Research Article

Pain Sensitisation in Women with Active Rheumatoid Arthritis: A Comparative Cross-Sectional Study

Nora Vladimirova,1 Anders Jespersen,1 Else Marie Bartels,1 Anton W. Christensen,1 Henning Bliddal,1,2 and Bente Danneskiold-Samsøe1,2

1The Parker Institute, Department of Rheumatology, Copenhagen University Hospital, Bispebjerg and Frederiksberg, 2000 Frederiksberg, Denmark
2The Health Science Faculty, University of Copenhagen, 2200 Copenhagen N, Denmark

Correspondence should be addressed to Else Marie Bartels; else.marie.bartels@regionh.dk

Received 9 March 2015; Revised 30 June 2015; Accepted 2 July 2015

Academic Editor: Burkhard Leeb

Copyright © 2015 Nora Vladimirova et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Objectives. In some rheumatoid arthritis (RA) patients, joint pain persists without signs of inflammation. This indicates that central pain sensitisation may play a role in the generation of chronic pain in a subgroup of RA. Our aim was to assess the degree of peripheral and central pain sensitisation in women with active RA compared to healthy controls (HC).

Methods. 38 women with active RA (DAS28 > 2.6) and 38 female HC were included in, and completed, the study. Exclusion criteria were polyneuropathy, pregnancy, and no Danish language. Cuff Pressure Algometry measurements were carried out on the dominant lower leg. Pain threshold, pain tolerance, and pain sensitivity during tonic painful stimulation were recorded.

Results. Women with active RA had significantly lower pain threshold (\(p < 0.01\)) and pain tolerance (\(p < 0.01\)) than HC. The mean temporal summation- (TS-) index in RA patients was 0.98 (SEM: 0.09) and 0.71 (SEM: 0.04) in HC (\(p < 0.01\)).

Conclusion. Patients with active RA showed decreased pressure-pain threshold compared to HC. In addition, temporal summation of pressure-pain was increased, indicating central pain sensitisation, at least in some patients. Defining this subgroup of patients may be of importance when considering treatment strategies.

1. Introduction

Rheumatoid arthritis (RA) is a chronic inflammatory disease causing synovial joint destruction, disability, and pain [1]. Despite better disease control, many RA patients rate their pain as a major burden, affecting their quality of life [2]. Although physicians often assume that inflammation is the main determinant of pain intensity, more than one in 10 of the patients in DAS28 remission [3] still report clinically significant pain levels [4]. The fact that pain persists despite absence of signs of inflammation may therefore indicate that other factors in the pain processing, for example, sensitisation, either peripherally or centrally, play an important role in the generation of pain in RA [5].

Central pain sensitisation is defined as “an increased responsiveness of nociceptive neurons in the central nervous system to their normal or subthreshold afferent input” [6]. Clinically, this results in hyperalgesia and allodynia and receptive field expansion [7]. It is plausible that subgroups of RA patients experience alterations in their central pain modulation which then becomes a primary cause of pain, possibly outlasting the inflammatory activity. In this subgroup, central pain sensitisation may contribute significantly to high tender joint counts and self-reporting of pain, causing overestimation of disease activity [8]. In RA patients, concomitant fibromyalgia has been found to be around 13–17% [9].

Pain in RA is usually assessed either by self-reported questionnaires or by simple palpation of the joints. Computerized Cuff Pressure Algometry (CPA) is a more precise method for quantitative pain testing which primarily assesses sensitivity of muscle and deep tissue. CPA is less influenced
by inter- and intraexaminer bias than handheld pressure algometry [10]. CPA is used to assess different aspects of pain in various rheumatic conditions [11–13]. Pain assessed by CPA has in a group of fibromyalgia shown to correlate to number of tender points, thereby creating a new approach to pain assessment in chronic pain patients [14].

Based on the hypothesis that some RA patients with a high DAS28 suffer from central pain sensitisation, our aim was to use CPA to study pain properties in patients with active RA. The main outcomes of interest were pain threshold and pain tolerance, indicating the degree of peripheral pain sensitivity and degree of temporal summation (i.e., central pain sensitisation) assessed by tonic painful cuff pressure stimulation [13].

2. Methods

2.1. Study Type and Ethics. This clinical study was cross-sectional and noninterventional carried out according to the Helsinki Declaration and accepted by the Capital Region of Denmark’s Ethical Committee (KF01-058/02). All assessments were performed the same day and in the same order by the same persons. All participants gave informed written consent to participate.

2.2. Participants. Patients were recruited from the Rheumatology Department, Bispebjerg and Frederiksberg Hospital, Denmark, from June 2011 to September 2012, and were screened according to the following inclusion criteria:

(i) age ≥ 18;
(ii) RA diagnosed by a rheumatologist according to the ACR criteria [1];
(iii) active RA according to the EULAR definition, DAS28 ≥ 2.6 [3].

Exclusion criteria were polyneuropathy, pregnancy, and patients not able to understand Danish or to follow instructions. Figure 1 shows the selection process of the patients enrolled in this study. Healthy controls (HC) were assessed at the clinic at an earlier study to provide data on healthy subjects, and the data was taken directly from this study [13]. The control subjects were all assessed at entry to be healthy, and the protocol for the pain measurements was the same as in the present study.

2.3. General Clinical Examinations. The following parameters were measured: A 28-joint count [3], plasma C-reactive protein (CRP) (measured at inclusion in heparin plasma with immunoturbidimetric absorption photometry (Roche/Hitachi cobas-C systems, Roche Diagnostics GmbH, D-68298 Mannheim), with a value ≤10 mg/L being considered normal concentration, and a detection limit of 0.3 mg/L), DAS28, Disease Activity Score in 28 joints based on CRP [3], and manual 18-tender points count [15]. Patient-reported outcomes were registered via computerized Health Assessment Questionnaire [16].

2.4. Cuff Pressure Algometry. The device consisted of a 61 cm tourniquet cuff (placed on the widest part of the lower leg on subject’s dominant side), a computer-controlled air compressor, and an electronic 10 cm visual analogue scale (VAS) (DoloCuff, Unique Electronics, Denmark) (Figure 2). The subject was instructed to rate the pain intensity continuously on the VAS, from the first sensation of pain after start of inflation and to press the pressure-release button at “the pain intensity strong enough to make one feel like interrupting or stopping it.” Based on a mean of three consecutive recordings the following variables were determined: (1) pain threshold (unit: kPa), defined as the moment of transition between strong and painful pressure (the first time the VAS exceeds 0) and (2) pain tolerance (unit: kPa), the pressure value at the termination of pressure inflation. Following this, degree of temporal summation was determined according to a predefined protocol [12, 13]. The cuff was filled to obtain a pain intensity based on the average pressure-pain threshold and pressure tolerance of the individual subject. This individualized stimulation was maintained for 10 minutes, while the subject continuously reported pain intensity on the electronic VAS. Based on highest VAS-pain rating, VAS-pain rating at end of stimulation, and Effective Stimulation Time, an index (TS-index) describing degree of temporal summation was calculated according to the following formula:

\[
TS\text{-index} = \log \left( \frac{\text{VAS}_{\text{end}}}{\text{VAS}_{\text{max}}} \times \frac{10}{\text{TIME}_{\text{stimulation}}} \times \text{VAS}_{\text{end}} \right)
\]
Figure 2: Cuff Pressure Algometry (CPA). The device consisted of a 61 cm tourniquet cuff, a computer-controlled air compressor, and an electronic 10 cm visual analogue scale (VAS). The cuff was placed on the widest part of the lower leg (not in relation to the joints). The measurements were carried out on the subject’s dominant side.

The TS-index increases with increasing degrees of temporal summation and expresses central pain sensitisation. VAS is measured on the 0–10 scale, and time is measured in minutes.

2.5. Statistical Methods. For data analyses, SAS software (v. 9.1.3 Service Pack 4; SAS Institute Inc., Cary, NC, USA) was applied. In order to evaluate the empirical distributions of the continuous outcomes, we tested for normal distribution looking at skewness/SE_{skew} and kurtosis/SE_{kurt}.

The PROC MEANS statement was used for summarizing the data. T-TEST procedure was used to compare the two groups, analysed as two independent samples. The CORR procedure, based on Spearman’s rank-order correlation, was used as a nonparametric measure of association between various variables.

3. Results

The median age of patients was 56 years (IQR 46–69) and 39 (IQR 32–44) of the controls. Characteristics of RA patients are shown in Table 1.

3.1. Cuff Algometry Data. Data from Cuff algometry measurements are shown in Table 2.

3.2. Clinical Features of the Patients. All RA patients had active disease at the time of assessment, with median DAS28CRP of 4.3 (IQR 3.6–5.0). The median number of swollen joints was 5 (IQR 3–11) and the number of tender joints was 7 (IQR 4–13). The disease duration varied from 4 to 444 months with a median of 33 months.

3.3. Pressure-Pain Thresholds and Tolerance. The mean pressure-pain threshold and pressure-pain tolerance within the RA group were significantly decreased (p < 0.01) compared with the healthy controls (Table 1).

3.4. Temporal Summation of Pain (TS-Index). The maximal value for the TS-index was 1.09 in HC; 11 out of 38 (29%) in RA group were significantly decreased (p < 0.01) compared with the healthy controls (Table 1).

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Median (interquartile range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>56 (46, 69)</td>
</tr>
<tr>
<td>Disease duration, months</td>
<td>33 (29, 45)</td>
</tr>
<tr>
<td>IgM RF seropositive, number (%)</td>
<td>23 (61)</td>
</tr>
<tr>
<td>Anti CCP-positive, number (%) (n = 36)</td>
<td>21 (58)</td>
</tr>
<tr>
<td>Bone erosions on X-Ray, number (%) (n = 37)</td>
<td>12 (32)</td>
</tr>
<tr>
<td>Clinical assessment of disease activity</td>
<td></td>
</tr>
<tr>
<td>DAS28</td>
<td>4.3 (3.6, 5.0)</td>
</tr>
<tr>
<td>Tender joints</td>
<td>7 (4, 13)</td>
</tr>
<tr>
<td>Swollen joints</td>
<td>5 (3, 11)</td>
</tr>
<tr>
<td>CRP mg/L</td>
<td>5.5 (0.5, 14)</td>
</tr>
<tr>
<td>VAS Global</td>
<td>50 (29, 64)</td>
</tr>
<tr>
<td>HAQ-score (n = 37)</td>
<td>0.88 (0.63, 1.75)</td>
</tr>
<tr>
<td>Tender points</td>
<td>14 (8, 18)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Medication</th>
<th>Numbers (% of total participating patient group)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients on DMARDs</td>
<td>34 (89)</td>
</tr>
<tr>
<td>Patients on DMARDs + biologics</td>
<td>14 (37)</td>
</tr>
<tr>
<td>Concomitant per oral prednisolone</td>
<td>9 (24)</td>
</tr>
<tr>
<td>Treatment with 80 mg i.m./i.art steroid within the last 4 weeks</td>
<td>5 (13)</td>
</tr>
</tbody>
</table>

RF: rheumatoid factor, anti-CCP: anticyclic citrullinated peptide, DAS 28: Disease Activity Score, VAS: visual analog scale, VAS Global: patient global assessment of disease severity as measured on a visual analog scale, DMARDs: disease-modifying antirheumatic drugs, i.m./i.art: intramuscular or intra-articular steroid injections.

Table 1: Patient characteristics and medication for the 38 participating rheumatoid arthritis patients.

Table 2: Cuff algometry parameters of patients and healthy control subjects.
In addition, post hoc analyses indicated that the degree of temporal summation was more pronounced in some of the RA patients, where 29% had a TS-index above the upper limit of that of HC. This suggests that degree of central sensitisation is more pronounced in a subgroup of RA patients [20].

4. Discussion

Earlier studies have reported concomitant fibromyalgia in a subpopulation of RA patients [9, 17, 18], but this is the first study investigating temporal summation as an indicator of central pain sensitization in RA patients with active disease according to having a DAS28 > 2.6. The study demonstrated that women with active RA (DAS28 > 2.6) showed a significantly decreased pressure-pain thresholds and tolerance on the muscles of the lower leg when compared to HC, indicating a higher degree of peripheral pain sensitivity [19]. Furthermore, temporal summation of pain characterized by the TS-index was significantly higher in RA patients compared with HC. These findings confirm that, as a group, RA patients suffer from nonarticular pain hypersensitivity and facilitated temporal summation, indicating central pain sensitisation [7].

In addition, post hoc analyses indicated that the degree of temporal summation was more pronounced in some of the RA patients, where 29% had a TS-index above the upper limit of that of HC. This suggests that degree of central sensitisation is more pronounced in a subgroup of RA patients [20].

Computerized Cuff Pressure Algometry has previously been applied in other chronic rheumatic pain conditions [12, 13, 21]. There are no previous reports concerning temporal summation assessed by CPA among RA patients but the present results suggest that CPA may be of value in the identification of RA patients with abnormal pain sensitisation.

In RA, pain is the major determinant of patient global assessment of disease activity [22]. In subjects with increased central pain sensitisation this may lead to increased pain reports and overestimates of disease activity [4, 8]. This would also in part explain the significant proportion of RA patients who do not respond to treatment with biologics, since these drugs target proinflammatory pathways [23]. This study indicates that degree of central sensitisation should be evaluated when planning a strategy for medical therapy of the individual RA patient. Interventions against such pain mechanisms may be supplementary, or even alternative to the interventions aimed at decreasing inflammation [24].

4.1. Limitations. Due to limited sample size, subgroup analyses were not possible. Our results, however, point towards a necessity of a large scale study assessing the relationship between treatment response and underlying pain mechanisms. Since this study was on women only, we cannot generalize our results to the pain condition in men with RA.

5. Conclusion

In patients with active RA, signs of central pain sensitisation were found as indicated by a higher TS-index compared to the group of healthy controls. This indicates that assessment of pain and the underlying mechanisms may be of importance when considering treatment strategies in the individual subjects with RA.

Conflict of Interests

The authors declare that there is no conflict of interests.

Authors’ Contribution

The first author, Nora Vladimirova, has written the paper in collaboration with the other authors, Anders Jespersen, Else Marie Bartels, Anton W. Christensen, Henning Bliddal, and Bente Danneskiold-Samsøe. The study design was made by Nora Vladimirova, Anders Jespersen, Else Marie Bartels, Henning Bliddal, and Bente Danneskiold-Samsøe. The study and the patient contacts have been conducted by Nora Vladimirova. The data handling was carried out by Nora Vladimirova, Anders Jespersen, Else Marie Bartels, and Anton W. Christensen. The paper was accepted by all authors prior to submission.

Acknowledgments

The study was supported by the OAK Foundation (OCAY-13-309). The authors wish to thank laboratory technician Jack Kvistgaard Olsen for help with the pain measurements.
References


