Relationship Between Fatigue and Inflammation, Disease Duration, and Chronic Pain in Psoriatic Arthritis: An Observational DANBIO Registry Study

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**ABSTRACT.** Objective. Fatigue is one of the most significant symptoms, and an outcome of great importance, in patients with psoriatic arthritis (PsA), but associations between underlying components of fatigue experienced by patients in relation to the disease have been sparsely investigated. The objectives were to describe the degree of fatigue in patients with PsA, and to examine important components associated with fatigue.

Methods. We performed a cross-sectional survey including patients registered in the Danish nationwide registry DANBIO from December 2013 to June 2014. Principal component analysis (PCA) was used to identify factors associated with fatigue.

Results. A total of 1062 patients with PsA were included in the study. A PCA reduced co-variables into 3 components explaining 63% of fatigue in patients. The first component, contributing to 31% of fatigue, was composed of inflammatory factors including swollen and tender joints, physician’s global assessment, elevated C-reactive protein (CRP), and high Pain Detect Questionnaire (PDQ) score. The second component, contributing to 17% of fatigue, consisted of increasing age and long disease duration. The third component, contributing to 15% of fatigue, consisted of high PDQ score, tender joint count, increasing age, and concomitant low CRP, suggestive of a chronic pain component consisting of central pain sensitization or structural joint damage.

Conclusion. Fatigue in patients with PsA may be driven by clinical inflammatory factors, disease duration, and chronic pain in the absence of inflammation. (First Release December 15 2019; J Rheumatol 2020;47:548–52; doi:10.3899/jrheum.181412)
Psoriatic arthritis (PsA) is a chronic inflammatory musculoskeletal disease with a prevalence of 0.2% in Denmark. The disease confers a considerable socioeconomic disease burden with decreased work productivity and increased healthcare use. Moreover, patients with PsA are characterized by a decreased quality of life compared to other patient groups and often fatigue is reported to be the factor limiting participation in daily activities.

Fatigue defined as sustained physical tiredness, mental exhaustion, and a lack of energy, is a well-known symptom of many chronic diseases and is often a crucial aspect in the management of such diseases. It is a common symptom in PsA and is deemed by patients to be one of the most significant symptoms. It is rated by patients as the worst symptom after pain and skin problems.

Though fatigue is considered an important outcome measure for patients with PsA, this outcome is not yet fully embedded in clinical practice or in the scientific thinking within this disease area, where reporting of fatigue as a patient-reported outcome is rare and studies on fatigue are limited.

However, the focus on fatigue is increasing and fatigue is now considered a core outcome according to the updated PsA core domain set from 2016. Studies have described the association between fatigue in patients with PsA and pain, female sex, physical disability, medication status, psychological distress, longstanding sick leave, and loss of ability to work. Further, biological agents have been shown to improve fatigue, suggesting a link between fatigue and inflammatory signaling. And so the inflammatory pathway is believed to be associated with several clinical manifestations of PsA. As for pain in PsA, it is traditionally considered to be of inflammatory origin, but despite better control of inflammation, some patients still report pain as a significant concern. This suggests that PsA may prompt central sensitization and thus be linked to other central mechanisms such as fatigue, and indicates why it is relevant to study fatigued Limiting participation in daily activities.

The objective of our study was to describe the degree of fatigue in patients with PsA in a nationwide study, and to examine important components associated with fatigue.

**MATERIALS AND METHODS**

**Study design and setting.** The study was designed as a cross-sectional survey including patients registered in the Danish nationwide registry, DANBIO. Recording of data in DANBIO was mandatory for patients in treatment with biological disease-modifying antirheumatic drugs (bDMARD), but DANBIO also contains treatment information on patients treated with conventional synthetic DMARD (csDMARD). PDQ was implemented on the DANBIO touch screens in Danish outpatient clinics at 22 of 24 departments of rheumatology for a period of 6 months (December 1, 2013, to June 1, 2014). The study was conducted in accordance with the STROBE statement (Supplementary Figure 1, available with the online version of this article) and according to a prespecified protocol available and published as open access at the official Website of the Parker Institute, a research unit that is part of Copenhagen University Hospital at Bispebjerg and Frederiksberg (www.parkerinst.dk). All patients registered as having PsA were invited to participate in the survey. Patients with a complete response to PDQ and a PDQ score above 0 were included in the analyses. Patient consent was obtained on the touch screen prior to the redirection to the PDQ. In accordance with Danish legislation, surveys do not require approval by ethics committees. Registrations and publications of data from clinical registries that do not pertain to human biological samples do not require patient consent or approval by ethics committees.

**Variables and outcome measures.** The VAS is a single-item scale (0–100 mm) to measure patient-reported pain, fatigue, and global health (VAS pain, VAS fatigue, VAS global health). The VAS scale has shown good reliability and performs as well as other questionnaires when assessing fatigue. In this study the VAS was used to measure patient-reported fatigue during the last week, with “0” representing “no fatigue” and “100” representing “worst imaginable fatigue.” We defined moderate-to-severe fatigue as fatigue scores ≥ 57 (chosen because 57 was the median VAS fatigue score for the population). PDQ is a mechanism-based pain classification instrument based on patient self-reported somatosensory signs and symptoms, assigning patients to one of 3 categories depending on the character of the experienced pain: neuropathic (PDQ score > 18), unclear (PDQ score 13–18), or nociceptive (PDQ score < 13). PDQ was originally developed to screen for a neuropathic pain component and based on pain phenotypic similarities to assess neuropathic pain features as a proxy of central sensitization. Statistical analysis. Patient characteristics were given with median and interquartile ranges (IQR) for continuous variables. Spearman’s p correlation coefficients were calculated to assess any potential association between fatigue scores and clinical indices. Two-sided p values < 0.05 were regarded as statistically significant.

To examine components explaining fatigue, a principal component analysis (PCA) was conducted. Variables were a priori selected based on clinical relevance with a predefined maximum allowed collinearity of 0.4. Variables included for further analysis consisted of age, disease duration, swollen/tender joint count (28 joints), pain detect score, C-reactive protein (CRP) level, and patient and doctor VAS global health scores (0–100 mm). Health Assessment Questionnaire (HAQ) scores were excluded from the PCA owing to collinearity. To assess the variability and association of components to fatigue in the entire population, multiple linear regression was conducted for VAS fatigue, with the 3 primary components identified in the PCA. A sensitivity analysis based on the PCA was constructed on VAS pain and stratification by sex, respectively, to explore any possible similarities or differences explaining fatigue when including PDQ score versus VAS pain and male versus female. IBM SPSS version 20 was used to carry out the analyses.

**RESULTS**

A total of 2388 patients were diagnosed with PsA in DANBIO, of which 2114 had a VAS fatigue score. Of these, 1062 chose to participate in the study and were included for analysis because they had a recorded PDQ score above 0. The

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The median VAS fatigue score was 57 mm for the population, and scores of 57 mm or more were considered moderate to severe fatigue. Patients with moderate to severe fatigue were predominantly female, and with higher 28-joint Disease Activity Score (DAS28)-CRP as well as higher VAS pain, VAS global health, PDQ score, and HAQ scores, compared with subjects with none to mild fatigue scores. Moreover, these patients had higher scores in physician’s global assessment (PGA), more tender and swollen joints, increased use of corticosteroids, and more often switched bDMARD (Table 1).

In the PCA (Supplementary Figure 2, available with the online version of this article), the clinical co-variables were reduced to 3 components explaining 63% of fatigue (Figure 1). The first component, contributing to 31%, was mainly constituted by inflammatory factors such as more swollen and tender joints, higher PGA, higher DAS28-CRP, and higher PDQ scores, whereas the second component mainly consisted of contributions from older age and longer disease duration, explaining 17% of fatigue. The third component, contributing to 15%, consisted of higher PDQ scores, more tender joint counts, increasing age, and concomitant low CRP.

The multiple linear regression analysis on the overall population with VAS fatigue as the dependent variable and the 3 identified components as independent variables showed an overall significant association of increasing fatigue with a correlation coefficient of 0.39 (p value < 0.001). For the first and third component the correlation coefficients were 0.73 and 0.35, respectively, with statistically significant p value < 0.001. For the second component, the regression coefficient was 0.06, with a p value of 0.45. In the sensitivity analysis, the PCA reduced the clinical co-variables to 3 major components explaining 64% of experienced fatigue (Supplementary Figure 3, available with the online version of this article). The components identified that included VAS pain in the analysis were almost identical to the components identified that included PDQ score. Comparing PCA performed on male versus female also resulted in similar components explaining 68% and 61% of experienced fatigue, respectively, though with a difference from the primary PCA in the inflammatory component; 36% in males and 29% in females (Supplementary Figure 4).

**DISCUSSION**

The median fatigue score in this population-based PsA cohort including patients treated with csDMARD and bDMARD was ≥ 57 mm VAS, underscoring the great importance of fatigue as a patient-reported disease manifestation. Our findings from the PCA in the population with fatigue above

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**Table 1. Patient characteristics.**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Fatigue: None to Mild, VAS Score &lt; 57, n = 520</th>
<th>Fatigue: Moderate to Severe, VAS Score ≥ 57, n = 542</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female, n (%)</td>
<td>253 (48.7)</td>
<td>358 (66.1)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Age, yrs</td>
<td>53.0 (44.0–62.0)</td>
<td>52.0 (42.8–60.0)</td>
<td>0.070</td>
</tr>
<tr>
<td>Disease duration, yrs</td>
<td>6.0 (3.0–11.5)</td>
<td>5.0 (2.0–10.0)</td>
<td>0.022</td>
</tr>
<tr>
<td>Previous use of DMARD, n (%):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>449 (86.3)</td>
<td>443 (81.7)</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>44 (8.5)</td>
<td>50 (9.2)</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>26 (5.0)</td>
<td>49 (9.0)</td>
<td></td>
</tr>
<tr>
<td>3+</td>
<td>1 (0.1)</td>
<td>0 (0.0)</td>
<td></td>
</tr>
<tr>
<td>Use of MTX, n (%)</td>
<td>316 (60.8)</td>
<td>313 (57.7)</td>
<td>0.319</td>
</tr>
<tr>
<td>Concomitant corticosteroid, n (%)</td>
<td>6 (1.2)</td>
<td>29 (5.4)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Biological treatment, status, n (%)</td>
<td>520</td>
<td>542</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Never treated with biologicals</td>
<td>272 (52.3)</td>
<td>279 (51.5)</td>
<td></td>
</tr>
<tr>
<td>In current treatment</td>
<td>224 (43.1)</td>
<td>195 (36.0)</td>
<td></td>
</tr>
<tr>
<td>Previous use</td>
<td>24 (4.6)</td>
<td>68 (12.5)</td>
<td></td>
</tr>
<tr>
<td>SJC, 0–28*</td>
<td>0.47 ± 1.3</td>
<td>0.94 ± 2.2</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>TJC, 0–28*</td>
<td>1.73 ± 3.6</td>
<td>5.0 ± 6.4</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>CRP, mg/l</td>
<td>3.0 (1.0–6.0)</td>
<td>4.0 (2.0–7.0)</td>
<td>0.008</td>
</tr>
<tr>
<td>Patient pain assessment, 0–100 mm VAS</td>
<td>25.0 (15.0–38.0)</td>
<td>66.0 (49.0–78.0)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PtPGA, 0–100 mm VAS</td>
<td>27.0 (15.0–43.0)</td>
<td>75.5 (61.0–86.0)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PGA, 0–100 mm VAS</td>
<td>7.0 (3.0–15.0)</td>
<td>14.0 (7.0–14.0)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PDQ score</td>
<td>9.0 (6.0–14.0)</td>
<td>17.0 (13.0–23.0)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>DAS28-CRP</td>
<td>2.3 (1.8–2.9)</td>
<td>3.5 (2.6–4.4)</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>HAQ score, 0–3</td>
<td>0.4 (0.1–0.8)</td>
<td>1.1 (0.8–1.6)</td>
<td>&lt; 0.001</td>
</tr>
</tbody>
</table>

Unless otherwise stated, data were given as median or interquartile range. * SJC and TJC given as mean ± SD. VAS: visual analog scale; DMARD: disease-modifying antirheumatic drugs; MTX: methotrexate; PDQ: Pain Detect Questionnaire; CRP: C-reactive protein; DAS28-CRP: 28-joint count Disease Activity Score using CRP; HAQ: Health Assessment Questionnaire; SJC: swollen joint count; TJC: tender joint count; PtGA: patient’s global assessment; PGA: physician’s global assessment.
the median suggested that fatigue was constituted by an inflammatory component, disease duration, and chronic pain in the absence of inflammation. Moreover, the multiple linear regression analysis showed that there was a significant and clinically relevant association with the 3 components and increasing fatigue in the entire population.

Conducting the PCA led to 3 components that affected and explained 63% of experienced moderate to severe fatigue in patients with PsA. The first component was driven by clinical inflammatory factors such as DAS28-CRP, PGA, and swollen and tender joints, revealing one of the underlying explanations of fatigue to be actual inflammatory disease activity – highlighting the importance of targeted treatment of PsA. The second component consisted of disease duration and age, leading our attention to the important aspect of a link between fatigue and disease chronicity. The third component was defined by an inverse relationship between low CRP and high pain indicators. High PDQ scores in the moderate to severe fatigue group suggested central pain sensitization, though the contribution from tender joints to the third component might be explained by a degree of structural damage as well. When substituting PDQ scores with VAS pain, the same components were identified, underscoring the experienced pain as an important driver of fatigue independent of cause or origin for the pain. PDQ scores were in general higher in patients with moderate to severe fatigue, implying a higher degree of centrally derived pain in this group. Chronic pain conditions are common within rheumatic diseases and this further indicates the importance of differentiating patients to provide them with the best possible care.

Previous studies showed that bDMARD and targeted treatments improved symptoms of fatigue in patients with PsA compared to placebo-controlled groups, indicating an inflammatory component in the type of fatigue also found in our present study. From the percentages experiencing no change in fatigue, one could consider whether this to a degree is treatment-refractory because of other components influencing experienced fatigue.

In line with previous research, our present study found that the moderate to severe fatigue group consisted of a statistically significant higher number of females, had higher pain scores, and higher HAQ scores. Additionally, the present study also found that concomitant use of corticosteroids and patients more often switching bDMARD were associated with having moderate to severe fatigue.

Limitations of our current study were (1) the incompleteness of baseline data; however, the proportion of missing data did not exceed 25% for any variable, and (2) the risk for

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**Figure 1.** Principal component analysis indicating 3 components explaining fatigue. The 3 components explaining fatigue include (1) clinical inflammatory manifestations, (2) chronicity, and (3) chronic pain. * High-impact variables contributing to the component. Each variable is presented with the corresponding loading factor. VAS: visual analog scale; CRP: C-reactive protein; PDQ: Pain Detect Questionnaire.
selection bias of the patients because recording of data in DANBIO was mandatory only for patients treated with bDMARD, which may lead to overrepresentation of patients with more severe disease who were taking highly effective therapies. Nonetheless, pain and fatigue remain of utmost importance to patients, and the current study offers new insights into the mechanisms leading to fatigue.

Our study showed a strong association between fatigue and clinically important features including inflammation, disease duration, and chronic pain, which are relevant to take into account when treating PsA. The 3 components explained in total 63% of the experienced fatigue in the moderate to severe fatigue population of patients with PsA.

ACKNOWLEDGMENT

We acknowledge our patient research partner for taking part in the process of preparing the study through discussions of fatigue in relation to PsA and for reading and commenting on the final article.

ONLINE SUPPLEMENT

Supplementary material accompanies the online version of this article.

REFERENCES