Factors that influence fatigue status in patients with severe rheumatoid arthritis (RA) and good disease outcome following 6 months of TNF inhibitor therapy: a comparative analysis

Patricia Minnock1 · Douglas J. Veale1,2,3 · Barry Bresnihan1,2,3 · Oliver FitzGerald1,2,3 · Gabrielle McKee4

Received: 18 June 2015 / Revised: 23 September 2015 / Accepted: 28 September 2015 / Published online: 9 October 2015 © The Author(s) 2015. This article is published with open access at Springerlink.com

Abstract The objective of the present study is to determine the factors associated with persistent fatigue in patients with severe rheumatoid arthritis (RA) and good disease response to 6 months of tumour necrosis factor inhibitor therapy. Eligible patients with either persistent (PF) or no fatigue (NF) were compared. Using validated questionnaires and bivariate analysis, this cross-sectional survey explored if clinical characteristics, pain, self-efficacy, sleep and mood/depression differed between groups. Patients with PF (PF; NF) (n=28; 28) reported significantly more overall pain (11.3±9.4 (0–33); 6.9±8.9 (0–33)), more recent and current pain intensity (41.4±26.6 (0–80) 24.4±26.6 (0–100) and depression (11.8±7.5 (1–35); 8.2 ±6.6 (0–26)), than the NF group. There was no significant difference between groups in self-efficacy and both groups experienced poor sleep quality (Pittsburgh Sleep Quality Index >5). Despite having good disease response, the PF group had significantly higher rheumatoid factor incidence, disease activity score-28, early morning stiffness duration and lower incidence of ever-failing disease-modifying anti-rheumatic drugs than the NF group. These findings enhance the fatigue literature in patients with RA prescribed tumour necrosis factor (TNF) inhibition therapy, identifying the potentially modifiable factors of pain and depression, previously demonstrated to be strongly associated with fatigue in non-biologic populations. In addition, this study highlights the association between persistent fatigue and an on-going state of low disease activity. This infers that more judicious disease management could minimise the symptom burden of pain and depression and consequentially fatigue.

Keywords Disease outcome · Fatigue · Pain · Rheumatoid arthritis · Self-efficacy

Fatigue is recognised as an important symptom in rheumatoid arthritis (RA) by patients and clinicians [1–3]. While not yet defined in clear terms, its multidimensional nature has been identified [4, 5]. Studies have shown that the causes and consequences of fatigue in RA can be attributed to illness-related aspects, physical functioning, cognitive/emotional functioning and social aspects [6]. It is acknowledged that RA fatigue likely results from a complex interplay between these factors [5, 7] and requires further study to determine causation and best management [3, 6, 7].

Debate around to what degree fatigue is a primary or secondary symptom of the autoimmune inflammatory process [8, 9] has occurred in parallel with the positive impact of biologic drug therapies on disease management and patient-reported
outcome [3, 8, 10], the Outcome MEasurement in Rheumatology Clinical Trials (OMERACT) patient perspective initiative [1, 11, 12] and the international endorsement of fatigue as a patient-centred outcome for inclusion in all patient studies [2]. Despite this elevated status, from patients’ perspectives, fatigue remains unmanageable and professional support remains rare [13].

The overarching aim of this study was to contribute to the rational development of appropriate prevention and management strategies for fatigue. This is contingent on identifying its drivers. Reasons for minimal insight into fatigue to date include (i) a lack of data from longitudinal observational studies exclusive to patients with RA, (ii) studies that were either confined to patients with only mild to moderately active disease [14] or (iii) studies that excluded populations from biologic registers on the basis of disease severity [9]. While researchers have identified key contributory factors, such as pain, sleep, self-efficacy and mood [9, 15], none has reported investigating the relevance of these factors in patients in different disease states of activity with differing fatigue levels, especially in the context of modern biological therapies [3]. Defining persistent fatigue as patient-reported presence of moderate or greater fatigue, the purpose of this study was to examine the factors associated with persistent fatigue in patients with established RA despite demonstrating a good disease response to 6 months of tumour necrosis factor (TNF) inhibitor therapy.

Methods

Study design and patient population This cross-sectional single-site study used an observational, naturalistic comparative design. The procedures followed in this study are in accordance with the ethical standards of the relevant Ethics and Medical Research Committees and with the Helsinki Declaration of 1975, as revised in 1983. The study population was patients with (1) confirmed RA [16], (2) minimal age 18 years and (3) who had demonstrated a moderate to good disease response following 6 months of TNF inhibitor treatment for moderate to high disease activity [8], according to the disease activity score-28 (DAS28)-based European Union of Leagues against Rheumatism (EULAR) Response Criteria (ERC) [17]. Patients were excluded from the study if they had a primary diagnosis of fibromyalgia syndrome or chronic fatigue syndrome.

The DAS28 is a standardised, composite outcome measure of disease activity calculated using a four variable formula (DAS28-4 (variable) (C-reactive protein (CRP))=0.56∗√(TJC28, 28 joint count for tenderness)+0.28∗√(SJC28, 28 joint count for swelling)+0.36∗ln (CRP)+1)∗0.014∗(GH, patient global health assessment on a 0–10 numeric rating scale)+0.96. The DAS28 is a continuous scale ranging from 0 to 9.4, indicating level of RA disease activity. DAS28 >5.1 reflects high disease activity, 5.1–3.2 reflects moderate disease activity, and <3.2 indicates low disease activity. Remission is achieved when a DAS28 score <2.6 is obtained. The ERC calculates both the degree of reduction in DAS28 as well as current DAS28 so as to classify individual patients into three categories of disease response: good (DAS28 improvement of >1.2), moderate (DAS28 improvement of >0.6–≤1.2) and non-response (DAS28 improvement of <0.6) [17]. Using the ERC, patients who had achieved at least a moderate improvement in their DAS28 in response to 6-month TNF inhibition therapy were identified.

To meet the analysis needs of this study, assuming a large effect size (0.8), alpha of 0.05 and power of 0.80, a sample size of 54 was required, with a minimum of 27 in each group [18]. Given the small sample size, the exact p value with the Z approximation test was reported; p values <0.05 were regarded as statistically significant. Holm’s sequential Bonferroni post hoc adjustment was used in the comparison tests to prevent type 1 error [19].

Data collection

Purposive consecutive sampling was used to recruit patients meeting the inclusion criteria over a period of 18 months. Sociodemographic details, clinical characteristics and clinical assessments of disease activity using the ACR core set of outcome measures (28-swollen joint count, 28-tender joint count, pain, patient global health assessment, functional assessment using the Health Assessment Questionnaire Disability Index (HAQ-DI) and biochemical acute phase reactant measure, C-reactive protein (CRP)) [20], along with fatigue levels, were derived from the case notes. In all scales, higher scores indicated worst outcome. Following telephone contact to inform patients of the study, questionnaires to assess pain, self-efficacy, sleep, mood and depression were posted to eligible participants enclosed with an invitation to partake in the study, a consent form and a prepaid addressed envelope. All persons gave their informed consent prior to their inclusion in the study.

Data collection tools

The sociodemographic and clinical characteristics derived from the case notes were gender, age, educational background, current smoking status, disease duration, rheumatoid factor (RF), haemoglobin level, disease-modifying anti-rheumatic drug (DMARD) history and early morning stiffness (EMS) duration in minutes. DAS28 and the derived ERC were the indices used to identify the study population.
Fatigue Fatigue was measured using a standard validated five-point verbal scale recording fatigue over the last week. This scale has previously demonstrated good validity and sensitivity in patients with RA [4]. Patients indicated their level of fatigue as being the following: none, mild, moderate, severe and very severe. These data were used to differentiate between those with and without persistent fatigue,

Pain Pain was assessed using the multidimensional Short-Form McGill Pain Questionnaire (SF-MPQ) [21]. It contains five separate pain scores. The first three (i) a sensory descriptor score of the pain experience, (ii) an affective descriptor score for pain over the past week were rated on a four-point intensity scale (none (0) – excruciating pain (4)). The remaining two captured (iv) overall past week pain intensity using a 100-mm visual analogue scale (VAS) and (v) current pain intensity captured verbally using five descriptive terms (0 = ‘no pain’ to 5 = ‘excruciating’ pain). In this study, internal consistency (Cronbach’s α 0.72) was found to be consistent with previous values shown in RA and fibromyalgia populations (Cronbach’s α 0.73 to 0.89) [22].

Self-efficacy Patient self-efficacy (SE) was measured using the Arthritis Self-Efficacy Scales (ASES), incorporating three subscales, which is extensively used in RA studies [23]. The ASES includes 20 items scored separately on a Likert scale of 1 (very uncertain) to 10 (very certain) from which the overall mean of the subscale items is derived and higher scores indicate greater confidence or SE. The three subscales are as follows: pain (five items), functioning (nine items) and other symptoms of SE (six items). The derived internal consistency (Cronbach’s α 0.79–0.84) was consistent with previously reported values (Cronbach’s α 0.76 to 0.89) [24].

Sleep The Pittsburgh Sleep Quality Index (PSQI), a 19-item self-rated questionnaire was used to measure seven different components of sleep, duration, quality, number of arousals, efficacy, disturbances, use of sleep medications and daytime dysfunction during the previous month. Its suitability for use in patients with RA is documented [25]. The PSQI has reported high construct validity [26], sensitivity and specificity. Scores range from 0 to 25; a total score of >5 is indicative of sleep disturbances. In this study, internal consistency among the seven component scores was somewhat lower (Cronbach’s α 0.65) than that reported (Cronbach’s α 0.83) in generic studies [26].

Mood Two separate scales were used to assess different aspects of mood and depression. The Profile of Mood States (POMS) short form, a widely used tool to assess transient and distinct mood states, has been used in a variety of RA fatigue studies [27]. This 37-item scale can be scored into five subscales (tension-anxiety, depression-dejection, fatigue-inertia, confusion-bewilderment and vigour). Each item is rated from 0—not at all to 4—extremely. Its discriminant validity is supported by being consistently more highly related to corresponding mood measures (for example, sad and depressed) (mean r=66.6), than non-corresponding mood scales (for example, vigour versus inertia) (mean r= 49.5), when compared with other scales [28]. Internal consistency in this study using Cronbach’s α ranged from 0.54 to 0.86 across the six items.

The Beck Depression Inventory II (BDI-II), an extensively tested and validated questionnaire, comprises a series of 21 questions developed to measure the intensity, severity and depth of depression. Each question assesses a specific symptom, 13 items assess psychological symptoms, and 8 items assess physical symptoms. Each item uses a four-point scale (0 to 3), and total score range is 0–63. A score of 0–13 is considered minimal, 14–19 mild, 20–28 moderate and 29–63 severe. Its appropriateness as a screening tool for depression in patients with RA has been demonstrated [29]. It has a high construct validity (Cronbach’s α=0.80) and discriminates between depressed and non-depressed groups [30]. Permission to use all scales was secured.

Analysis The sample was subdivided into two groups for comparative purposes. Those who self-reported moderate or greater fatigue at time of recruitment were labelled the persistent fatigue group while those who reported mild or no fatigue were labelled the no-fatigue group. To enhance sample homogeneity, the two groups were matched approximately for gender, age and disease duration. Analysis was conducted using the statistical package SPSS 16.0 for Windows. Numbers and percentage counts were used to report nominal data (education/smoking status/RF); mean and standard deviations were reported on the normally distributed variables (DAS28) and medians and range for skewed data such as haemoglobin (HB) and early morning stiffness (EMS) duration. Inferential statistics to test for differences between subgroup means included chi-square, independent sample t tests and Mann-Whitney U test as appropriate. The significance level was originally set at 0.05 and then subjected to a Holm’s sequential Bonferroni adjustment as multiple tests which are indicated at the bottom of the relevant table.

Results

Patient demographic and clinical details All eligible patients who were invited to take part in the study accepted (n=64), and all patients gave their informed consent prior to their inclusion in the study. Of these 64 patients, 47 % (n=28) qualified as the persistent fatigue group. The other 53 % (n=
Table 1  Sociodemographic and clinical characteristics of patients with good disease outcome and either persistent or no fatigue

| Patients characteristics | Persistent fatigue | No fatigue | p value |
|--------------------------|--------------------|------------|---------|
| n (study, control)       | n (valid %)        | n (valid %) |         |
| Female gender* (n=28, 28) | 22 (79)            | 23 (82)    | ≤0.213  |
| Age (mean±SD (range), years)b (n=28, 28) | 58±11 (26–77) | 58±11 (23–81) | ≤0.716  |
| Disease duration (mean±SD (range), years)b (n=28, 28) | 14±11 (0–36) | 14±12 (0–39) | ≤0.949  |
| Smoking statusa (n=27, 24) | Current 9 (32) | 7 (29) | ≤0.139  |
| Educational backgrounda (n=26, 21) | Primary 7 (25) | 4 (20) |         |
| | Secondary 12 (43) | 7 (35) |         |
| | Third level 6 (21) | 9 (45) |         |
| Rheumatoid factora (n=28, 28) | Positive 15 (54) | 4 (14)a | =0.006* |
| | Negative 13 (46) | 23 (82) |         |
| Ever-failed DMARDa (n=28, 27) | Yes 18 (64) | 24 (86)a | =0.036* |
| | No 10 (36) | 3 (11) |         |
| Ever-failed biologic (n=28, 28) | No 0 | 0 |         |
| Haemoglobin levels (median (range), g/dl) (n=22, 13) | 6 months 13 (9–15) | 13.4 (9–17) | ≤0.452  |
| HAQ-DI (mean±SD) (n=26, 16) | 6 months 1.11±(0.62) | 0.76±(0.55) | =0.07  |
| Early morning stiffness duration (median, min)c (n=27, 26) | 6 months 10 (0–180) | 10 (0–30) | =0.001* |
| Swollen joint count (mean±SD) (n=28, 16) | 6 months 2.2±(2.9) | 1.0±(1.7) | <0.135  |
| Tender joint count (mean±SD) (n=28, 16) | 6 months 3.2±(4.7) | 1.0±(2.3) | <0.028* |
| Pain (mean±SD) (n=28, 16) | 6 months 4.4±(2.2) | 2.3±(1.0) | <0.001* |
| Patient global health (mean±SD) (n=28, 16) | 6 months 4.9±(2.2) | 2.3±(1.0) | <0.0001* |
| C-reactive protein (mean±SD)c (n=28, 16) | 6 months 4 (0–42) | 4 (0–27) | <0.718  |
| C-reactive protein n (%) | 6 months | | |
| <2 | 2 (7.1) | 1 (5.9) |         |
| <10 | 21 (75.0) | 12 (70.6) |         |
| >10 | 5 (17.9) | 4 (17.9) |         |
| DAS-28 (mean±SD, min-max)c (n=28, 16) | 6 months 3.3±(1.1) | | |
| (1.7–5.8) | 2.4±(0.7) | =0.002* |         |
| (1.2–4.4) | | |         |
| DAS-28 (n=28, 16) | 6 months | | |
| >3.2 | 15 (62.5) | 1 (37.5) |         |
| <2.7 | 9 (7.1) | 13 (92.9) |         |

*Significance at the <0.05

a Chi-square test.
b Independent sample t test

c Mann-Whitney U test

HAQ-DI Health Assessment Questionnaire Disability Index, DMARD disease-modifying anti-rheumatic drugs
36) of patients demonstrated a good fatigue outcome, qualifying as the no-fatigue group, from which 28 patients were purposively selected based on age, gender and disease duration.

The majority of the patients were female. There was no significant difference between the groups’ smoking status, educational background or HB. Significantly higher RF-positive status, EMS duration and mean DAS28 and a significantly lower incident of ever failing a disease-modifying anti-rheumatic drug (DMARD) were seen in the persistent fatigue group. While the difference between groups for the HAQ-DI was not statistically significant, it did exceed the known minimal clinical important difference [31] (Table 1).

A significant difference between fatigue groups was demonstrated in total pain experienced during the previous week ($u=0.241, p=0.021$). Sensory ($u=0.251, p=0.02$), pain intensity VAS ($u=0.223, p=0.023$) and current pain intensity subscales ($u=0.231, p=0.009$) were also significant (Table 2).

The three subscales for SE pain, functioning and other symptoms did not differ significantly between fatigue groups (Table 3). There was no significant difference in the total sleep score or the sleep subscales between groups (Table 4). There was also no difference between groups in relation to use of sleep medications. The sleep score greater than 5 recorded in both groups was clinically indicative of poor sleep quality.

The mean BDI-II Score±SD (range) of $11.8±7.5$ (1–35) for the persistent fatigue group was shown to be statistically different from that of the no-fatigue group $8.2±6.6$ (0–26), $U=265$ and exact $p=0.037$ (two-tailed). Total scores on the BDI-II were in what is considered to be the mild (0–13) to moderate range (14–19) for depression. Mean mood (POMS) total scores or subscales did not differ significantly between groups (Table 5).

**Discussion**

This study identified factors peculiar to patients with RA and persistent fatigue, despite a good disease response to 6-month TNF inhibitor therapy, by comparing those with persistent fatigue to those with no fatigue. There were no significant differences in key or minor characteristics between the groups at initiation of TNF inhibitor treatment, as reported elsewhere [8]. However, there were some statistically or clinically significant differences in group characteristics at 6 months post-initiation of TNF inhibitor therapy. The HAQ-DI did exceed the known minimal clinical important difference between groups even though this was not statistically significant [31]. This is an important observation, as functional health status is repeatedly reported to influence fatigue outcome in RA [6, 32]. The proportion of patients within the persistent fatigue group who were RF-positive was significantly greater than that in the no-fatigue group. The persistent fatigue group also reported significantly longer EMS duration; this finding is in keeping with a published longitudinal study which showed a significant association between stiffness on awakening and higher fatigue [33]. Furthermore, qualitative study has shown that patients regard the absence of stiffness as an important criterion for remission [34]. The lower incidence of ever-failing DMARDs found in the persistent fatigue group was contrary to what might have been expected.

**Table 2** Comparative analysis of pain experience in patients with good disease outcome and either persistent or no fatigue (Mann-Whitney $U$ test)

| Short-Form McGill Pain Questionnaire (SF-MPQ) | Persistent fatigue | No fatigue | $p$ value | Level of significance $^a$ |
|------------------------------------------------|--------------------|------------|-----------|--------------------------|
| Descriptors (during last week)                | Mean±standard deviation (range) | $p$ value | Level of significance $^a$ |
| Mean sensory ($n=27, 28$) (0–33)              | $8.0±6.4$ (0–24)    | $5.2±6.5$ (0–23) | 0.02* | 0.013                     |
| Mean affective ($n=27, 28$) (0–12)           | $2.7±3.3$ (0–12)    | $1.8±2.8$ (0–10) | 0.235 | 0.05                      |
| A: Total descriptor score ($n=27, 28$) (0–45) | $11.3±9.4$ (0–33)   | $6.9±8.9$ (0–33) | 0.021* | 0.016                     |
| B: Past week: VAS pain intensity ($n=27, 26$ (0–100 mm)) | $41.4±26.6$ (0–80) | $24.4±26.6$ (0–100) | 0.023* | 0.025                     |
| C: Current: pain intensity ($n=27, 28$)      |                     |            |           |                          |
| No pain ($n=27, 28$)                          | 3 (11)              | 9 (32)     | 0.009* | 0.01                      |
| Mild                                           | 6 (22)              | 11 (39)    |           |                          |
| Discomforting                                  | 13 (48)             | 5 (18)     |           |                          |
| Distressing                                    | 3 (11)              | 2 (7)      |           |                          |
| Horrible                                       | 2 (7)               | 0          |           |                          |
| Excruciating                                   | 0                   | 1 (3)      |           |                          |

$^a$Significant

$^a$ Level of significance as set using Holm’s sequential Bonferroni adjustment
Inclusion criteria for this study specified the demonstration of a moderate to good disease response to TNF inhibitor therapy, according to the ERC, as opposed to a specific DAS28 target. However, the mean DAS28 score (≈3.3) reflected a low disease activity state for patients with persistent fatigue. However, the statistically significant and even lower mean DAS28 score demonstrated in 93% of those with no fatigue reflected a state of clinical remission (DAS28 <2.6) [17], versus only 37% of the persistent fatigue group. These data also indicate an association between persistent fatigue and other unfavourable disease characteristics [35], including a positive RF [36]. Assertions within the literature suggest that DAS28 <2.6 is more representative of minimal disease activity than of remission [37]. Overall, these data show an association between persistent fatigue and an ongoing state of low disease activity. It therefore highlights the need to look beyond disease remission for optimal disease management to improve symptom burden and patient outcomes.

With respect to the patient-reported outcome, pain findings confirm that in patients with disease severity which warranted TNF inhibitor therapy, those with persistent fatigue experienced more pain than those with no fatigue [3, 6, 38]. This interrelationship was previously reported from studies with either mixed diagnostic groups or non-biologic-treated populations [3, 9]. Further, the nature of this pain was reported to be more physical in nature as opposed to being affective/emotionally driven. Although the recent systematic review of fatigue and factors related to fatigue in RA found conflicting evidence with respect to the relationship between pain and

| Table 3 | Comparative analysis of arthritis self-efficacy in patients with good disease outcome and either persistent or no fatigue (Mann-Whitney U test) |
|---------------------------------|----------------|----------------|----------------|----------------|
| Subscales (range 1–10) (n=28)   | Persistent fatigue | No fatigue | p value | Significance levela |
|---------------------------------|----------------|----------------|----------------|----------------|
| **Self-efficacy pain subscale:** How certain are you that you can… | | | | |
| 1 Decrease your pain quite a bit? | 5.9±2.0 | 6.1±3.0 | | |
| 2 Continue most of your daily activities? | 6.5±2.5 | 6.9±2.9 | | |
| 3 Keep arthritis pain from interfering with your sleep? | 6.1±3.0 | 6.7±2.8 | | |
| 4 Make small/moderate reduction in pain? | 5.5±2.7 | 4.5±2.9 | | |
| 5 Make large reduction in your arthritis pain by using methods other than taking extra medication? | 4.3±2.7 | 4.8±3.2 | | |
| **Mean pain** | 5.6±1.9 | 6.1±2.3 | 0.359 | 0.025 |
| **Self-efficacy function subscale:** As of now how certain are you that you can… | | | | |
| 1 Walk 100 ft on flat ground in 20 s? | 6.7±3.3 | 7.6±3.1 | | |
| 2 Walk ten steps downstairs in 7 s? | 6.3±3.3 | 6.7±3.1 | | |
| 3 Get out of armless chair quickly without using your hands for support? | 5.7±3.4 | 6.7±3.1 | | |
| 4 Button and unbutton three medium-size buttons in a row in 12 s? | 6.8±3.1 | 7.7±2.4 | | |
| 5 Cut two bite-size pieces of meat with a knife and fork in 8 s? | 6.8±3.2 | 7.5±2.7 | | |
| 6 Turn an outdoor tap all the way on and off? | 6.6±2.9 | 6.8±3.1 | | |
| 7 Scratch your upper back with both your right and left hands? | 5.1±3.3 | 5.8±3.3 | | |
| 8 Get in and out the passenger side of car without assistance from another person and without physical aids? | 7.2±2.5 | 7.7±2.9 | | |
| 9 Put on long-sleeved front-opening shirt or blouse (without buttoning) in 8 s? | 7.2±2.9 | 8.2±2.5 | | |
| **Mean function** | 6.7±2.4 | 7.1±2.5 | 0.413 | 0.05 |
| **Self-efficacy other symptom subscale:** How certain are you that you can… | | | | |
| 1 Control your fatigue? | 4.9±2.4 | 6.1±2.3 | | |
| 2 Regulate your activity so as to be active without aggravating your joints? | 6.2±2.6 | 7.0±2.4 | | |
| 3 Do something to help yourself feel better if feeling blue? | 6.1±2.7 | 7.4±2.7 | | |
| 4 Manage arthritis pain during daily activities? | 6.4±2.8 | 7.2±2.5 | | |
| 5 Manage arthritis symptoms so that you can do the things you enjoy doing? | 5.7±2.7 | 7.0±2.8 | | |
| 6 Deal with frustration of arthritis? | 5.9±2.8 | 7.1±2.8 | | |
| **Mean other symptom** | 5.8±2.4 | 7.2±2.2 | 0.022 | 0.017 |

*a Level of significance as set post-Holm’s sequential Bonferroni adjustment
fatigue, nonetheless, pain was one of three variables with the strongest evidence for a relationship with fatigue [6]; this study supports the dominant interrelationship between pain and fatigue. As patients in this study had demonstrated a good disease response to 6 months of TNF inhibitor treatment at time of assessment, these findings in a new treatment group add to the body of evidence of previous work illustrating that physical pain is a more important driver of persistent fatigue in RA than is inflammation [3, 38, 39]. This study result therefore underscores why proactive pain management should continue to be a priority in comprehensive patient care.

SE as a previously reported contributor to fatigue [6] was not substantiated by this study. In a previous systematic review, five studies, undertaken in largely non-biologic populations, supported SE and only one demonstrated no interrelationship between SE and fatigue [6]. Nonetheless, as in this current study, the systematic review failed to demonstrate that SE majorly influenced fatigue status in RA. This current lack of association may, in part, be explained by the superior clinical efficacy of biologic medications.

This study found that patients with persistent fatigue had significantly more depression than those with no fatigue. While published evidence is conflicting with respect to the contribution of depression towards the symptom of fatigue in RA; nonetheless, strong evidence of a relationship between these two symptoms exists [6]. While this association between fatigue and depression is recognised in RA [6], so too is the existence of a bidirectional causal pathway [40]. One previous study found a steep increase in depression as functional limitation increased [41], while another TNF inhibition study in RA showed that both fatigue and pain significantly impacted on changes in depression status and that clinical remission improved symptoms of depression [42]. There is a risk that the interrelationship between pain and fatigue [43] is minimised by some literature that highlights the mediating effect of mood disorders [6]. With respect to symptom management for this patient group, such an assumption could result in suboptimal pain and fatigue management. Moreover, physiological pain and depression, identified as problematic within the persistent fatigue group, are potential modifiable drivers of fatigue.

Similar patient numbers per subgroup used sleep-inducing medication, and both groups experienced a poor sleep quality. In keeping with previous work [44], this was attributed to physical discomforts, namely, that joint pain and limitation

| Profile of mood states (range) | Persistent fatigue Mean (SD), range n=28 | No fatigue Mean (SD), range n=28 | p value |
|--------------------------------|------------------------------------------|---------------------------------|---------|
| Depression-dejection (0–32)   | 4.8±5.4 (0–19)                           | 3.3±5.2 (0–20)                  | 0.102   |
| Vigour-activity (0–24)         | 7.0±5.6 (0–18)                           | 8.5±5.7 (0–20)                  | 0.293   |
| Anger-hostility (0–28)         | 3.9±4.5 (0–18)                           | 2.4±2.9 (0–9)                   | 0.134   |
| Tension-anxiety (0–24)         | 4.1±3.8 (0–13)                           | 3.6±4.3 (0–16)                  | 0.351   |
| Confusion-bewilderment (0–20)  | 2.1±1.9 (0–8)                            | 2.1±3.0 (0–10)                  | 0.324   |
| Fatigue-inertia (0–20)         | 5.7±4.9 (0–20)                           | 5.6±5.8 (0–21)                  | 0.469   |
| POMS Total (0–100)             | 13.7±19.8 (14–59)                        | 8.6±18.4 (18–49)                | 0.306   |
| Beck Depression Inventory (BDI)| n=28                                      | n=28                           |         |
| Level of depression (0–63)     | 11.8±7.5 (1–35)                          | 8.2±6.6 (0–26)                  | 0.037*  |

*Significant at <0.05 level
due to pain mediated the association between arthritis and insomnia [45]. However, findings from this study do not suggest any association between persistent post-treatment fatigue and sleep quality.

It is important to highlight study limitations inherent in a naturalistic study design. This study was conducted in a real-life clinical situation with an inability to address completeness of data and study attrition. Study replication sufficiently powered to support multiple regression analysis is recommended. Further, the comparative study of persistent post-treatment fatigue was restricted to the single disease entity of RA. While a high response rate was achieved, the generalisation of results to non-RA patients is not appropriate. It is also acknowledged that the uniqueness of a single-site academic centre which had poor representation of minority groups restricts the representativeness of findings. Data on possible confounding variables such as daily physical activities, physical exercise capacity, thyroid function and morning (08.00 h) adrenocorticotropic hormone (ACTH)/cortisol values was not collected. This comparative study adds further to the body of knowledge on factors related to persistent fatigue in patients with RA [6, 7] and to the limited body of knowledge with respect to fatigue in TNF inhibitor-treated patients [3, 32]. It firmly highlights potential modifiable factors to improve fatigue, namely, pain and depression. In addition, it highlights the association between persistent fatigue and an ongoing state of low disease activity. This infers that more judicious disease management could minimise the symptom burden of pain and depression and consequently fatigue.

Compliance with Ethical Standards This human study was approved by the appropriate ethics committees and has therefore been performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. All persons gave their informed consent prior to their inclusion in the study.

Disclosures None.

Funding This study was supported by a Health Research Board of Ireland National Council for Nursing and Midwifery Junior Clinical Scientist Award.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. Kirwan JR, Minnock P, Adebayo A, Bresnihan B, Choy E, de Wit M et al (2007) Patient perspective: fatigue as a recommended patient centered outcome measure in rheumatoid arthritis. J Rheumatol 34: 1174–1177

2. Aletaha D, Landewe R, Karonitsch T, Bathon J, Boers M, Bombardier C et al (2008) Reporting disease activity in clinical trials of patients with rheumatoid arthritis: EULAR/ACR collaborative recommendations. Ann Rheum Dis 67:1360–1364

3. Campbell RC, Batley M, Hammond A, Ibrahim F, Kingsley G, Scott DL (2012) The impact of disease activity, pain, disability and treatments on fatigue in established rheumatoid arthritis. Clin Rheumatol 31:717–722

4. Hewlett S, Hehir M, Kirwan JR (2007) Measuring fatigue in rheumatoid arthritis: a systematic review of scales in use. Arthritis Rheum 57:429–439

5. Nicassio PM, Ormseth SR, Custodio MK, Irwin MR, Olmstead R, Weissman MH (2012) A multidimensional model of fatigue in patients with rheumatoid arthritis. J Rheumatol 39:1807–1813

6. Nikolau S, Bode C, Taal E, van de Laar MA (2013) Fatigue and factors related to fatigue in rheumatoid arthritis: a systematic review. Arthritis Care Res (Hoboken) 65:1128–1146

7. Hewlett S, Chalder T, Choy E, Cramp F, Davis B, Dures E et al (2011) Fatigue in rheumatoid arthritis: time for a conceptual model. Rheumatology (Oxford) 50:1004–1006

8. Minnock P, McKee G, Bresnihan B, FitzGerald O, Veale DJ (2014) How much is fatigue explained by standard clinical characteristics of disease activity in patients with inflammatory arthritis? A longitudinal study. Arthritis Care Res (Hoboken) 66:1597–1603

9. Bergman MJ, Shahouri SS, Shaver TS, Anderson JD, Weidensaul DN, Busch RE et al (2009) Is fatigue an inflammatory variable in rheumatoid arthritis (RA)? Analyses of fatigue in RA, osteoarthritis, and fibromyalgia. J Rheumatol 36:2788–2794

10. Wolfe F, Michaud K (2004) Fatigue, rheumatoid arthritis, and anti-tumor necrosis factor therapy: an investigation in 24,831 patients. J Rheumatol 31:2115–2120

11. Minnock P, Kirwan J, Bresnihan B (2009) Fatigue is a reliable, sensitive and unique outcome measure in rheumatoid arthritis. Rheumatology (Oxford) 48:1533–1536

12. Nicklin J, Cramp F, Kirwan J, Urban M, Hewlett S (2010) Collaboration with patients in the design of patient-reported outcome measures: capturing the experience of fatigue in rheumatoid arthritis. Arthritis Care Res (Hoboken) 62:1552–1558

13. Repping-Wuts H, van Riel P, van Achterberg T (2008) Rheumatologists’ knowledge, attitude and current management of fatigue in patients with rheumatoid arthritis (RA). Clin Rheumatol 27:1549–1555

14. Repping-Wuts H, Fransen J, van Achterberg T, Bleijenberg G, van Riel P (2007) Persistent severe fatigue in patients with rheumatoid arthritis. J Clin Nurs 16:377–383

15. van Hoogmoed D, Fransen J, Bleijenberg G, van Riel P (2010) Physical and psychosocial correlates of severe fatigue in rheumatoid arthritis. Rheumatology (Oxford) 49:1294–1302

16. Arnett FC, Edworthy SM, Bloch DA, McShane DJ, Fries JF, Cooper NS et al (1988) The American Rheumatism Association 1987 revised criteria for the classification of rheumatoid arthritis. Arthritis Rheum 31:315–324

17. Fransen J, van Riel PL (2005) The Disease Activity Score and the EULAR response criteria. Clin Exp Rheumatol 23:S93–S99

18. Faul FEE, Buchner A, Lang A-G (2009) Statistical power analyses using G*Power 3.1: tests for correlation and regression analyses. Behav Res Methods 41:1149–1160

19. Holm S (1979) A simple sequentially rejective multiple test procedure. Scand J Stat 6:65–70

20. Felson DT, Anderson JJ, Boers M, Bombardier C, Furst D, Goldsmith C et al (1995) American College of Rheumatology. Preliminary definition of improvement in rheumatoid arthritis. Arthritis Rheum 38:727–735

21. Melzack R, Katz J (eds) (2001) The McGill Pain Questionnaire: appraisal and current status. The Guilford Press, New York
22. Burckhardt CS, Bjelle A (1994) A Swedish version of the short-form McGill pain questionnaire. Scand J Rheumatol 23:77–81
23. Brady TJ (2011) Measures of self-efficacy: Arthritis Self-Efficacy Scale (ASES), Arthritis Self-Efficacy Scale-8 Item (ASES-8), Children’s Arthritis Self-Efficacy Scale (CASE), Chronic Disease Self-Efficacy Scale (CDSES), Parent’s Arthritis Self-Efficacy Scale (PASE), and Rheumatoid Arthritis Self-Efficacy Scale (RASE). Arthritis Care Res (Hoboken) 63(Suppl 11):S473–S485
24. Lorig K, Chastain RL, Ung E, Shoor S, Holman HR (1989) Development and evaluation of a scale to measure perceived self-efficacy in people with arthritis. Arthritis Rheum 32:37–44
25. Wells GA, Li T, Kirwan JR, Peterson J, Aletaha D, Boers M et al (2009) Assessing quality of sleep in patients with rheumatoid arthritis. J Rheumatol 36:2077–2086
26. Carpenter JS, Andrykowski MA (1998) Psychometric evaluation of the Pittsburgh Sleep Quality Index. J Psychosom Res 45:5–13
27. Belza BL (1995) Comparison of self-reported fatigue in rheumatoid arthritis and controls. J Rheumatol 22:639–643
28. Nyenhuis DL, Yamamoto C, Luchetta T, Terrien A, Parmentier A (1999) Adult and geriatric normative data and validation of the profile of mood states. J Clin Psychol 55:79–86
29. Krug HE, Woods SR, Mahowald ML (1997) The importance of identifying depression in patients with rheumatoid arthritis: evaluation of the Beck depression inventory. J Clin Rheumatol 3:248–257
30. Beck AT, Steer RA, Carbin MG (1988) Psychometric properties of the Beck Depression Inventory: twenty-five years of evaluation. Clin Psychol Rev 8:77–100
31. Pope JE, Khanna D, Norrie D, Ouiemt JM (2009) The minimally important difference for the health assessment questionnaire in rheumatoid arthritis clinical practice is smaller than in randomized controlled trials. J Rheumatol 36:254–259
32. Druce KL, Jones GT, Macfarlane GJ, Basu N (2015) Patients receiving anti-TNF therapies experience clinically important improvements in RA-related fatigue: results from the British Society for Rheumatology Biologics Register for Rheumatoid Arthritis. Rheumatology (Oxford) 54:964–971
33. Stone AA, Broderick JE, Porter LS, Kaell AT (1997) The experience of rheumatoid arthritis pain and fatigue: examining momentary reports and correlates over one week. Arthritis Care Res 10:185–193
34. van Tuyl LH, Lems WF, Boers M (2014) Measurement of stiffness in patients with rheumatoid arthritis in low disease activity or remission: a systematic review. BMC Musculoskelet Disord 15:28
35. Smolen JS, Landewe R, Breedveld FC, Dougados M, Emery P, Gaujoux-Viala C et al. (2010) EULAR recommendations for the management of rheumatoid arthritis with synthetic and biological disease-modifying antirheumatic drugs. Ann Rheum Dis
36. Furst DE, Keystone EC, Braun J, Breedveld FC, Burmester GR, De Benedetti F et al (2011) Updated consensus statement on biological agents for the treatment of rheumatic diseases, 2010. Ann Rheum Dis 70(Suppl 1):i2–i36
37. Aletaha D, Ward MM, Machold KP, Nell VP, Stamn T, Smolen JS (2005) Remission and active disease in rheumatoid arthritis: defining criteria for disease activity states. Arthritis Rheum 52:2625–2636
38. Minnock P, Bresnihan B (2008) Fatigue is related to poor pain outcomes in women with established rheumatoid arthritis. Clin Exp Rheumatol 26:707–708
39. Druce KL, Jones GT, Macfarlane GJ, Basu MN. (2015) Determining pathways to improvements in rheumatoid arthritis fatigue: results from the BSRBR-RA. Arthritis Rheumatol.
40. Wolfe F, Michaud K (2009) Predicting depression in rheumatoid arthritis: the signal importance of pain extent and fatigue, and co-morbidity. Arthritis Rheum 61:667–673
41. Margaretten M, Barton J, Julian L, Katz P, Trupin L, Tommer C et al (2011) Socioeconomic determinants of disability and depression in patients with rheumatoid arthritis. Arthritis Care Res (Hoboken) 63:240–246
42. Kekow J, Moots R, Khandker J, Melin J, Freundlich B, Singh A (2011) Improvements in patient-reported outcomes, symptoms of depression and anxiety, and their association with clinical remission among patients with moderate-to-severe active early rheumatoid arthritis. Rheumatology (Oxford) 50:401–409
43. van Dartel SA, Repping-Wuts JW, van Hoogmoed D, Bleijenberg G, van Riel PL, Fransen J (2013) Association between fatigue and pain in rheumatoid arthritis: does pain precede fatigue or does fatigue precede pain? Arthritis Care Res (Hoboken) 65:862–869
44. Louie GH, Tektonidou MG, Caban-Martinez AJ, Ward MM (2011) Sleep disturbances in adults with arthritis: prevalence, mediators, and subgroups at greatest risk. Data from the 2007 National Health Interview Survey. Arthritis Care Res (Hoboken) 63:247–260
45. Minnock P (2011) Mixed methods research design to determine clinical characteristics, contributory factors and patients’ perceptions of fatigue in inflammatory arthritis. School of Nursing and Midwifery. [PhD Thesis], Dublin: Trinity College Dublin.