and the physician, HAQ-Di and acute phase reactants. Also included were additional ACR 50 and 705 results, DAS 28, the criteria for disease activity and improvement of the European League Against Rheumatism (EULAR) at each visit and at the end of the study, EULAR referral criteria and recording of treatment discontinuation due to adverse events.

The criteria for discontinuation of patients in the study were applied to all those who did not achieve ACR 20 improvement at
Patients were randomized into 2 blocks using a table of random numbers, without the intervention of the research group (1:1): the target for the LFN group and a control group of MTX. For the target LFN group, a loading dose of 100 mg/day for 3 consecutive days was given, based on the average half-life of the drug, and administered at a weekly dose of 100 mg. For the MTX group, a fixed low dose of 10 mg weekly was administered; for both groups, placebos were administered in numerical form in an equivalent manner to achieve the blinding of patients and medical researchers.

Statistical Analysis

The primary objective of the study was to compare the efficacy and safety of a weekly dose of 100 LFN mg compared to the effect achieved with low dose of MTX 10 mg weekly. The efficacy was measured by ACR 20 improvement criteria as a study endpoint at 52 weeks of treatment. Variables also included were ACR 50 and 70 improvement, EULAR improvement criteria, and an independent evaluation of the ESR and HAQ-Di variables.

Results

Of the 90 patients evaluated for study entry, 5 were excluded, and the 85 remaining were randomized into 2 groups as follows: a group of 43 patients were assigned to LFN and 42 to MTX (Fig. 1). Both groups of patients were assessed at least once during follow up from the baseline visit. The demographics and disease characteristics were similar for both groups (Table 1). Three patients were treated with DMARDs prior to enrollment, 2 for the LFN group. One of them who received LFN 20 mg/day and hydroxychloroquine for 2 months discontinued treatment for 7 months before being randomized to the LFN group. A second patient, with an irregular treatment, took MTX for a month, 3 months after being randomized to the LFN group. Finally, in a patient receiving conventional LFN, diffuse alopecia developed after 2½ months and treatment was suspended; the patient was sent to our hospital and included in pre-randomization, after no treatment was given for 3 months, to the MTX group. Sixty-three patients completed 52 weeks of treatment, 31 in the LFN (72%) and 32 in the MTX group (74.4%). Early discontinuation of patients at week 16 occurred more often in the LFN than in the MTX group (19.4 vs 5%), respectively. At the end of the study, the total of patients who left were 21, either by loss to follow up or adverse events. Twelve cases occurred (27.9%) in the LFN and 10 patients (23.8%) in the
Table 1
Disease Characteristics and Demographic Data.

<table>
<thead>
<tr>
<th></th>
<th>Leflunomide Group±SD</th>
<th>Methotrexate Group±SD</th>
<th>(P^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients (No.)</td>
<td>43</td>
<td>42</td>
<td>–</td>
</tr>
<tr>
<td>Age, years</td>
<td>42.8 (±11.7)</td>
<td>42.1 (±10.8)</td>
<td>.76</td>
</tr>
<tr>
<td>Female, %</td>
<td>88.3</td>
<td>85.7</td>
<td>.56</td>
</tr>
<tr>
<td>Duration of the disease, months</td>
<td>25.2 (±6.8)</td>
<td>20.9 (±3.5)</td>
<td>.57</td>
</tr>
<tr>
<td>Tender joint count, 0–28</td>
<td>11.1 (±5.1)</td>
<td>11.5 (±6.1)</td>
<td>.74</td>
</tr>
<tr>
<td>Swollen joint count, 0–28</td>
<td>8.9 (±4.8)</td>
<td>7.5 (±4.9)</td>
<td>.17</td>
</tr>
<tr>
<td>Global disease score by the patient (activity), 0–100 mm, VAS &amp;</td>
<td>43.1 (±15.1)</td>
<td>44.5 (±15.1)</td>
<td>.67</td>
</tr>
<tr>
<td>HAQ-Di</td>
<td>0.96 (±0.09)</td>
<td>0.83 (±0.07)</td>
<td>.27</td>
</tr>
<tr>
<td>DAS 28</td>
<td>5.8 (±0.96)</td>
<td>5.6 (±0.88)</td>
<td>.24</td>
</tr>
<tr>
<td>Erythrosedimentation rate, mm/h</td>
<td>41 (±95.3)</td>
<td>39 (±90.7)</td>
<td>.53</td>
</tr>
<tr>
<td>Prior DMARD treatment</td>
<td>2 (4.6%)</td>
<td>1 (2.3%)</td>
<td>.10</td>
</tr>
</tbody>
</table>

SD: standard deviation; VAS: visual analog scale; DMARD: disease modifying antirheumatic drugs.

\(P^a\): no statistically significant differences were seen between groups; \(P\leq .05\).

MTX group. Discontinuation due to lack of efficacy was found in 2 patients in the LFN group (5.2%) and in 4 cases with MTX (12.1%) (Fig. 1).

The ACR improvement criteria were assessed at weeks 8, 24, and 52. In patients assigned to LFN, 28 (80%) achieved ACR 20 at week 24 and in 29 cases (93.5%) at week 52. For the MTX group the results showed that 30 patients (83%) achieved ACR 20 at week 24 and 25 (78.1%) at week 52; comparing the two groups, there was no statistically significant difference (Fig. 2). Evaluating the results of the study end point for ACR 50 and ACR 70 we found no significant differences by comparing the groups for these variables.

The independent variables were evaluated and the results of the HAQ-Di at baseline scored 0.96 for the LFN group and 0.83 for MTX (\(P=27\)). The final evaluation of study data showed a score of 0.23 for LFN and 0.39 for MTX, with a reduction of 0.7 and 0.43, respectively for each study group, with a marginal difference when evaluating this data (\(P=0.05\)) in the LFN group.

EULAR criteria for improvement and remission were evaluated at week 52 of the study. Initial DAS 28 results of the LFN group were 5.83 and 3.45 at 52 weeks (2.38 reduction points). For the MTX group a baseline score of 5.60 was seen, 3.67 at study end, with a net reduction of 1.93 points. There were also no statistically significant differences when comparing results between the two groups (\(P=43\)). The standard cutoffs to define improvement in EULAR DAS 28 were as follows: <3.2 points=good response, from 3.2 to 5.1 moderate response >5.1 points no response (Fig. 4).

Applying the EULAR remission criteria (<2.6 points), 7 patients of the LFN group and 6 of the MTX group achieved remission.

Fig. 2. Percentage of patients reaching the ACR 20 response criteria at 24 and 52 weeks. There were no statistically significant differences between groups.

Fig. 3. Percentage of patients who achieved ACR improvement. No statistically significant difference was seen when comparing the two groups.

Fig. 4. This graph shows the results and final response in both groups at week 52 of the study. LFN group presented a good response in 51.5% compared to 37.5% of the MTX patients. Applying EULAR remission criteria (<2.6 points), 7 patients of the LFN group and 6 of the MTX group achieved remission.
Three cases of hypertension were detected, 2 in the MTX group and 1 in the LFN group. This study included 16 patients, 8 in the LFN group and 8 in the MTX group. Table 2 presents the Infectious Diseases Registered.

<table>
<thead>
<tr>
<th>Disease</th>
<th>LFN=43</th>
<th>MTX=42</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper respiratory tract infections</td>
<td>6 (13.9%)</td>
<td>12 (28.5%)</td>
</tr>
<tr>
<td>Urinary infection</td>
<td>3 (6.9%)</td>
<td>3 (7.1%)</td>
</tr>
<tr>
<td>Gastroenteritis</td>
<td>1 (2.3%)</td>
<td>1 (2.4%)</td>
</tr>
<tr>
<td>Herpes zoster</td>
<td>0 (0%)</td>
<td>1 (2.4%)</td>
</tr>
<tr>
<td>Vulvovaginitis</td>
<td>1 (2.3%)</td>
<td>3 (7.1%)</td>
</tr>
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</table>

A greater number of upper respiratory tract infections were seen in the MTX than in the LFN group.

**Safety**

Serious adverse events were considered by investigators in 9 cases, 2 dermatological reactions occurred in patients in the LFN group, one of them developing severe rash, another erythema multiforme on the trunk. Six patients had elevated liver enzymes 2.5 times above the normal range, four of them received LFN and 2 belonged to the MTX group and they were all withdrawn from the study. (Fig. 1).

Less important events recorded for both groups included vasculitis, pruritus, alopecia, and headache. Presence of infections was observed in both groups, with a slightly higher percentage in patients with MTX treatment (Table 2). Gastrointestinal adverse events (GI) are described in Table 3, where episodes of diarrhea were more often present in the LFN group. The data recorded regarding alterations in liver function tests were as follows: 7 and 17 cases had elevated liver enzymes in the LFN and MTX groups. Four (9.3%) in the LFN group remained >2.5 times the normal range, so patients were withdrawn from the study. In 3 other patients, despite the high values referred to above, these returned to normal levels during the study. For the MTX group, 2 patients had elevations for which they were eliminated, another 7 returned to normal baseline levels without recurrence at study end; there were 2 cases of leukopenia, 2 with anemia and one with thrombocytopenia. There were no adverse events that would jeopardize the lives of patients in any of the 2 groups.

**Discussion**

In daily practice, rheumatologists have a need for RA treatment regimens that are effective and safe, in addition to being flexible in their administration, in order to maintain adherence and compliance to treatment and thus achieve the goals and objectives of clinical improvement or remission of disease.

LFN is a non-biological DMARDs belonging to the isoxazole class; after administration it is rapidly converted to its active metabolite A77 1726; this metabolite induces its therapeutic effect by inhibiting the enzyme dihydrorotate dehydrogenase. This is an important key enzyme in pyrimidine de novo production in T lymphocytes. This molecule has a long plasma life of about 2 weeks (14–18 days). Published studies indicate that the ACR 20 improvement criteria in patients with RA treated with MTX monotherapy ranges from 40% to 60% at 6 and 12 months of follow up. On the other hand, it is well known that treatment of RA patients at doses of 20 LFN mg/day has shown benefit in clinical response similar to MTX and other DMARDs as SSZ. LFN is a non-biological DMARDs belonging to the isoxazole class; after administration it is rapidly converted to its active metabolite A77 1726; this metabolite induces its therapeutic effect by inhibiting the enzyme dihydrorotate dehydrogenase. This is an important key enzyme in pyrimidine de novo production in T lymphocytes. This molecule has a long plasma life of about 2 weeks (14–18 days). Published studies indicate that the ACR 20 improvement criteria in patients with RA treated with MTX monotherapy ranges from 40% to 60% at 6 and 12 months of follow up. On the other hand, it is well known that treatment of RA patients at doses of 20 LFN mg/day has shown benefit in clinical response similar to MTX and other DMARDs as SSZ. Published studies indicate that the ACR 20 improvement criteria in patients with RA treated with MTX monotherapy ranges from 40% to 60% at 6 and 12 months of follow up. On the other hand, it is well known that treatment of RA patients at doses of 20 LFN mg/day has shown benefit in clinical response similar to MTX and other DMARDs as SSZ.

**Table 2**

<table>
<thead>
<tr>
<th>Disease</th>
<th>LFN=43</th>
<th>MTX=42</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastritis</td>
<td>12 (27.9%)</td>
<td>11 (26.1%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>9 (20.9%)</td>
<td>1 (2.3%)</td>
</tr>
<tr>
<td>Abdominal Distension</td>
<td>2 (4.6%)</td>
<td>6 (14.2%)</td>
</tr>
<tr>
<td>Nausea</td>
<td>3 (6.9%)</td>
<td>6 (14.2%)</td>
</tr>
<tr>
<td>Other</td>
<td>2 (4.6%)</td>
<td>1 (2.3%)</td>
</tr>
</tbody>
</table>

Gastrointestinal events such as gastritis and diarrhea were more frequently reported in the LFN than in the MTX group, as seen in the literature; however, abdominal distension was more common in the MTX group.

**Fig. 5.** The behavior of the average serum levels of liver enzymes for each group, indicating that over time the LFN group had a tendency to maintain optimal levels.
In addition, our group previously performed an open 6 month clinical trial at LFN weekly dose of 100 mg in patients with active RA. Fifty patients were enrolled in the study, starting treatment with a loading dose of 100 mg/day for 3 days followed by a weekly dose of 100 LFN mg for a period of 6 months. After 12 weeks, 75% of patients had achieved ACR 20 improvement response and 58% achieved an ACR 50. At study end, 74% achieved ACR 20, 64% of patients achieved ACR 50 and ACR 70 28% improvements.

Adverse events reported in the study ranged from 2% to 16%, which included headache, rash, hair loss, elevated liver enzymes and diarrhea. It was concluded that clinical benefit in response to a weekly regimen of 100 mg of LFN is associated with minor adverse events already reported previously.

The results obtained in this study show that both drugs, at doses lower than those recommended, help compliance and adherence to treatment in a very acceptable percentage, emphasizing that the low dose of MTX of 10 mg/week is only presently recommended at baseline, increasing it if tolerated quickly and in a stepwise fashion. We observed that at week 52, retention of patients was 31 patients (72%) and 32 cases (76%) for the LFN and MTX groups, respectively, with an overall retention of 74%, a situation that differs from reports by other authors, which present more than a 50% loss in studies to LFN at a standard dose; a similar number is reported in patients with long-term treatment with MTX.

The results of ACR improvement, HAQ-DI and ESR did not differ between groups, stressing that the dose of MTX used is currently considered suboptimal and not comparable for assessment of the efficacy of MTX in this study, as the current recommendations of EULAR point out, where a rapid increase up to 20 or 25 mg/week is indicated in order to reduce clinical activity. Of patients who completed the study in the LFN group, 28 achieved an ACR 20 response (90.3%) at week 52 (Fig. 2); however, applying the calculation of patients intended to treat (ITT), the ACR 20 response was 67.4%. Two patients were eliminated from the LFN group for not achieving ACR 20 improvement, compared to MTX where 4 cases did not achieve it.

Regarding adverse events, those seen in the LFN group were similar to those reported in the literature, affecting the skin with erythematous urticaria, alopecia and diarrhea. Liver toxicity was apparently lower in the LFN group, and only one patient remained with persistent enzyme elevation; in this case we ruled out viral hepatitis, and only found fatty liver by conventional ultrasound. The few non-serious adverse events identified were probably related to the low dose of LFN employed.

We conclude that the weekly dose of 100 mg of LFN provides an adequate and sustained response in patients who respond to this drug, allowing for greater adhesion and compliance than reported in the literature for conventional treatment, despite apparently showing fewer reported adverse events compared with the recommended standard dose. It currently constitutes the loading dose in common practice and is not commonly used as described here; therefore, unfortunately some countries have recalled tablets with LFN 100 mg. This scheme also opens the possibility of its use as monotherapy or in combination with other DMARDs, including MTX as an attractive option avoiding polypharmacy. Moreover, the weekly dose of 100LFN mg/week represents a savings for patients, using a lower dose of the drug while maintaining its effectiveness and this situation applies only in countries where there is no health system that allows full coverage of the population.

Finally, we emphasize that the lack of efficacy observed in patients in the MTX group could be a reflection of the low dose used for the purposes of this study, but by no means constitute a recommendation by the authors for use in daily clinical practice.

Studies with larger populations and longer durations will ratify the results we obtained in this study.

Financing

This study or the researchers had no financial relationship with the pharmaceutical industry. The drugs employed were obtained through the institute (ISSEMyM) where research was carried out.

Conflict of Interest

The authors have no conflict of interest to declare.

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References


