Original research

Trajectories of fatigue in actively treated patients with established rheumatoid arthritis starting biologic DMARD therapy

Sella Aarrestad Provan, Brigitte Michelsen, Joseph Sexton, Tillmann Uhlig, Hilde Berner Hammer

ABSTRACT

Objectives To define fatigue trajectories in patients with rheumatoid arthritis (RA) who initiate biological DMARD (bDMARD) treatment, and explore baseline predictors for a trajectory of continued fatigue.

Methods One-hundred and eighty-four patients with RA initiating bDMARDs were assessed at 0, 1, 2, 3, 6 and 12 months. Swollen and tender joint counts, patient reported outcomes (PROMs), blood samples and ultrasound examinations were collected at each time point. Fatigue was assessed by the fatigue Numeric Rating Scale (0–10) from the Rheumatoid Arthritis Impact of Disease (RAID) questionnaire. Clinically significant fatigue was predefined as fatigue ≥4. Three trajectories of interest were defined according to level of RAID fatigue: no fatigue (≤3 at 5/6 visits), improved fatigue (≤4 at start, but ≤3 at follow-up) and continued fatigue (≥4 at 5/6 visits). Baseline variables were compared between groups by bivariate analyses, and logistic regression models were used to explore baseline predictors of continued vs improved fatigue.

Results The majority of patients starting bDMARD therapy followed one of three fatigue trajectories, (no fatigue; n=61, improved; n=33 and continued fatigue; n=53). Patients with continued fatigue were more likely to be anti–B27 antibody and/or rheumatoid factor positive and had higher baseline PROMs compared to the other groups, while there were no differences between the groups for variables of inflammation including. Patient global, tender joint count and anxiety were predictors for the continued fatigue trajectory.

Discussion A trajectory of continued fatigue was determined by PROMs and not by inflammatory RA disease activity.

INTRODUCTION

Rheumatoid arthritis (RA) is an inflammatory joint disease that can potentially affect all synovial joints, causing joint destruction and disability. Fatigue is a common complaint among patients with RA and has a major impact on the burden of disease. In a recent Danish study, 61% of the patients with RA reported having moderate to severe fatigue. Fatigue was identified by patients with RA as an important patient-reported outcome measure (PROM) during the development of the RA Impact of Disease (RAID) questionnaire. The aetiology of fatigue is multifactorial and is likely to be influenced both by disease-related inflammation, by pain, cognitive factors, depressive symptoms and personal issues in the individual patient’s life.

Disease activity in RA is assessed by composite measures of disease activity that usually include measures of swollen and tender joints, inflammation and patient-reported global assessment. Disease remission, at which disease activity falls to a pre-specified level, is the treatment target. RA is usually treated by BMJ.

Key messages

What is known?

► Fatigue in RA is known to be multifactorial.

What this study adds?

► The longitudinal course of fatigue of the majority of patients with longstanding RA who start bDMARD therapy could be classified into three trajectories.

► Only 18% of patients starting a bDMARD were categorised into the improved fatigue trajectory, in this heterogeneous cohort.

► The trajectory of continued fatigue was predicted by PROMs and not by inflammatory RA disease activity or ultrasonographic joint evaluation.

How this impact on clinical practice?

► Patients who experience improved fatigue most frequently did so within the first months after the start of bDMARD therapy.

► PROMs are important determinants of continued fatigue and should be the focus of future interventions targeting this symptom.
with conventional synthetic disease-modifying anti-rheumatic drugs (csDMARDs) from the time of diagnosis, and if remission is not reached within the desired time frame, dose escalation or change to a biologic DMARD (bDMARD) such as a TNFalpha-inhibitor (TNFi) is recommended.1

Relief of fatigue may improve quality of life, and treatment with TNFi has been shown to affect fatigue levels in a substantial proportion of patients. However, fatigue has been shown to persist at a clinically significant level in the majority of patients who are in remission.5 There is little knowledge regarding the factors predicting continued fatigue in patients treated with a bDMARD.

The objectives of this study were to define trajectories of fatigue in patients with RA who initiate bDMARD treatment and to define baseline predictors for continued fatigue.

PATIENTS AND METHODS

Patients

This study uses data from a previously described cohort of patients with RA fulfilling the American College of Rheumatology 1987 criteria6 who were consecutively enrolled when they initiated or changed biologic DMARD treatment in the period from January 2010 to June 2013.7 The patients were comprehensively assessed at baseline and after 1, 2, 3, 6, and 12 months. Patients who had answered the RAID questionnaire, including the question regarding fatigue, were selected for the current analyses.7 The study (Anzctr.org.au identifier ACTRN12610000284066) was approved by the Norwegian Regional Committee for Medical and Health Research Ethics South East (reference number 2009/1254) and the patients gave their written informed consent according to the Declaration of Helsinki.

Clinical, laboratory and ultrasound assessments

At each visit clinical examinations including examiner’s global disease activity (0–100 VAS), 28 tender and swollen joint counts were performed by study nurses and blood samples were taken for laboratory assessments of erythrocyte sedimentation rate (ESR), (CRP) mg/L, Rheumatoid factor (RF) and anti-cyclic-citrullinated protein (CCP) antibodies. In addition, calprotectin (µg/L), a major leukocyte protein shown to be sensitive for rheumatic inflammation, was measured in EDTA plasma with an ELISA from CALPRO AS (Lysaker, Norway) according to the instructions of the manufacturer.

An experienced sonographer (HBH) performed all the ultrasonography (US) examinations (using a dedicated Siemens Antares Excellence version machine, 5–13 MHz probe with PD frequency 7.3 MHz and PRF 391 Hz, with no updates of the software during the study), blinded from the clinical assessments and laboratory markers from the same time points, as well as from previous US results. Grey scale (GS) ultrasound reflecting synovial hypertrophy and power Doppler (PD) ultrasound reflecting vascularity in the synovium, were scored semiquantitatively on a 4-point scale (0=no, 1=minor, 2=moderate, 3=major presence) of 36 joints (bilateral wrist (radiocarpal, midcarpal, radioulnar joints scored separately), metacarpophalangeal 1–5, proximal interphalangeal 2–3, elbow, knee, ankle (tibiotalar), metatarsophalangeal 1–5) with the US atlas as reference,7 as well as four tendon sheaths (bilateral extensor carpi ulnaris and tibialis posterior). The sonographer has previously shown high reliability for US assessments of these joints and tendons.8

Patient-reported outcome measures (PROMs)

The patients were examined when initiating bDMARD therapy and assessed at baseline and after 1, 2, 3, 6 and 12 months. The PROMs included the RAID score, which has been developed in liaison with patients and includes fatigue, sleep, pain, physical function, physical and psychological well-being as well as coping, all scored on a 10 points numeric rating scale (NRS).3 The fatigue question is anchored at no fatigue and total exhaustion. In addition, patients scored their joint pain (0–100 visual analogue scale (VAS)), patient’s global disease activity VAS (0–100), modified health assessment questionnaire (MHAQ, 0–3, two main questions from the pain-catastrophizing assessment7 and short form-36 mental health scale score (SF36MH). Anxiety and depression were assessed by the use of the Hospital Anxiety and Depression scale (HADS). The score of widespread pain was self-reported by indicating presence of pain at 19 pre-specified sites during the past 7 days.

Identification of fatigue trajectories

Prior to data analyses, we had defined three trajectories to be of interest: no clinically significant fatigue, improved fatigue and continued clinically significant fatigue. Clinically significant fatigue was defined as RAID fatigue after discussions and a review of the literature (≥4).9 The authors SAPR, TU and HBH examined the fatigue trajectory of each individual patient in line-plots. Using a visual data-driven approach we sought to categorise the maximum number of participants by examining changes in fatigue in relation to the cut-off over time while also constructing mutually exclusive categories (online supplemental table 1). We decided to place the cut-off between improved and permanent fatigue as change in RAID fatigue from ≥4 to ≤3 before the 5th visit at 6 months. Patients who did not fit into the aforementioned trajectories were categorised as ‘increasing fatigue’ if there was no clinically significant fatigue at baseline but then a progression to clinically significant fatigue that persisted over several visits. Patients were categorised as ‘no discernible pattern’ if the fatigue level fluctuated over and below the cut-off for clinically significant fatigue, and as ‘missing’ if data was missing after the 2nd or 3rd visit.
Baseline variables were compared across groups using independent samples T-test, Wilcoxon rank-sum or Chi² for normally distributed, non-parametric distribution or dichotomised data, respectively.

Logistic regression models were used to explore baseline predictors of continued vs improved fatigue. Each variable was entered into univariate models, adjusted for age and gender. Variables that predicted the outcome with a p≤0.1 were entered into the multivariable model. Variables that were highly correlated (Pearsons correlation co-efficient >0.7) were examined in separate the multivariable models. Variables were removed according to level of significance (backward stepwise), until the final model only included significant predictors at p<0.05. Age and gender were forced into the model. Possible confounding was explored in the final model by re-entering the discarded variables to look for significance in the final model and examining the main effects for change in significance or a 25% or more change in the co-efficient.

Disease activity indicators and PROMs were plotted as estimated marginal means in figures across the trajectories, and differences between groups assessed by mixed models with maximum likelihood, adjusted for age, gender and time.

Last observation carried forward replaced missing data if data from one visit, but not the first visit, was missing (<5% of the different variables). Patients with >1 missing visit were not included in the analyses. All calculations were performed by the use of STATA version 14, and p<0.05 was considered statistically significant.

**RESULTS**

Two hundred and eight patients had completed the RAID questionnaire for fatigue, 24 of these were missing from >1 visit and were excluded from the study.

Longitudinal trajectories of fatigue were examined in 184 patients (online supplemental table 1). The definition of three fatigue trajectories that allowed for the categorisation of the maximum number of patients into the three trajectories were: No fatigue (61 patients): fatigue score ≤3 at baseline and at 4 out of 5 follow-up visits. Improved fatigue (33 patients): fatigue score ≥4 at baseline but ≤3 at the 6- and 12-month visits. Continued fatigue (53 patients): fatigue level ≥4 at 5 out of 6 visits. In addition, no discernible pattern was found in 34 patients and 3 patients had an increasing level of fatigue. The fatigue levels of four trajectories, adjusted for age and gender are presented in figure 1. 24 patients left the study after the second (4 patients) third (7 patients) or fourth visit (13 patients), these had comparable age, gender distribution and disease activity, to the participants remaining in the study. More details regarding the categorisation into trajectories are presented in the online supplemental table 1.

Baseline fatigue was significantly higher in the continued fatigue group compared to the no fatigue and improved fatigue groups, and was also higher in the group with improved fatigue compared to no fatigue.

Table 1 describes demographics and clinical factors of the three groups. Patients with continued fatigue were more often female, were more likely to be anti-CCP and RF-positive, but had lower levels of education compared to patients with no clinically significant fatigue, and were more likely to be anti-CCP positive and have lower education than patients who had improved fatigue. Patients with continued fatigue had higher levels of all PROMs compared to patients with no fatigue, and also reported higher levels of most PROMs compared to patients who improved from their fatigue. Patients who had improved fatigue had higher levels of patient global, sleep disturbance, widespread pain and pain catastrophising compared to patients with no fatigue. The average change (SD) in fatigue from baseline to 12-months was 0.49 (1.24) for the no-fatigue, 4.07 (0.28) for the improved fatigue and 1.49 (2.16) for the continued fatigue trajectory.

In logistic multivariable regression the trajectory of continued fatigue vs improved fatigue was predicted by baseline calprotectin, patient’s global, tender joints and anxiety (table 2). The HADS anxiety and depression scales were strongly correlated (Pearsons r=0.85) and were therefore explored in different models. The presented model was chosen based on the log likelihood.

Figure 2 illustrates the trajectories for the three groups of interest, showing the continued fatigue group to have significantly higher levels of anxiety level, sleep disturbance, pain and pain catastrophizing in contrast to no higher level of inflammation as assessed by CRP and ultrasound PD at all visits. The main co-efficients of the mixed model analyses are shown in online supplemental table 1. The mixed models show that the different trajectories are best differentiated by PROMs and not measures of inflammation.

**DISCUSSION**

The present study adds to the limited knowledge of longitudinal fatigue trajectories in patients with RA. We
defined and applied three different fatigue trajectories that categorise over 70% of the patients. PROMs were found to be highly significant predictors for the development of continued fatigue.

In a cohort of patients with longstanding RA we have found that 53/206 (26%) continued to have a clinically significant fatigue after the first year of bDMARD therapy. This group was defined by a combination of a high frequency of anti-CCP and RF-positive patients in univariate analyses, and long disease duration in bivariate analyses, but there was no evidence of higher baseline inflammati...
trajectory, this is not in agreement with the findings of the same systematic review\textsuperscript{11} which concluded that disease duration did not influence fatigue.

This study presents an expansive data-collection of 6 prospective visits during a 12-month follow-up. A weakness of the study is that fatigue was only measured by a single instrument, RAID question on fatigue is however identical to the first severity question in the Bristol Numerical Fatigue Rating Scale which has been found to be reliable\textsuperscript{16}. Another weakness is that LOCF was used to impute variables and we cannot be certain that all variables were missing completely at random. However, only a very small proportion of variables were missing and all examinations were performed by the author HBF giving us some knowledge of the logistical reasons for missingness. Lastly, the study included 208 patients and the further division into trajectories posed a limitation on the number of variables that could be included in the regression models and on the statistical methods chosen to build the regression models.

The inclusion of repeated ultrasonography examinations allowed for a comprehensive assessment of inflammatory disease activity at all visits providing a novel and important confirmation of the lack of association between joint inflammation and fatigue, and this is a strength of the study. The cohort is a heterogeneous sample of real-life clinic and differs from cohorts of the more stringent pharmaceutical trials. This is another strength of the study. We have chosen to use a clinical rather than statistical approach to define the trajectories of interest. We believe that this strength in the identification of trajectories of fatigue in relation to a pre-defined cut-off is of relevance to both clinicians and patients. In order to categorise the maximum number of patients we allowed fatigue levels to cross the pre-defined cut-off for one out of six visits in the trajectory of no fatigue and continued fatigue and we acknowledge that this methodology is more open for individual interpretation. We show that a substantial proportion of patients still have clinically significant fatigue after 1 year of bDMARD and that the majority of patients who improve do so within the first months of bDMARD therapy.

**CONCLUSION**

The longitudinal course of fatigue in patients with long-standing RA who started bDMARD therapy follows one of three trajectories in the majority of patients. The development of continued fatigue was determined by PROMs and not by inflammatory RA disease activity.

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**Table 2** Baseline predictors for continued fatigue vs improved (logistic regression analyses)

<table>
<thead>
<tr>
<th>Variables</th>
<th>Univariate</th>
<th>Multivariable</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>OR (95% CI)†</td>
<td>OR (95% CI)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>1.01 (0.98–1.05)</td>
<td>1.02 (0.98–1.06)</td>
</tr>
<tr>
<td>Female gender</td>
<td>1.79 (0.52–6.15)</td>
<td>1.57 (0.34–7.35)</td>
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<tr>
<td>Higher Education</td>
<td>0.30 (0.11–0.81)*</td>
<td></td>
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<tr>
<td>Anti-CCP positive</td>
<td>5.51 (1.02–29.60)*</td>
<td></td>
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<tr>
<td>RF positive</td>
<td>1.72 (0.56–5.29)</td>
<td></td>
</tr>
<tr>
<td>Disease duration</td>
<td>1.03 (0.98–1.10)</td>
<td></td>
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<tr>
<td>RA disease activity baseline</td>
<td></td>
<td></td>
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<tr>
<td>Swollen joints (28)</td>
<td>1.01 (0.92–1.10)</td>
<td></td>
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<tr>
<td>CRP (mg/L)</td>
<td>0.99 (0.97–1.01)</td>
<td></td>
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<tr>
<td>Calprotectin</td>
<td>0.77 (0.57–1.04)</td>
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<tr>
<td>Sum PD</td>
<td>0.98 (0.95–1.02)</td>
<td></td>
</tr>
<tr>
<td>Sum BM</td>
<td>0.99 (0.97–1.02)</td>
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<tr>
<td>PROMs</td>
<td></td>
<td></td>
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<tr>
<td>Patient global VAS</td>
<td>1.04 (1.02–1.07)**</td>
<td>1.03 (1.01–1.06)*</td>
</tr>
<tr>
<td>Tender joints 28</td>
<td>1.21 (1.08–1.35)</td>
<td>1.19 (1.04–1.35)*</td>
</tr>
<tr>
<td>RAID sleep</td>
<td>1.26 (1.06–1.51)*</td>
<td></td>
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<tr>
<td>Widespread pain</td>
<td>1.08 (0.98–1.20)</td>
<td></td>
</tr>
<tr>
<td>HADS anxiety</td>
<td>1.17 (1.03–1.33)*</td>
<td>1.23 (1.05–1.45)*</td>
</tr>
<tr>
<td>HADS depression</td>
<td>1.26 (1.06–1.49)</td>
<td></td>
</tr>
<tr>
<td>Pain Catastrophizing</td>
<td>1.23 (0.89–1.69)</td>
<td></td>
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</tbody>
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†Univariate model adjusted for age and gender.

\(p<0.10 \, ^*p<0.05 \, ^{**}p<0.001\).

No fatigue: Fatigue \(\leq 3\) at baseline and at 4/5 visits. Improved fatigue: Fatigue \( \geq 4\) at baseline, but \(\leq 3\) at 6- and 12-months visit. Continued: Fatigue \( \geq 4\) at 5/6 visits. GS, Grey scale; HADS, Hospital anxiety and depression scale; PD, Power Doppler; VAS, Visual analogue scale.

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**Figure 2** Longitudinal change in disease activity and PROMS across trajectories of fatigue Comparison of the longitudinal course of disease activity and patient-reported outcomes (PROMs) using mixed models via maximum likelihood. VAS, Visual analogue scale; HADS, Hospital anxiety and depression scale.
REFERENCES