Effectiveness and safety of leflunomide for pulmonary and extrapulmonary sarcoidosis

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Effectiveness and safety of leflunomide for pulmonary and extrapulmonary sarcoidosis

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**How does this advance the field?**

The use of steroid-sparing medications in sarcoidosis is widely recommended but the choice of which one is based on very little published data. The therapeutic effectiveness of leflunomide and its capacity to facilitate reduction of steroid dose are currently not widely accepted. The experience reported here is the largest description of the clinical effect of a steroid-sparing medication for sarcoidosis and it validates a role for leflunomide as a useful steroid-sparing option.

**What are the clinical implications?**

Our experience demonstrates that leflunomide is a viable option for pulmonary and extrapulmonary sarcoidosis. It is useful for patients who have steroid side effects, who have not tolerated methotrexate, or who have suboptimal responses to other medications. Periodic monitoring for side-effects is mandatory.
Abstract

Background:

Leflunomide has been reported as an alternative therapy in sarcoidosis. However, the published data are limited.

Methods:

We performed a retrospective chart review of the tolerance and effects leflunomide therapy in patients with sarcoidosis.

Results:

Seventy-six patients were included. The most common reasons for initiation were progression of disease or failure of other immunomodulator therapy. Side effects attributable to leflunomide were noted in 34% of subjects, prompting discontinuation in 17%. The lungs were a target of therapy in 33 (44%) and extrapulmonary organs were a target in 45 (59%). The mean change in forced vital capacity in the six months prior to leflunomide was -0.1 ± 0.3 L, and it was +0.09 ± 0.3 L in the following six months (p=0.01). For extra-pulmonary target organ response, 51% had a good response and 32% partial response. The median corticosteroid dose at initiation was 10 mg (25th, 75th percentile 5, 20 mg) at baseline, and 0 mg (0, 10 mg) at 6 month follow-up (p<0.001).

Conclusions:

Leflunomide is a viable alternative agent for pulmonary and extra-pulmonary sarcoidosis. Leflunomide appears to facilitate reduction of steroid dose and can be considered as
monotherapy or as add-on therapy in cases of progressive disease.
Introduction:  

Treatment options for sarcoidosis are expanding rapidly, as medications approved by regulatory agencies for use in other immune-mediated inflammatory diseases have been adopted by the sarcoidosis community. The data supporting the use of several steroid-sparing agents are mainly limited to small case series, and there is a need for further description of clinical experiences with steroid-sparing therapies.

Leflunomide is an oral anti-lymphocyte agent that has been approved by the Food and Drug Administration (FDA) since 1998 for treatment of rheumatoid arthritis. Its putative mechanism of action involves the inhibition of dihydroorotate dehydrogenase, a key enzyme in the de novo synthesis of uridine monophosphate (dUMP). Since activated, but not memory, T-lymphocytes depend on de novo pyrimidine production for membrane biosynthesis, clonal expansion and terminal differentiation into effector cells, leflunomide represses lymphocyte responses only for actively-stimulated lymphocyte clones\textsuperscript{1,2}. In the absence of sufficient intracellular dUMP, p53-mediated apoptosis is triggered in activated, but not resting, lymphocytes\textsuperscript{3}.

Granulomatous inflammation in sarcoidosis requires antigen-specific CD4\textsuperscript{+} T-lymphocytes\textsuperscript{4, 5}. Inhibition of lymphocyte activation and proliferation is therefore an attractive therapeutic strategy. Baughman et al. previously reported a favorable experience in a single-center retrospective review of leflunomide for 32 patients with failure of or toxicity from methotrexate\textsuperscript{6}. We have been using leflunomide for sarcoidosis since 2004 for pulmonary and extrapulmonary
manifestations. We conducted a retrospective chart review to assess the effectiveness of leflunomide in our population as well as to report our experiences with tolerance and toxicity.

**Materials and Methods:**

We identified all sarcoidosis patients for whom leflunomide was prescribed between January 2004 and March 2009 through review of the electronic medical record. All patients met standard criteria for the diagnosis of sarcoidosis\(^5\). Organ involvement was classified according to criteria proposed in the A Case Control Etiologic Study of Sarcoidosis formulation\(^7\). Clinical records were reviewed to determine the target organ(s) precipitating the use of leflunomide. Patients for whom two immunomodulatory drugs (e.g., leflunomide and infliximab) were started at the same time were excluded from the effectiveness analysis, but patients for whom leflunomide was added to a stable dose of other medications were included. This study was approved by the Cleveland Clinic Institutional Review Board under approval number 09-873.

The effects of therapy on extrapulmonary disease were assessed according to the criteria described by Baughman et al\(^6\). The therapeutic assessments were obtained from the chart as documented by the treating physician and were re-reviewed by an investigator. *Complete response* was defined as greater than 90% improvement of the maximal disease involvement; *partial response* required greater than 50% reduction in the maximal disease involvement; patients with less than 50% improvement in disease and/or progressive disease in one or more organs were classified as “*no response*”. The analysis of effectiveness was made at the first visit occurring after six months on therapy, but no later than nine months. Only patients who had
follow-up for at least six months after initiation of leflunomide were included for effectiveness analysis. For toxicity follow-up, we included all data up to the most recent clinic visit.

The effect of leflunomide on lung function was assessed by comparing the change in lung function over the six month period prior to initiation of leflunomide to that occurring at six months after initiation using a paired t-test. Pulmonary function testing included for this analysis had to be performed within 12 weeks after the six month time-point to be included. All spirometry data were obtained at our center using the modified spirometry maneuver described by Stoller et al.

A paired t-test or Mann-Whitney test was used to test the effects of leflunomide on outcomes, according to the distribution of data. SAS 9.1 software was used for statistical analysis.

Results:

We identified 76 patients with a diagnosis of sarcoidosis who received a prescription for leflunomide. The demographic characteristics are described in Table 1: the majority of the patients were females, European-American, and current or former smokers. Seventy (92%) of the patients exhibited lung involvement, mostly commonly Scadding radiographic Stage 2 or 3.

We classified the main reason(s) for starting leflunomide according to the prescribing physician (Table 2). The most common single rationale for starting leflunomide was insufficient response to prior therapy. More than one reason could be present. In 13 (17%) of the patients, there was more than one reason for the use of leflunomide. At the time leflunomide was prescribed, 58
(76%) of patients were receiving oral corticosteroids, with a median prednisone dose of 10 mg (25th, 75th percentile 5, 20 mg) daily. Sixty-five (86%) of the patients had been on other non-steroid immunomodulators, most commonly methotrexate (58 patients, 77%), for a mean period of 23±39 months.

Our standard practice is to initiate leflunomide at 20 mg daily; only three patients were loaded with higher doses initially (100 mg daily for three days). The three patients who were loaded with 100mg did not experience toxicities. The mean duration of leflunomide therapy in our cohort was 16 ± 13 months. All but four patients who remained on leflunomide received 20 mg daily, one patient received 30 mg and the remainder 10 mg. Of the 76 patients, 54 (71%) remained on leflunomide and had at least six months of follow-up at our center. Of the remaining 22 patients, three patients were lost to follow-up after the medication was prescribed, three patients did not start the medication due to insurance difficulties, and the remaining 16 patients discontinued the medication within six months. The reasons for discontinuation included gastrointestinal intolerance (4 patients), other side effects (10 patients), and patient preference (2 patients).

Side effects were common (34% of subjects), but usually minor; 14 patients (20%) eventually stopped leflunomide due to toxicities or side-effects (Table 3). The most common side effects were diarrhea (25%) and elevated liver enzymes (7%). Diarrhea responded to dose reduction to 10 mg daily in four patients when that was tried. None of the patients developed persistent hepatic enzyme derangements or evidence of liver failure. Other possible toxicities noted in our cohort included peripheral neuropathy, arthralgia, blurred vision and hair loss (Table 3). Eight
patients developed symptoms suggestive of lower respiratory tract infection while on leflunomide. The two who were managed at our institution both required hospitalization and had radiographic and clinical evidence of pneumonia. It was not possible to distinguish whether the remaining six patients had bronchitis or pneumonia with the available records. All eight subjects were treated with antibiotic therapy, had resolution of their symptoms and leflunomide was continued without difficulties in six of them.

Of the 54 patients who completed at least 6 months of treatment and had adequate follow-up, 41 patients were on prednisone at the time leflunomide was started. At six months, 13 patients were weaned entirely off systemic corticosteroids. Overall, the median (25th, 75th percentile) prednisone dose at initiation was 10 mg (5, 20 mg) and 0 mg (0, 10 mg) at 6 months follow-up (p<0.001). Thirty-six (87%) of 41 patients who used any corticosteroids during the study period were able to reduce the dose by at least 50%, whereas two subjects (5%) required increased prednisone. Concomitant immunomodulators had been stopped in 16 of the 35 (45%) patients who were using them prior to starting leflunomide therapy.

The lungs were considered to be a target for initiation of leflunomide in 33 (44%) of patients, of whom 24 completed at least six months follow up. We assessed the effect of leflunomide on lung function in this group by comparing the change in forced vital capacity in the six month period prior to initiation of leflunomide to that occurring at six months after initiation (Figure 1). Prior to leflunomide, the mean change in FVC was -0.1 ± 0.3 L; after starting the medication, there was a mean gain of +0.09 ± 0.3L (p<0.01, paired t-test). The effect of leflunomide on change in the FVC slope was not dependent on the initial FVC, radiographic stage, disease duration, age,
gender or smoking status, but the overall numbers are too small to exclude any relationship between these variables and the outcome. The mean decrease of DLCO in the six months prior to leflunomide was $-0.8 \pm 3.1$ mL/min/mm Hg; in the six months after starting leflunomide the mean gain was $+0.6 \pm 2.8$ mL/min/mm Hg ($p=0.16$, Figure 1b). Response rates for FVC and DLCO were nearly identical when the patients were stratified by the reason for starting leflunomide.

Leflunomide was prescribed for treatment of 45 extrapulmonary organ targets in 38 patients; of these, 28 patients with 37 target organs completed at least 6 months treatment with leflunomide. The most frequent extrapulmonary manifestations were cutaneous (32% of the patients assessed for effectiveness), ocular (21%) and sinonasal disease (16%). No patient had concomitant improvement in a target organ and progressive disease in a different organ. Of the 37 target organs, 19 (51%) had a complete response and 12 (32%) a partial response. There was a trend for a better response in subjects on combination methotrexate/leflunomide therapy than for leflunomide monotherapy (Fischer exact test, $p=0.004$) (Table 4).

**Discussion:**

The decision to treat sarcoidosis is based on the clinical phenotype of the disease, its perceived effect on organ function and quality of life, and discussions with the patient\textsuperscript{10,11}. There are no FDA approved drugs for the treatment of sarcoidosis. Most authors recommend corticosteroids as the mainstay of treatment\textsuperscript{5,10,11}, but there is a growing recognition that the chronic use of corticosteroids may be overly burdensome for some patients due to their toxicities\textsuperscript{12,13}. Some data suggest that the use of corticosteroids is associated with impaired quality of life, even when
taking disease severity into account. In chronic sarcoidosis, it may be worthwhile to have risk benefit assessment to prioritize steroid-sparing therapies more heavily in clinical decision-making, though this hypothesis has not been formally tested.

Unfortunately, once the decision to use steroid-sparing therapies is made, there are little data to guide the clinician. Methotrexate is the most studied alternative therapy, and the choice of most sarcoidosis experts. We use methotrexate in our center as the preferred second-line option. Other commonly touted non-biologic therapies include azathioprine, leflunomide, mycophenolate and anti-malarial drugs. However, there are no data comparing any of these options and the scientific data supporting their use in sarcoidosis is generally extremely weak.

For the past several years, we have routinely preferred leflunomide as the third line agent after methotrexate (when biologic therapies are not indicated) in patients with progressive disease and in those with toxicities from the other medications. This preference has been based on our anecdotal experience with these options.

The first successful use of leflunomide for sarcoidosis was reported in 2003 by Majitha et al for sinonasal sarcoidosis. Subsequently, Baughman et al. described their experience in 32 subjects, with a partial or complete response present in 78%. Of note, patients intolerant to MTX were usually successfully treated with leflunomide in that series. We observed a similar pattern: of the 33 patients who were started on leflunomide because of toxicity from other immunomodulatory medications, 20 tolerated the leflunomide well. Of these, 13 of 24 patients who switched from methotrexate because of toxicity tolerated leflunomide. Compared to Baughman’s cohort, our patient group was younger (mean age 50 vs. 40 yrs) and had a higher frequency of pulmonary disease as a treatment target. The incidence of leflunomide toxicity that precipitated discontinuation was higher in our series (18% vs. 9%); this finding may relate to
more aggressive dosing in our population—more than 90% of our patients received 20 mg daily, whereas 56% of the patients in the prior series were treated with 10 mg daily. Other factors that may account for the higher incidence of side-effects in our series include the longer follow-up period, different use of concomitant medications, clinician preferences, or other patient-related factors. It is difficult to compare the response rates between the two studies, but overall they appear to be roughly similar, with complete or good responses noted in 82% of extra-pulmonary organs in our series versus 78% in the prior experience.

The safety of leflunomide has been well studied in rheumatoid arthritis both as monotherapy and in combination with other immunomodulators\textsuperscript{2,17,18,19}. In the rheumatoid arthritis population, 50 to 70% of patients remained on leflunomide at the end of one year\textsuperscript{18,20}; the most common reasons for stopping leflunomide in those studies were side effects (40%), lack of efficacy (33%), or both (26%)\textsuperscript{21}. We had a 60% retention of leflunomide at 12 months in our patients, including 100% retention in three elderly patients aged >65 years. A recent prospective, open-label observational series of 334 subjects treated with leflunomide noted that diarrhea (3.0%), nausea (2.4%), hypertension (1.8%), and headache (1.5%) were the most common toxicities, with serious adverse drug reactions in four patients (1.2%)\textsuperscript{22}. The overall incidence of diarrhea has been reported to be up to 24% in some studies, but only 2.2% required discontinuation of the medication\textsuperscript{2}. We found that side-effects occurred commonly in our patients (34%), with a similar incidence of diarrhea (25%) in our population. When it was attempted, four patients responded very well to dose reduction. Our design biases the results toward reporting toxicity, since the effectiveness analysis included only those patients with six months follow-up but the toxicity could occur at any point during the follow-up period.
The most serious reactions we noted were lower respiratory tract infections and peripheral neuropathy. Review of the eight episodes of lower respiratory tract infection in our population revealed that most of them treated on an outpatient basis except two, who had severe pneumonia, requiring hospitalization and withdrawal of immunosuppressive medications. Of these eight patients, six were on leflunomide and prednisone and two were on the combination of leflunomide and methotrexate. It is possible that some of our subjects actually developed pneumonitis from the medication, a potential toxicity that has been reported in small series in Japan and New Zealand\textsuperscript{18,23,24}. However, six of the eight patients continued on the same immunomodulator including leflunomide with no further evidence of untoward pulmonary events.

New symptoms of length-dependent peripheral neuropathy occurred in two of our subjects, on average four months after starting the medication. Peripheral neuropathy has been reported as a complication of leflunomide\textsuperscript{25}. Our standard approach when patients complain of neuropathic symptoms includes immediately stopping the medication and active removal with cholestyramine. The neuropathic symptoms persisted in both patients, but partially abated after stopping the medication with no evidence of progression. In both cases, the symptoms remain mild. However, this is a potentially very serious complication if it is not recognized and addressed promptly.

Our study was statistically significant for prednisone dose reduction and FVC change after initiation of treatment with leflunomide. Although the mean change in FVC was only 200 mL,
which could be construed as clinically unimportant, the data demonstrate reversal of established declining FVC, which is likely to be relevant in pulmonary sarcoidosis. When comparing the change in FVC between the patient being treated with leflunomide and leflunomide plus methotrexate in outcome there was no statistical difference.

There are several weaknesses inherent in our study design. The data are retrospective and uncontrolled. The instrument used for grading disease response is subjective and has not been validated. In support of the grading system, the clinicians caring for the patients were generally able to reduce the dose of concomitant medications. Also, there was no prospective attempt to comprehensively collect adverse events. However, most of the patients we reviewed followed up closely in our center and routinely notified us of any new symptoms. Monitoring blood tests are almost exclusively sent to us after initiating new medications. Therefore, it is likely that almost all significant toxicities were included in our sample.

**Conclusion:**

Leflunomide is a viable alternative immunosuppressive agent in the treatment of sarcoidosis with benefits for both pulmonary and extra-pulmonary diseases. We noted that both pulmonary and extrapulmonary sarcoidosis responded favorably to leflunomide, contemporaneous with substantial reductions of steroid and non-steroid medications. The frequency of side effects in this cohort suggests that clinicians prescribing leflunomide should monitor patients closely and consider the use of cholestyramine wash-out when significant toxicity occurs. Our data demonstrate that leflunomide can be useful both as a steroid-sparing agent and for patients failing other therapies. There is a need for prospective comparative studies to evaluate the
relative efficacy of leflunomide monotherapy or combination therapy compared with other alternatives.

**Author contributions**

D. Sahoo contributed to the design of the study, data collection, analysis, and drafting of the manuscript.

D. Bandyopadhyay contributed to the data collection, analysis, and drafting of the manuscript.

M. Xu contributed to the design of the study and data analysis.

K. Pearson contributed to the data collection and review of the final manuscript.

J.O. Parambil contributed to clinical characterization of the subjects and contributed to and approved the final manuscript.

C.A. Lazar contributed to clinical characterization of the subjects and contributed to and approved the final manuscript.

J.T. Chapman contributed to clinical characterization of the subjects and contributed to and approved the final manuscript.

D.A. Culver oversaw the design of the study, data analysis, drafting of the final manuscript and approval of the final manuscript.
References:


Table 1: Study population

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>N=76</th>
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</thead>
<tbody>
<tr>
<td>Age at initiation of drug (years)</td>
<td>49 ± 10</td>
</tr>
<tr>
<td>Median duration of sarcoidosis before initiation of treatment (years)</td>
<td>5.0 ± 7.8</td>
</tr>
<tr>
<td>Gender (% female)</td>
<td>60</td>
</tr>
<tr>
<td>Race (%)</td>
<td></td>
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<tr>
<td>European-American</td>
<td>55</td>
</tr>
<tr>
<td>African-American</td>
<td>45</td>
</tr>
<tr>
<td>Tobacco use (%Smoker/Ex-smoker)</td>
<td>53</td>
</tr>
<tr>
<td>CXR stage (%)</td>
<td></td>
</tr>
<tr>
<td>Stage 0</td>
<td>7</td>
</tr>
<tr>
<td>Stage 1</td>
<td>19</td>
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<tr>
<td>Stage 2</td>
<td>35</td>
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<tr>
<td>Stage 3</td>
<td>28</td>
</tr>
<tr>
<td>Stage 4</td>
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</table>

Denotes mean ± standard deviation, unless stated otherwise
Table 2: Reason for initiation of leflunomide

<table>
<thead>
<tr>
<th>Reason for initiation (N=76)</th>
<th>Number (%)</th>
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<tbody>
<tr>
<td>Poor response to prior medication</td>
<td>60 (79%)</td>
</tr>
<tr>
<td>• Pulmonary</td>
<td>33 (44%)</td>
</tr>
<tr>
<td>• Extra- Pulmonary</td>
<td>45 (59%)</td>
</tr>
<tr>
<td>Toxicity from therapy</td>
<td>13 (17%)</td>
</tr>
<tr>
<td>Patient preference to taper steroids</td>
<td>3 (4%)</td>
</tr>
<tr>
<td>More than one of the above reasons</td>
<td>13 (17%)</td>
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Table 3: Side effects of leflunomide

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<th>Side Effect (n = 68*)</th>
<th>Number (%) **</th>
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<tr>
<td>None</td>
<td>45 (66%)</td>
</tr>
<tr>
<td>Diarrhea/nausea/bloating</td>
<td>17 (25%)</td>
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<tr>
<td>Hepatic enzyme elevation</td>
<td>5 (7%)</td>
</tr>
<tr>
<td>Neuropathy</td>
<td>2 (3%)</td>
</tr>
<tr>
<td>Hair loss</td>
<td>2 (3%)</td>
</tr>
<tr>
<td>Visual disturbance</td>
<td>1 (1%)</td>
</tr>
<tr>
<td>Arthralgia</td>
<td>1 (1%)</td>
</tr>
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</table>

*8 patients did not take leflunomide after initial prescription

** Sum is more than 100% since some patients had more than one side effect
Table 4: Target organ and treatment response for extra pulmonary organ at 6-9 months

<table>
<thead>
<tr>
<th></th>
<th>Complete (&gt;90%)</th>
<th>Partial (50-89%)</th>
<th>No response (&lt;49%)</th>
<th>Cannot assess</th>
<th>Total</th>
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<td>5</td>
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<td>5</td>
<td>2</td>
<td>0</td>
<td>1</td>
<td>8</td>
</tr>
<tr>
<td>Sinonasal</td>
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<td>2</td>
<td>0</td>
<td>0</td>
<td>6</td>
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<td>Cardiac</td>
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<td>1</td>
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<td>0</td>
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<td>1</td>
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<td>2</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
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<td>0</td>
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<td>0</td>
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<td>0</td>
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<td>0</td>
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<td>1</td>
</tr>
<tr>
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<td>1</td>
</tr>
<tr>
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<td>0</td>
<td>0</td>
<td>1</td>
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<tr>
<td>Total*</td>
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<td>B</td>
<td>13</td>
<td>3</td>
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A = Leflunomide, B = Leflunomide + Methotrexate
CNS=central nervous system; PNS= peripheral nervous system; MSK=musculoskeletal

* Fischer exact test p=0.004 between A&B
Figure 1: Change in FVC and DLCO from initiation of leflunomide to six-nine month follow-up; rectangles represent inter quartile range, horizontal line within box is the median and dot denotes mean value. Change in FVC was statistically significant (p < 0.01) while DLCO was not (p= 0.16)