Managing fatigue in sarcoidosis – a systematic review of the evidence

Dr Chris Atkins – NIHR Clinical Research Fellow¹
Professor Andrew M Wilson – Professor of Respiratory Medicine¹

¹Norwich Medical School, University of East Anglia, Norwich, Norfolk, UK, NR4 7TJ

Postal Address (for both authors):
Bob Champion Research and Education Building, Norwich Medical School, Norwich, Norfolk, UK, NR4 7UQ.

Contact for correspondence:
Dr Chris Atkins, Research fellow, Bob Champion Research and Education Building, Norwich Medical School, Norwich, Norfolk, UK, NR4 7UQ.
Telephone: 01603 591594
Email: c.atkins@uea.ac.uk
Abstract

Background – Fatigue is a common manifestation of sarcoidosis, often persisting without evidence of disease activity. First-line therapies for sarcoidosis have limited effect on fatigue. This review aimed to assess the treatment options targeting sarcoidosis-associated fatigue.

Search Methods – Medline and Web of Science were searched in November 2015; the bibliographies of these papers, and relevant review papers, were also searched. Studies were included if they reported on the efficacy of interventions (both pharmacological and non-pharmacological) on fatigue scores in sarcoidosis patients.

Results – Eight studies were identified that fulfilled the inclusion criteria. These studies evaluated six different interventions (infliximab, adalimumab, ARA 290, methylphenidate, armodafinil and exercise programmes). Evidence to support a treatment effect of anti-TNF-alpha therapies (adalimumab and infliximab) and neurostimulants (methylphenidate and armodafinil), but within five of the studies the risk of bias was high within most domains and the remaining three studies included only small numbers of participants and were short in duration.

Conclusions – Trial evidence for treating fatigue as a manifestation of sarcoidosis is limited and requires further investigation. Anti-TNF-alpha therapies may be beneficial in patients with organ-threatening disease. Neurostimulants have some trial evidence supporting improvements in fatigue but further investigation is needed before they can be recommended.

Registry number - CRD42015030079

Abstract word count – 199 words

Keywords (MeSH) – Sarcoidosis; Fatigue; Fatigue syndrome, chronic; Antibodies, monoclonal; Exercise therapy; Central nervous system stimulants
Introduction

Fatigue is a common problem in patients with sarcoidosis. Its prevalence varies between studies, from 50%\(^1\) to over 80\% of patients\(^2\), with a significant adverse impact on quality of life\(^3\). The aetiology is poorly understood and likely to be multifactorial, encompassing active inflammation, cytokine release, depression, sleep disturbance and/or small-fibre neuropathy\(^4\). Furthermore, systemic treatments used to treat sarcoidosis can themselves cause fatigue, including corticosteroids \(^5\), \(^6\). Management of this symptom can be challenging for physicians. Chronic fatigue in stable sarcoidosis patients was described as long ago as 1993, with management suggested to include “unremitting sympathy”\(^7\); despite being identified as a feature of sarcoidosis for so long it remains a poorly understood and often forgotten problem.

Identifying and treating any underlying and reversible causes of fatigue should be the first priority for any physician faced with a sarcoidosis patient with significant fatigue, which can include including sleep disordered breathing\(^8\), \(^9\) and periodic limb-movement syndromes\(^9\). Once these have been excluded, strategies to improve fatigue should be considered. The evidence available for treatments of fatigue in sarcoidosis patients is limited. Few trials have specifically investigated treating fatigue, although measurement of fatigue is recommended as an outcome measure for trials involving sarcoidosis patients by the World Association of Sarcoidosis and Other Granulomatous Diseases (WASOG)\(^10\).

Various potential management strategies for sarcoidosis-fatigue have been suggested previously, including both pharmacological and non-pharmacological interventions. Tumour necrosis factor (TNF)-alpha is released by alveolar macrophages and is involved in the initial pathogenesis of sarcoidosis\(^11\), as well as influencing prognosis\(^12\). Anti TNF-alpha therapies have been investigated as a treatment for refractory sarcoidosis; amongst these trials, measurements of fatigue pre- and post-treatment have been taken. ARA 290\(^13\), a novel peptide modelled on erythropoietin
that has anti-inflammatory and tissue protective activities, has also been investigated for its potential effects on symptoms of small-fibre neuropathy, including cognitive failure and fatigue(14).

The use of neurostimulants has shown promise in treating other causes of fatigue. Methylphenidate, and its d-isomer dexamethasone, act by inhibiting dopamine and noradrenaline transporters, elevating dopamine and noradrenaline levels within the brain(15). Methylphenidate is known for its use in attention-deficit hyperactivity disorder (ADHD)(16), but has been trialled for fatigue in other situations (including chemotherapy(17), post-radiotherapy(18), HIV(19), and Parkinson’s disease(20)) with some benefit. Modafinil and its enantiomer armodafinil are neurostimulants that are used for promoting wakefulness in narcolepsy. They have a complex profile of neurochemical effects which are different to those of amphetamines(21).

Finally, physical activity programmes, including pulmonary rehabilitation in chronic obstructive pulmonary disease (COPD) patients, have been shown to have wide reaching benefits, including improving dyspnoea scores, health-related quality of life, anxiety and depression(22). The use of pulmonary rehabilitation in patients with interstitial lung diseases is recommended by guidelines(23, 24), with evidence of benefit in this patient group similar to that seen in COPD patients(25). Recent trials investigating the use of such programmes in sarcoidosis patients and their impact on fatigue.

The objective of this systematic review was to examine the evidence for treatments or management strategies of fatigue in sarcoidosis patients who are experiencing significant fatigue, as well as evaluating the quality of the evidence that presently exists, and to present the results of a qualitative analysis of the available data.

**Methods**

**Publication Search**

This systematic review was registered with PROSPERO (registration number CRD42015030079) and was performed in accordance to the Preferred Reporting Items for Systematic Reviews and Meta-
Analysis (PRISMA) statement(26). The protocol can be accessed on the PROSPERO database using the registration number. The electronic databases Medline (using PubMed) and Web of Science were searched using the following search strategy: “sarcoid” OR “sarcoidosis” OR sarcoidosis [MeSH Terms] OR Sarcoid* (truncation) AND “fatigue” OR “chronic fatigue” OR “chronic fatigue syndrome” OR fatigue [MeSH Terms] OR fatigue syndrome, chronic [MeSH Terms] AND “treatment” OR “management” OR “clinical trial”.

Bibliographies of appropriate papers, including review articles, was undertaken to identify relevant additional sources. The search included all trials published to the end of November 2015. The title and abstract of all papers identified were reviewed for relevance, with irrelevant studies and reports excluded. Remaining papers were reviewed in full.

Selection Criteria

Papers that were considered suitable for inclusion were any studies that evaluated the effect of an intervention or management strategy (pharmacological or non-pharmacological) on fatigue in sarcoidosis patients. All trial designs (case series, case-control, cross-over and parallel-arm randomised controlled trials) were included in the qualitative synthesis with the exception of single case reports, and so trials where no comparator group or comparator intervention were included. This ensured the broadest collection of evidence. The studies must have (1) evaluated sarcoidosis patients exclusively, or if the trial included other diseases then the sarcoidosis cohort had to evaluate sarcoidosis separately in the results, (2) evaluated the efficacy of the intervention on an assessment/measurement score of fatigue, (3) reported quantitative results for reduction in fatigue score between pre- and post-intervention results and (4) be presented in full-text form and in English.
Data Collection

Data extraction was performed independently by the two authors (CA and AMW) using a checklist of data to extract. In addition, the Cochrane Collaboration’s tool for assessing risk of bias was also used to assess the methodological quality of each trial and identify possible sources of bias at a study level. The following data was collected: Main author’s last name, year of publication, study design, number of participants, severity of sarcoidosis in participants by chest X-ray staging (if given), intervention (including dose), duration of intervention, measurement score of fatigue used, change in fatigue outcome, number of participants reaching minimum clinically important difference on outcome score (if given), number of participants completing the intervention and adverse events reported within the trial. The summary measure of interest was mean improvement of fatigue score however a meta-analysis of data was not possible due to the heterogeneity of interventions and study designs included. Therefore a narrative review of the data is presented.

Results

Search results and characteristics of eligible trials

The search strategy identified 150 records through Medline and 126 records through Web of Science. Two further papers of interest was identified through bibliography searches. Titles and abstracts were reviewed for relevance, with studies excluded if they did not include patients with sarcoidosis, were evaluating a potential intervention, or were review articles or case reports. A short list of 29 studies was identified, of which eight met the criteria for inclusion. The flow diagram (PRISMA 2009) of screening articles is shown in figure 1. Only three randomised controlled trials (either parallel arm or cross-over) were identified, the remaining articles consisted of two case-control studies, three case series and one retrospective case review. Three papers evaluated formal
physical activity programs, two evaluated systemic treatment with tumour necrosis factor alpha (TNF-alpha) inhibitors, two evaluated symptom-targeted therapy with neurostimulants (dexamethasone and armodafinil), and one trial looked at a novel molecule aimed at treating small-fibre neuropathy. All but one of the trials used the ‘Fatigue Assessment Scale’ (FAS), a ten-point questionnaire with a maximum score of 50 points. This scale has been validated in sarcoidosis patients(27), has a well-defined cut-off score for fatigue of >21 points(27), and known minimal clinically important differences (MCID) of four points or 10% reduction(28). A total of 185 patients were included across all the trials. The details of the included studies are shown in table 1 (study information, safety and efficacy) and table 2 (risks of bias within studies). In addition, three papers that did not meet the inclusion criteria but provided useful information are discussed.

Systemic therapy: anti-TNF-alpha treatment and anti-inflammatories

Three trials evaluated interventions which have systemic or disease-modifying effects. Two investigated anti-TNF-alpha drugs (adalimumab(29) and infliximab(30)) and one investigated ARA 290(13) Only one trial, investigating ARA 290(13), was a blinded, randomised-controlled trial; neither of the studies investigating anti-TNF-alpha therapies were of a randomised design.

Erckens and colleagues(29) followed 26 sarcoidosis patients with refractory uveitis who had been commenced on Adalimumab (40mg subcutaneously once weekly) over a 12 months treatment period. All patients had previously received prednisolone and methotrexate. Fatigue was measured using the FAS, with 21 patients (80.7%) having a baseline FAS score >21 points. After the 12 month treatment period, there was a mean reduction in FAS score of 2.2 points and an improvement in FAS score in 14 of the 21 patients with a baseline FAS score >21 points; although it is not stated what constituted an improvement in the FAS and whether these patients met the MCID for the FAS score. The risk of bias is high with this trial design, as there is no comparator group to determine if this is a
placebo-effect or natural progression of the disease, however, there is a suggestion that fatigue is improved by anti-TNF-alpha therapy.

In another study from the Netherlands, Van Rijswijk and colleagues (30) retrospectively reviewed 48 sarcoidosis cases who required treatment with Infliximab (5mg/kg intravenously at 0, 2, 6, 10, 14 and 18 weeks). All patients had previously received immunosuppression; 30 had received prednisolone and methotrexate, 12 received prednisolone only and 1 received methotrexate only. Data involving quality of life measures, including fatigue scores (within the ‘Checklist of Individual Strength’ questionnaire), had been collected pre-treatment in the most recent 27 cases. The fatigue score improved by a mean of 5.3 points from a baseline of 49.4 points (p=0.003), although it is not clear if this is a clinically important improvement. Furthermore, as this was a retrospective review, and without a comparator or placebo group, it is not possible to definitively attribute the change in fatigue scores to the intervention and does not conclude the benefits of TNF-inhibition in treating fatigue.

Heij and colleagues (13) undertook a small double-blind, randomised controlled trial investigating the safety and efficacy of ARA 290 for symptoms of small fibre neuropathy in sarcoidosis patients. Only 12 patients were included in the active treatment arm. Of these patients, four were receiving steroids and 1 received a systemic anti-inflammatory drug. Participants were excluded if they received an anti-TNF drug within six months of the trial. Fatigue was measured using the FAS score, though this was a secondary outcome. The baseline groups were imbalanced with regards to fatigue (mean FAS 37.9 in the ARA 290 group, 33.6 in the placebo group), but despite the higher levels of fatigue in the treatment arm at baseline there was an identical reduction in FAS scores over the four week trial period in both arms. There was no evidence that ARA 290 improved fatigue scores, but the trial was not powered for this outcome and was very short. It did show improvement in its primary outcome (the small fibre neuropathy screening list) as well as being well tolerated; if a
larger and longer trial investigating the drug were to be performed, measuring change in fatigue would be important to ensure that a treatment effect was not missed in this trial.

**Neuro-stimulants**

Two randomised-controlled trials, both using a cross-over design, investigated the use of neurostimulants for treating fatigue in sarcoidosis. One trial investigated the use of armodafinil[31] and one trial investigated dexmethylphenidate[32]. In each trial the change in fatigue was the primary outcome being measured.

Lower and colleagues[32] treated ten patients with sarcoidosis-related fatigue with dexmethylphenidate and assessed its response by measuring fatigue using the FAS score. Participants received up to 10mg twice daily of dexmethylphenidate or matched placebo and were investigated weekly for eight weeks per arm (treatment and placebo). All patients were receiving at least one systemic agent for their sarcoidosis. Treatment effect in the intervention arm was seen after five weeks of therapy, with a mean reduction of five points in the FAS score after eight weeks of treatment. The number of patients meeting the MCID was not reported. The drug was well tolerated, with no withdrawals. The small scale of the trial means that the results should be interpreted with caution, but suggests clinical benefit.

In a cross-over trial by Lower and colleagues[31], fifteen patients with stable sarcoidosis received up to 250mg of armodafinil daily, all of whom underwent polysomnography and multiple sleep latency testing pre- and post-intervention. All participants had received systemic treatment for sarcoidosis (prednisolone, methotrexate, leflunomide, hydroxychloroquine, azathioprine or anti-TNF therapy) and nine were receiving continuous positive airway pressure therapy. There was a mean reduction of 4.5 points in the FAS score (the primary outcome) and nine patients (64%) exhibited a reduction of four points or more. Participants receiving armodafinil did have a prolonged sleep onset latency
compared with placebo which was statistically significant, although it is not clear if this was a clinically significant difference as no participants discontinued the medication due to insomnia. The paper did not report randomisation or allocation procedures, although it was double-blind and appeared an appropriate design for the study’s aims. The small number of patients and short period of the trial suggest that further evidence is needed to confirm the efficacy of armodafinil in this setting.

*Non-pharmacological treatment strategies*

Three recent papers have investigated the effect of structured physical training programmes on fatigue in sarcoidosis. Marcellis and colleagues(33) undertook an observational case-series study of twenty-four sarcoidosis patients in the Netherlands suffering with fatigue and/or impaired exercise tolerance. The intervention consisted of both upper- and lower-extremity peripheral muscle resistance training, with progressively increasing resistance through the training period, and endurance training, consisting of either treadmill or walking or cycling on an ergometer. Eighteen patients completed the entire training regime, with six not completing the programme. A statistically significant improvement was observed in FAS scores at the completion of the intervention: mean baseline and post-exercise FAS scores of 29.7 and 27.0 points respectively. Of the eighteen participants who completed the exercise programme, six patients (33.3%) had a reduction of four points in their FAS score; when using the alternative MCID of a 10% reduction, nine patients (50%) met this criteria. This study had no comparator group, it did not state how many patients were approached or screened for inclusion, and patients entering the trial and completing the exercise programme are likely to have been motivated to undertake such an intervention. Although the results from this study suggest that the intervention is beneficial for sarcoidosis-associated fatigue, it is unlikely to be beneficial in all patients.
Strookappe and colleagues (34) investigated physical activity programmes in a similar cohort of Dutch sarcoidosis sufferers in a retrospective observational study. From an initial cohort of 147 sarcoidosis patients who had undergone physical performance assessment, 49 patients undertook a 12-week supervised exercise programme. This was similar to that described by Marcellis and colleagues, with the peripheral muscles strengthening individualized for each patient. Twenty-one (42.9%) of the group were receiving steroids. A comparison was made with 41 sarcoidosis patients who had chosen not to undertake the programme but undertook identical physical assessments. Following re-assessment at the end of the 12-week period there was a statistically significant within-group improvement in FAS scores in those who received physical therapy (29.8 pre, 25.6 post, p=0.009) whereas the comparator group had a non-significant reduction (pre 30.3, post 28.6, p=0.408). In the exercise programme group, 74.4% of patients had an improvement in their FAS score of four points or more, although 48.5% of the comparator group showed the same reduction despite not receiving any intervention. The within-group results suggest that physical training may be beneficial for fatigue, though the results should be interpreted with caution. As with the other trials investigating exercise programmes on fatigue, participants within the trial would have self-selected as a group keen to undertake the intervention. Even given this source of bias, patients who chose not to receive the intervention reported a clinically significant improvement in their fatigue. Finally, although the authors state that blinding was not possible in this trial, it would be possible to blind assessors to patient groups when assessing physical and psychological parameters pre- and post-intervention; whether this occurred was not stated in the paper.

Another study from Strookappe and colleagues (35) investigated the physical training in patients with end-stage sarcoidosis-related pulmonary fibrosis, alongside a cohort with idiopathic pulmonary fibrosis (IPF). Twelve patients with stage IV sarcoidosis participated in a 12-week exercise programme, similar to those described in the previous trials (33, 34). Patients were recruited from the same centre and during the same period as those included in another study by the same authors (34), though it is not specified whether these patients were also included in the results from
the earlier study. Despite this, the study was included as it reported the results of a sub-group of patients with end-stage pulmonary sarcoidosis, which were not reported separately in the other paper. FAS score was measured pre- and post-intervention. Baseline mean FAS score was 25.1 in the sarcoidosis group, with only six patients (50%) having clinically significant fatigue. Following the programme, four of the six patients reported improved fatigue, although the paper does not state whether participants that improved met the MCID for the FAS score. This study did not have a non-intervention group, and only a small number of participants with fatigue at baseline were included. The same source of biases existed as described with the other two studies investigating this intervention, making it difficult to be confident of the effect of physical training programmes for sarcoidosis patients suffering fatigue.

*Excluded Studies of Interest*

Two studies investigating anti-TNF-alpha therapy in large cohorts were not included in the systematic review; the first had no measurement of pre-treatment fatigue scores (14) and the second failed to describe a quantitative change in fatigue (36). One article describing methylphenidate use for sarcoidosis-associated fatigue was excluded because it was not a full article and did not describe a baseline fatigue score. Furthermore, one cross-sectional study has suggested that the anti-malarial hydroxychloroquine may have benefits on fatigue.

One study investigating cognitive failure and sarcoidosis-associated fatigue in 343 patients was excluded as baseline assessment of fatigue occurred after patients had already been established on treatment (14). The study was a six-month cross-sectional assessment of patients who had already received various therapies, including TNF-inhibition in 42 patients. The results showed an improvement in FAS scores in the anti-TNF alpha therapy group over six months (baseline FAS 32.8 +/- 7.31, six month change -4.90 +/- 5.57) when compared with patients on no treatment (baseline FAS 28.6 +/- 7.94, six month change 0.44 +/- 5.13), or on corticosteroids with or without
antimetabolite (methotrexate) therapy (baseline FAS 28.2+/-7.81, six month change +1.19+/-4.87).

The patients in the anti-TNF-alpha therapy group had higher fatigue scores on their initial questionnaires compared with the other groups, and after six months there was no difference in fatigue scores between patients receiving other forms of therapy. However, without baseline characteristics in each group before commencing treatment it is not possible to directly compare the results of the groups, or establish the effect of treatment on fatigue scores.

A further study investigating TNF-inhibitor treatment in sarcoidosis patients (adalimumab or infliximab) recorded pre-treatment fatigue levels in 111 patients was excluded because no numerical data for change in fatigue measure (FAS) were included(36). All patients had received prednisolone and methotrexate before receiving TNF blockade, and had evidence of ongoing disease activity despite treatment. Of the 111 patients included, 100 (90.1%) reported a FAS score >21 (mean baseline FAS 33.0) and 59 reported severe fatigue (FAS score >34). After 12 months of therapy 60 patients who were fatigued at baseline had improvement in their fatigue score; unfortunately the definition of improvement and the scale of change in the FAS score required to be classified as a responder are not stated, therefore it is not possible to evaluate whether the intervention was clinically effective from these results.

Methylphenidate was used in a series of five patients with severe sarcoidosis-associated fatigue that was described in a letter from Wagner and colleagues(37). Five patients received 10mg twice daily of methylphenidate. There was no formal measure of fatigue severity at baseline, but the paper describes a statistically significant reduction on the “Symptoms of Fatigue” scale after one month. There were positive reports from four of the five patients, with two reporting that they felt as if their lives were “back to normal”. The five patients continued on methylphenidate long-term; at two-years, all five of the patients remained on methylphenidate and reported continuing improvement in fatigue, although no formal fatigue scoring was performed. The authors concluded that further
studies in larger groups of patients are required, though at the time of writing only the two small cross-over studies of armodafinil and dexamethylphenidate have been undertaken.

Although no papers directly investigated the use of chloroquine or hydroxychloroquine for fatigue, a possible effect of the drug on fatigue scores was noted in a cross-sectional study comparing two cohorts of sarcoidosis patients(2). This paper was not included in this systematic review as it did not report change in fatigue scores pre- and post-treatment, but the authors noted that patients receiving hydroxychloroquine (n=22) had lower fatigue scores than patients on other agents in the absence of any other differences in disease activity or severity. The lack of pre- and post-treatment fatigue scores, as well as the small number of patients receiving the agent, mean that conclusions about the effectiveness of hydroxychloroquine for treating fatigue cannot be directly drawn from these results.

Discussion

The evidence base for treating fatigue in sarcoidosis remains weak. Only eight trials were identified, all of which have been presented here. Of these studies, all were either small or were of poor quality study design which led to the possibility of inherent biases affecting the results. This makes it difficult to draw strong conclusions about the benefits of each therapy.

In patients with clinically significant fatigue with evidence of disease activity despite use of first line immunosuppressants, anti-TNF-alpha therapy may be indicated. In the absence of active, organ-threatening disease the risks and potential side-effects of these drugs make them difficult to recommend for treating fatigue alone.

Physical exercise programmes appear to lead to improvements in fatigue scores, but in the one trial that had a comparison group(34) almost half of the controls demonstrated clinically significant improvements in fatigue without any intervention. The patients who did enrol on the exercise
programmes were likely to have been motivated to undertake this and therefore most likely to benefit. Nevertheless, improvements were seen in physical measures beyond fatigue and so in patients with physical limitation and fatigue who express an interest in undertaking physical therapy a structured exercise programme may provide benefits.

The management of fatigue in patients with quiescent disease is often a clinical challenge, especially given the potential side effects of disease-modifying medications. Neurostimulants such as modafinil or methylphenidate may be appropriate in these cases. The two trials investigating these interventions were well designed, but only included a small number of patients. Long-term use of these medications has been safe in other conditions (ADHD) but the trials investigating their use in sarcoidosis have been very short. Further evidence to investigate the longer-term safety and efficacy of these medications is required.

Chloroquine and hydroxychloroquine are considered effective for treating cutaneous sarcoidosis(38). Its use in patients with sarcoidosis-associated fatigue who require corticosteroid therapy has been suggested in a previous review(39). The possible effectiveness of treating fatigue with these agents is interesting, but evidence from trials investigating pre- and post-intervention fatigue scores is needed before stronger recommendation can be made for its use.

Future research considerations

The main limitation of the evidence base for managing fatigue is the lack of trials of sufficient sample size or duration to make firm recommendations for managing patients with fatigue in clinical practice. Much of the data available are from observational studies or studies of less than 30 patients. Any future trials investigating therapies for treating sarcoidosis should include fatigue as an outcome measure given the frequency and significance of fatigue in sarcoidosis cohorts.

In patients with quiescent disease, where fatigue is the primary symptom driving treatment decisions, more randomised placebo controlled clinical trials are required. The need to eliminate any
placebo effect is important; in one of the trials included in this review (13) almost identical changes in fatigue from baseline were seen in both intervention and placebo arms. Designing these trials appropriately to inform clinical decision making is therefore the primary concern. The randomised trials that have already been performed have been very short, either four- or eight-weeks duration. Clinical use of agents such as neurostimulants would likely be over many months and future trials should therefore assess change in fatigue scores over a much longer period of time than previously seen, at least six months.

There are limitations to this review. The review included only English language papers, although no papers in other languages were found in the search strategy. Although efforts were made to contact authors regarding missing data or unclear elements of trial design there remains gaps in the data presented here. Furthermore, the existing data is limited; the studies included involved only a small number of participants or followed a methodology that would include intrinsic bias.

Conclusions

The available data for treating sarcoidosis-associated fatigue is limited. Anti TNF-alpha therapies appear to improve fatigue but all data comes from observational trials without placebo arms. The neurostimulants dexamphetamine and armodafinil both appeared to improve fatigue scores compared with placebo but the trials were very small and short. Given the frequency that fatigue occurs in sarcoidosis, and the importance of this symptom for patients, larger and longer trials are necessary to help inform management decisions. The increasing awareness of fatigue as a problematic manifestation of sarcoidosis will hopefully ensure that any future trials investigating interventions for sarcoidosis will include measures of fatigue.

4,229 words
**Funding:** This report is independent research supported by the National Institute for Health Research (NIHR Doctoral Research Fellowship, Dr Chris Atkins, DRF-2015-08-190). The views expressed in this publication are those of the authors and not necessarily those of the NHS, the National Institute for Health Research or the Department of Health.

**Conflict of interest statement:** Both authors confirm that there are no conflicts of interest.
Table 1 – Overview of trials in sarcoidosis patients including change in fatigue as an outcome measure

<table>
<thead>
<tr>
<th>Author and Year</th>
<th>Study Design</th>
<th>Intervention</th>
<th>Dose</th>
<th>Comparator</th>
<th>Duration</th>
<th>No. participants</th>
<th>% Male</th>
<th>Age (mean ± SD)</th>
<th>Disease stage (0/I/II/III/IV)</th>
<th>Measurement of fatigue</th>
<th>INTERVENTION: Pre- and Post-fatigue scores (mean ± S.D. unless stated)</th>
<th>COMPARATOR: Pre- and Post-fatigue scores (mean ± S.D. unless stated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erckens et al. 2011 (29)</td>
<td>Case series</td>
<td>Adalimumab</td>
<td>40mg S/C weekly</td>
<td>None</td>
<td>12 months</td>
<td>26</td>
<td>36.6%</td>
<td>51 ± 15</td>
<td>16/4/4/2/0</td>
<td>FAS</td>
<td>Pre: 37.9 ± 2.6 Post: 33.3 ± 2.8 Change: -5.3 ± 8.5</td>
<td>N/A</td>
</tr>
<tr>
<td>Heij et al. 2012 (13)</td>
<td>RCT</td>
<td>ARA 290</td>
<td>2mg IV weekly</td>
<td>Placebo</td>
<td>4 weeks</td>
<td>22 (10 placebo)</td>
<td>50%</td>
<td>48.1 ± 2.7</td>
<td>Not stated</td>
<td>FAS</td>
<td>Pre: 37.9 ± 2.6 Post: 33.3 ± 2.8 Change: -5.3 ± 8.5</td>
<td>Pre: 33.6 ± 2.3 Post: 29.8 ± 3.3</td>
</tr>
<tr>
<td>Van Rijswijk et al. 2013 (30)</td>
<td>Retrospective case review</td>
<td>Infliximab</td>
<td>5mg/kg IV at 0,2,6,10,14 and 18 weeks</td>
<td>Placebo</td>
<td>18 weeks</td>
<td>27‡</td>
<td>60%</td>
<td>48.9 ± 10.1</td>
<td>Not stated</td>
<td>FAS</td>
<td>Pre: 49.4 ± 9.2 Change: -5.3 ± 8.5</td>
<td>Pre: 33.6 ± 2.3 Post: 29.8 ± 3.3</td>
</tr>
<tr>
<td>Lower et al. 2013 (31)</td>
<td>Cross-over RCT</td>
<td>Armodafinil</td>
<td>Up to 250mg twice daily</td>
<td>None</td>
<td>8 weeks per arm</td>
<td>15</td>
<td>33.3%</td>
<td>54 (range 35-62)</td>
<td>5/7/14/5/14</td>
<td>FACIT-F (range)</td>
<td>Pre: 28.5 ± 5.4 Post: 27.7 ± 5.7 (6 of the 12 patients had FAS &gt;21 at baseline)</td>
<td>Pre: 37 (14-43) Change: -4.5 (-20, 5) FACIT-F (range): Pre: 23 (10-47) Change: +9 (12-26)</td>
</tr>
<tr>
<td>Strookappe et al. 2015 (35)</td>
<td>Case series</td>
<td>Physical activity programme</td>
<td>12-week physical activity programme</td>
<td>None</td>
<td>12 weeks</td>
<td>12</td>
<td>91.7%</td>
<td>53.2 ± 11.7</td>
<td>0/0/0/0/12</td>
<td>FAS</td>
<td>Change: +3.5 (-9,14)</td>
<td>Change: +3.5 (-9,14)</td>
</tr>
<tr>
<td>Marcellis et al. 2015 (33)</td>
<td>Retrospective, Observational</td>
<td>Physical activity programme</td>
<td>13-week physical activity programme</td>
<td>None</td>
<td>13 weeks</td>
<td>24</td>
<td>75%</td>
<td>49.4 ± 10.5</td>
<td>4/11/22/0/4</td>
<td>FACIT-F (range)</td>
<td>Change: 5 (-17,11)</td>
<td>Change: 5 (-17,11)</td>
</tr>
<tr>
<td>Strookappe et al. 2015 (34)</td>
<td>Case series</td>
<td>Physical activity programme</td>
<td>12-week physical activity programme</td>
<td>41 patients who chose not to participate in programme</td>
<td>12 weeks</td>
<td>49</td>
<td>57.1%</td>
<td>47.6 ± 11.3</td>
<td>0-1 = 29.2% II-III = 66.6% IV = 4.2%</td>
<td>FACIT-F (range)</td>
<td>Change: 5 (-17,11)</td>
<td>Change: 5 (-17,11)</td>
</tr>
</tbody>
</table>
†Group values at the end of the placebo phase not given, data only presented as average values across placebo phase.
‡48 patients included in study but quality of life scores (including fatigue) only available in 27.
FAS = Fatigue Assessment Scale; FACIT-F = Functional Assessment of Chronic Illness Therapy – Fatigue; MCID = Minimal Clinically Important Difference

<table>
<thead>
<tr>
<th>Statistical difference vs. comparator</th>
<th>FAS: p=0.0295 (between groups)</th>
<th>P &lt; 0.01 compared with baseline</th>
<th>Non-significant between groups</th>
<th>FAS: p=0.003 compared with baseline</th>
<th>FACIT-F: p=0.0040 (between groups)</th>
<th>Non-significant compared with baseline</th>
<th>P = 0.003 compared with baseline</th>
</tr>
</thead>
<tbody>
<tr>
<td>Within group (intervention): p=0.009</td>
<td>Not stated</td>
<td>Not clear; 14/21 less fatigued but doesn’t state MCID used.</td>
<td>Not stated</td>
<td>64% treated with armodafinil improved FAS by 4 points (MCID); only 7% in placebo</td>
<td>66% of patients with FAS &gt;21 at baseline had improvement in FAS, not specified if MCID used</td>
<td>6 (3%) improve FAS by 4 points (MCID); 9 (50%) improved FAS by 10%</td>
<td>74.4% of intervention group and 48.5% of comparator group reduced FAS by 4 points (MCID).</td>
</tr>
</tbody>
</table>

| Number of participants with clinically significant improvement | Not stated | Not clear; 14/21 less fatigued but doesn’t state MCID used. | Not stated | 64% treated with armodafinil improved FAS by 4 points (MCID); only 7% in placebo | 66% of patients with FAS >21 at baseline had improvement in FAS, not specified if MCID used | 6 (3%) improve FAS by 4 points (MCID); 9 (50%) improved FAS by 10% | 74.4% of intervention group and 48.5% of comparator group reduced FAS by 4 points (MCID). |

| Drop outs/ Side Effects | No withdrawals 4 patients required lower afternoon dose | No withdrawals. 1 severe injection site reaction | No drop outs | 3 patients discontinued within 6 infusions: 1) Allergic reaction 2) Progression of dyspnoea 3) Hepatitis (due to methotrexate) | 1 withdrawal due to severe anxiety | No withdrawals | 6 patients withdrew; 3 = problems other than sarcoid 2 = health insurance problems 1= No reason | Not stated |

| Evidence of treatment effect in fatigue | Yes | Yes | No | Yes | Yes | Unclear | Yes | Yes |

**Notes:**
- Statistical difference vs. comparator
- FAS: Fatigue Assessment Scale
- FACIT-F: Functional Assessment of Chronic Illness Therapy – Fatigue
- MCID: Minimal Clinically Important Difference
### Table 2 – Risks of bias within trials in sarcoidosis patients reporting fatigue as an outcome measure

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sequence Generation</strong></td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>HIGH RISK</td>
<td>HIGH RISK</td>
</tr>
<tr>
<td>Random sequence computer-generated</td>
<td>No randomisation (NOT RCT)</td>
<td>Computer generated randomisation code</td>
<td>No randomisation (NOT RCT)</td>
<td>No statement on randomisation procedure</td>
<td>No randomisation (NOT RCT)</td>
<td>No randomisation (NOT RCT)</td>
<td>No randomisation (NOT RCT)</td>
<td>No randomisation (NOT RCT)</td>
</tr>
<tr>
<td><strong>Allocation Concealment</strong></td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>LOW RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
</tr>
<tr>
<td>Pharmacy-controlled allocation</td>
<td>No concealment, all patients receive drug</td>
<td>Pharmacy-controlled allocation</td>
<td>No concealment, all patients received drug</td>
<td>Only allocating pharmacist unblinded</td>
<td>Double-blind with low risk of breaking blinding</td>
<td>No blinding</td>
<td>No blinding</td>
<td>No blinding</td>
</tr>
<tr>
<td><strong>Blinding of participants, personnel and outcome assessors</strong></td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>LOW RISK</td>
<td>HIGH RISK</td>
<td>HIGH RISK</td>
<td>HIGH RISK</td>
</tr>
<tr>
<td>Double-blind with low risk of breaking blinding</td>
<td>No blinding</td>
<td>Only allocating pharmacist unblinded</td>
<td>No blinding</td>
<td>Double-blind with low risk of breaking blinding</td>
<td>No blinding</td>
<td>No blinding</td>
<td>No blinding</td>
<td>No blinding</td>
</tr>
<tr>
<td><strong>Incomplete outcome data</strong></td>
<td>UNCLEAR</td>
<td>LOW RISK</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
<td>LOW RISK</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
</tr>
<tr>
<td>No description of missing data points or handling of missing data</td>
<td>No description of missing data points or handling of missing data</td>
<td>No description of missing data compensated for by taking forward last value</td>
<td>No description of missing data points or handling of missing data</td>
<td>No description of missing data points or handling of missing data</td>
<td>Low risk of bias from physical function pre- and post.</td>
<td>No description of missing data points or handling of missing data</td>
<td>No description of missing data points or handling of missing data</td>
<td>No description of missing data points or handling of missing data</td>
</tr>
<tr>
<td><strong>Selective outcome reporting</strong></td>
<td>LOW</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
<td>LOW</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
<td>UNCLEAR</td>
</tr>
<tr>
<td>All outcomes reported</td>
<td>No protocol available</td>
<td>No protocol available</td>
<td>No protocol available</td>
<td>All outcomes reported</td>
<td>No protocol available</td>
<td>No protocol available</td>
<td>No protocol available</td>
<td>No protocol available</td>
</tr>
<tr>
<td><strong>Other sources of bias</strong></td>
<td>NO</td>
<td>Study design (case series) limits conclusions – no comparator group to eliminate placebo effect.</td>
<td>Baseline imbalance in FAS and health status score (SF36) between groups – significantly lower fatigue scores in placebo arm.</td>
<td>Retrospective review – data collected pre- and post-intervention but high risk of bias from retrospective nature</td>
<td>No other clear causes of bias identified; cross-over ensures groups balanced, patients are own controls. Small sample.</td>
<td>Not an RCT, also participants enrolling on programme would self-select as motivated people, generalisable.</td>
<td>Not an RCT, also participants enrolling on programme would self-select as motivated people, generalisable.</td>
<td>Patients choosing the intervention would be a self-selecting cohort; controls not randomised but refused intervention</td>
</tr>
<tr>
<td><strong>Overall risk of bias</strong></td>
<td>LOW</td>
<td>HIGH – Study design (case series) means no blinding, randomisation or comparator.</td>
<td>LOW – Well designed RCT but not powered to look at change in fatigue.</td>
<td>HIGH – Design (retrospective case series) has no blinding, randomisation or comparator.</td>
<td>UNCLEAR – Issues with description of randomisation allocation and concealment mean study at risk of bias</td>
<td>HIGH - Study design (case series) means no blinding, randomisation or comparator.</td>
<td>HIGH - Study design (case series) means no blinding, randomisation or comparator.</td>
<td>HIGH - Study selection intervention group, high risk of bias given control group declined intervention</td>
</tr>
</tbody>
</table>

---

Note: The table above summarizes the risks of bias within trials in sarcoidosis patients reporting fatigue as an outcome measure. Each column represents a different aspect of bias, such as sequence generation, allocation concealment, blinding of participants, incomplete outcome data, selective outcome reporting, and other sources of bias. The ratings range from LOW RISK to HIGH RISK, indicating the level of risk associated with each aspect. The table highlights specific issues for each trial, such as the method of randomization, the quality of blinding, and the handling of missing data.
References:

37. Wagner MT, Marion SD, Judson MA. The effects of fatigue and treatment with methylphenidate on sustained attention in sarcoidosis. Sarcoidosis Vasculitis and Diffuse Lung Diseases. 2005;22(3):235-.
