EXTENDED REPORT

Development and psychometric validation of a patient-reported outcome measure to assess fears in rheumatoid arthritis and axial spondyloarthritis: the Fear Assessment in Inflammatory Rheumatic diseases (FAIR) questionnaire

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ABSTRACT

Objectives To develop and validate an outcome measure for assessing fears in patients with rheumatoid arthritis (RA) and axial spondyloarthritis (axSpA).

Methods Fears were identified in a qualitative study, and reformulated as assertions with which participants could rate their agreement (on a 0–10 numeric rating scale). A cross-sectional validation study was performed including patients diagnosed with RA or axSpA. Redundant items (correlation >0.65) were excluded. Internal consistency (Cronbach’s α) and factorial structure (principal component analysis) were assessed. Patients were classified into fear levels (cluster analysis). Associations between patient variables and fear levels were evaluated using multiple logistic regression.

Results 672 patients were included in the validation study (432 RA, 240 axSpA); most had moderate disease activity and were prescribed biologics. The final questionnaire included 10 questions with high internal consistency (α: 0.89) and a single dimension. Mean scores (±SD) were 51.2 (±25.4) in RA and 60.5 (±22.9) in axSpA. Groups of patients with high (17.2%), moderate (41.1%) and low (41.7%) fear scores were identified. High fear scores were associated with high Arthritis Helplessness Index scores (OR 6.85, 95% CI (3.95 to 11.87)); high Hospital Anxiety and Depression Scale anxiety (OR 5.80, 95% CI (1.19 to 4.22)) and depression (OR 2.37, 95% CI (1.29 to 4.37)); low education level (OR 3.48, 95% CI (1.37 to 8.83)); and high perceived disease activity (OR 2.36, 95% CI (1.10 to 5.04)).

Conclusions Overall, 17.2% of patients had high fear scores, although disease was often well controlled. High fear scores were associated with psychological distress. This questionnaire could be useful both in routine practice and clinical trials.

INTRODUCTION

Chronic inflammatory rheumatic diseases (CIRDs) such as rheumatoid arthritis (RA) and axial spondyloarthritis (axSpA) have a major impact on quality of life.1 They interfere with many aspects of daily functioning, including recreational activities, work, family life and relationships.2 These aspects of disease burden are frequently underestimated or unrecognised by the patient’s family and friends, as well as the treating physician.2,3 In addition, these diseases may be associated with considerable psychological distress, including anxiety or depression.4–6 Several studies, including a recent qualitative study in France,7 have shown that patients with CIRDs have specific fears about how their disease will progress, limitations in daily activities, being a burden on others and treatment.4–9 Although these aspects are important to patients, they are currently difficult to assess due to the lack of a specific evaluation tool.

Although several patient-reported outcome (PRO) measures assess emotional status or anxiety levels,10–13 many of these are generic and none, to our knowledge, specifically assess fears.14 A questionnaire focusing on CIRD-related fears would potentially be useful both in the context of everyday care (eg, to help understand patients’ motivations and reluctance towards treatments) and in clinical trials, since such fears may have an impact on the efficacy of a study drug.15 Current recommendations on PRO development and validation include grounding PROs in the patient perspective, and performing adequate psychometric validation of all such measures.16–20

The objectives of the present study were to develop a PRO for fear assessment in patients with RA or axSpA, and to perform a preliminary psychometric validation of the resulting instrument.

METHODS

This study was part of a larger programme of research on patient perceptions in chronic progressive rheumatic diseases. The programme was supervised by a steering committee (the authors of this manuscript), composed of rheumatologists, psychologists, methodologists and representatives of the scientific staff of the Arthritis Fondation Courtin and of UCB Pharma, who jointly funded the programme.

Development of a preliminary questionnaire

In a previously published qualitative study,7 25 patients with RA and 25 with axSpA participated...
in semistructured interviews about their perceptions of the diseases. Interviews were transcribed verbatim, and the data extracted inductively from the interview transcripts. Fears about the future course of the disease, the impact of disease and its treatment were frequently expressed, and appeared to be shared in common between patients with axSpA and those with RA.

In the present study, all fears that were expressed by >5% of patients in the qualitative study were used. Non-redundant statements were then rephrased as assertions over two working sessions involving members of the Steering Committee and a patient research partner (a member of the EULAR PARE (People with Arthritis and Rheumatism) programme). The agreement with each item was assessed on a scale of 0–10 (‘totally disagree’ to ‘totally agree’). The questions were then tested in a sample of 10 patients with RA and 10 with axSpA for linguistic validation, and cognitive debriefing was performed during individual face-to-face interviews with trained interviewers. This preliminary questionnaire contained 23 items related to fears.

Validation study
This was a prospective, cross-sectional study in patients with RA or axSpA in France. Participants were recruited by hospital and community rheumatologists between July 2014 and October 2015.

Participants
All rheumatologists currently practising in France were invited to participate in the study through post and email. Each participating rheumatologist was expected to invite up to 20 consecutive patients with RA or axSpA attending a routine consultation who were aged >18 years, and had a diagnosis of RA according to the ACR/EULAR (American College of Rheumatology/European League Against Rheumatism) classification criteria, or of axSpA according to the ASAS (Assessment in Spondyloarthritis International Society) classification criteria. Patients with psoriatic arthritis or other CIRDs, and those who were unable to complete a questionnaire, were excluded.

Data collection
Patients were asked to complete the preliminary questionnaire, as well as the patient global assessment of overall disease activity (scored from 0 to 10), the Hospital Anxiety and Depression Scale (HADS), the Arthritis Helplessness Index (AHI) and, for patients with axSpA, the Bath Ankylosing Spondylitis Disease Activity Index (BASDAI). The patient also provided information on sociodemographic indicators, health insurance coverage, disease duration and family history of rheumatic disease. In parallel, the rheumatologist provided information on the patient’s disease activity, as measured by the 28-item Disease Activity Score calculated with erythrocyte sedimentation rate (DAS28(ESR)) for RA, and an overall assessment of disease activity scored from 0 to 10. Information on current treatment was also collected.

In order to assess the reproducibility of the questionnaire, 30 randomly selected patients were provided with two questionnaires and invited to complete and return the second one 2 weeks later.

Finalisation and psychometric validation of the Fear Assessment in Inflammatory Rheumatic diseases questionnaire

Finalising the questionnaire
The number of items on the fear dimensions was reduced to avoid redundancy. Interitem Pearson’s correlation coefficients between each pair of items were determined across the entire data set, and pairs presenting an r>0.65 were considered redundant. In such cases, the item considered most clear in wording by the Steering Committee was retained. In addition, items only relevant to a subgroup of patients, such as those relating to pregnancy (only applicable to women of childbearing age) or to professional activity (only applicable to people in work), were eliminated. The finalised questionnaire was translated from French into English through two independent forward and backward translations and reconciliation of the translated texts.

Preliminary validation
All patients for whom both patient and physician questionnaires had been received were considered. Missing values were replaced according to a Missing at Random hypothesis. When the proportion of missing data was <5%, individual missing items were replaced with the median value of the corresponding

| Table 1 Patient characteristics |
|--------------------------------|
| RA (n=432) | axSpA (n=240) | Total (n=672) |
| Age (years) | n=368 | n=207 | n=575 |
| Gender | | | |
| Male | n=373 | n=208 | n=581 |
| Female | 276 (74.0%) | 94 (45.2%) | 370 (63.7%) |
| Male | 97 (26.0%) | 114 (54.8%) | 211 (36.3%) |
| Professional activity | n=424 | n=237 | n=661 |
| In employment | 162 (38.2%) | 167 (70.5%) | 329 (49.8%) |
| Student | 2 (0.5%) | 1 (0.4%) | 3 (0.5%) |
| Unemployed | 8 (1.9%) | 19 (8.0%) | 27 (4.1%) |
| Retired | 201 (47.4%) | 30 (12.7%) | 231 (34.9%) |
| Others | 51 (12.0%) | 20 (8.4%) | 71 (10.7%) |
| Disease duration (years) | n=358 | n=203 | n=561 |
| Disease activity | n=427 | n=236 |
| DAS28 | 2.6±1.2 | – | – |
| BASDAI | – | 3.3±2.2 | – |
| Physician global assessment of disease activity (NRS) | n=419 | n=232 | n=651 |
| Patient global assessment of disease activity (NRS) | n=382 | n=216 | n=598 |
| Treatments | n=326 | n=238 | n=564 |
| None | 5 (1.5%) | 7 (2.9%) | 12 (2.1%) |
| Corticosteroids alone | 6 (1.1%) | – | 6 (1.1%) |
| NSAIDs alone | – | 36 (15.1%) | 36 (6.4%) |
| Synthetic DMARDs ± corticosteroids | 61 (18.7%) | – | 61 (10.8%) |
| Synthetic DMARDs ± NSAIDs | – | 15 (6.3%) | 15 (2.7%) |
| Biological DMARDs (alone or in combination) | 252 (77.3%) | 173 (72.7%) | 425 (75.4%) |
| Others | 2 (0.6%) | 7 (2.9%) | 9 (0.7%) |

Data are presented as mean values±SD for continuous variables, and as frequency counts (%) for categorical variables.
variable. When the proportion exceeded 5%, multiple imputation methods based on Markov chains and Monte Carlo simulations were used. Score distribution was assessed using mean±SD and median with IQR scores for each disease population.

The factorial structure of the questionnaire was determined using principal component analysis, and eigenvalues calculated. A confirmatory factor analysis was then performed to determine goodness of fit, restricted to dimensions with eigenvalue > 1. The internal coherence was assessed with Cronbach’s α coefficient.

Test–retest stability of the PRO was evaluated by determining the Pearson’s correlation coefficient for total scores between two questionnaires completed at 2 weeks’ interval by 30 respondents. Coefficient > 0.70 were considered to represent a strong correlation, and coefficients 0.50–0.70, a moderate correlation. The discriminative validity of the PRO was assessed by evaluating the relationship between the scores and other study variables expected to be related to the PRO score, such as HADS anxiety score, helplessness (AHI score) or disease activity score. Anxiety/depression and helplessness were expected to be moderately to strongly correlated with fears, whereas disease activity was expected to be only moderately correlated.

Identification of patient clusters and characteristics associated with fears

Subgroups of patients were identified according to their fear scores using descending cluster analysis (Ward method). Optimal thresholds to distinguish between high and low fear score clusters were identified using receiver operating characteristic (ROC) curves based on the Youden index (optimal sensitivity and specificity).

Univariate, then multivariate logistic regression was used to identify patient variables (including demographic, social and economic characteristics; disease status, and anxiety/depression and helplessness levels) independently associated with the highest compared with the lowest fear score cluster. Variables identified in the univariate analysis (p < 0.20) were entered into a backward stepwise multiple logistic regression model.

All statistical analyses were performed using SAS V9.2.

Ethics

The study was performed in accordance with Good Epidemiological Practice and relevant French guidelines for patient surveys. Verbal informed consent was obtained from all participating patients.

RESULTS

Participants

All 1618 rheumatologists in France were contacted: 134 agreed to participate in the study, and 100 enrolled at least 1 patient. Twenty were exclusively community based, 51 exclusively hospital based and the remaining 29 had a mixed practice. A total of 796 patients were enrolled, of whom 672 (84.4%) were retained for analysis (see online supplementary figure 1). Patient characteristics are presented in table 1. Disease was moderately active, and use of biologics exceeded 70% in both the RA and axSpA patient populations.

Finalisation of the Fear Assessment in Inflammatory Rheumatic diseases questionnaire

Factorial analysis of the initial 44-item questionnaire (which dealt with both fears and opinions) revealed two highly correlated fear clusters. Subsequently, clusters were extracted and a 4-item questionnaire was developed and tested on 240 patients (100 from each disease population). A confirmatory factor analysis was then performed to determine goodness of fit, restricted to dimensions with eigenvalue > 1. The internal coherence was assessed with Cronbach’s α coefficient.

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correlated dimensions related to fears: one to disease outcome, and the other to treatment. After exclusion of redundant items with interitem correlation coefficients >0.65 (see online supplementary table 1), the final scale comprised 10 items (table 2 and online supplementary tables 2 and 3). Each item is scored on a 10-point numerical rating scale ranging from 0 (no fear) to 10 (strong fear). The total score ranges from 0 to 100 and was calculated as the sum of the 10 individual item scores.

**Psychometric validation**

Internal consistency was high (Cronbach’s α coefficient: 0.89). Principal component analysis identified a single dimension (eigenvalue: 5.1), which accounted for 51.2% of variance in the item scores. Confirmatory factor analysis matching the data to a unidimensional factorial structure revealed a goodness-of-fit index of 0.91. Twenty-eight patients (13 RA and 15 axSpA) provided two questionnaires completed 2 weeks apart. The test–retest correlation coefficient was ≥0.81. Total FAIR (Fear Assessment in Inflammatory Rheumatic diseases) scores were correlated with HADS anxiety (r=0.47; p<0.001) and depression (r=0.40; p<0.001) scores, and with AHI scores (r=0.50; p<0.001) (see online supplementary figure 2).

**Distribution of scores in patients with RA and axSpA**

The mean and median FAIR scores were 54.9±24.9 and 57 (IQR: 35–75), respectively. Scores were higher in patients with axSpA (60.5±22.9; 65 (43–79)) than in patients with RA (51.8±25.4; 52 (33–71)). The distribution of PRO scores for the full data set is presented in figure 1. The mean item scores on the FAIR scale are presented in table 2 for the total study population, for patients with RA and for patients with axSpA. Mean fear scores were consistently higher for all items in patients with axSpA compared with those with RA.

**Subgroups of patients**

Hierarchical cluster analysis identified three groups of patients characterised by high (cluster 1; n=116; 17.2%; mean score 87.0±7.9), moderate (cluster 2; n=276; 41.1%; mean score 65.8±11.4) and low levels of fear (cluster 3; n=280; 41.7%; mean score 31.1±14.7) (figure 2). These three clusters accounted for 68.3% of the variance in the data set. The most discriminating cut-off threshold to distinguish the high fear cluster from the other two was 77 (sensitivity: 0.90; specificity: 0.91). The most sensitive cut-off threshold to distinguish the low fear cluster from the other two was 51 (sensitivity: 0.92; specificity: 0.93). The area under the ROC curve was >0.97 in both cases (see online supplementary figure 3).

Multiple logistic regression analysis was used to determine patient characteristics independently associated with high fear scores, discriminating between patients in cluster 1 and those in cluster 3 (figure 3). Cluster 1 (high fear scores) was associated with higher global rating of disease activity by the patient, high AHI helplessness scores and high HADS anxiety and depression scores. With respect to sociodemographic variables, low education level, not working and living alone were also associated with higher FAIR score, as was immigrant status. No significant effects of disease type (axSpA vs RA) or age were observed. With respect to the patients in cluster 2 (moderate fear scores), the same variables were identified, although the ORs were lower.

**DISCUSSION**

This large national survey of patients with RA or axSpA generated two principal results. First, almost one-fifth (17.2%) of evaluated patients had high fear scores, despite both diseases being typically well managed, and these scores were associated with psychological distress. Thus, the fears identified in this study may reflect psychological distress, and need to be addressed even in patients who have moderate to low disease activity. Second, we have developed the FAIR questionnaire: a disease-specific,
Factors associated with fear scores (multiple logistic regression)

| Factor                                | OR     | 95% CI   |
|---------------------------------------|--------|----------|
| AHI Score >20 vs <20                   |        |          |
| HADS Anxiety Score 8-10 vs <8         |        |          |
| HADS Depression score ≥8              |        |          |
| Disease activity evaluated by the patient |       |          |
| 30-60 vs ≤30                          |        |          |
| >60 vs ≤30                            |        |          |
| Place of birth                         |        |          |
| Foreign vs France                     |        |          |
| Education level                       |        |          |
| Primary vs tertiary                   |        |          |
| Secondary vs tertiary                 |        |          |
| Professional activity                 |        |          |
| In employment vs retired              |        |          |
| Unemployed vs retired                 |        |          |
| Entourage                             |        |          |
| Living alone vs with others           |        |          |

Figure 3 Variables independently associated with a high or moderate FAIR (Fear Assessment in Inflammatory Rheumatic diseases) score versus a low score in patients with rheumatoid arthritis (RA) and axial spondyloarthritis (axSpA). High scores indicate those >75 (shown in red); moderate scores, 50–75 (shown in blue); and low scores, <50. Data are presented as ORs (95% CIs). AHI, Arthritis Helplessness Index; HADS, Hospital Anxiety and Depression Scale.

psychometrically validated PRO to measure disease and treatment-related fears in patients with RA or axSpA. This instrument demonstrated acceptable psychometric properties: notably unidimensionality, high internal coherence, good discriminant validity and adequate test–retest stability. The FAIR is short (10 items), simple to score and may be a useful tool both in routine practice and clinical trials.

The strengths of this study include the size of the study population, the high level of patient involvement in the development of the questionnaire and the psychometric validation of this instrument in line with the recommended guidelines. Limitations include a potential cultural bias, since the items were derived from a qualitative survey of patients in France, and potential redundancy with existing disease-specific PROs for CIRDs. Aspects will need to be evaluated in future studies. Although some questions within the questionnaire may seem redundant, statistical tests were used to remove truly redundant questions, and all questions underwent validation with patients.

In this study, it was possible to classify patients according to their level of fear using the FAIR score. Fear scores did not appear to be related to objective disease activity scores (DAS28(ESR) or BASDAI), although patients with high perceived disease activity (>6) were more frequently classified in the high fear cluster. In contrast, a strong association was observed between FAIR scores and scores on the AHI (≥20) or HADS (≥10 for anxiety and ≥8 for depression), all of which are non-specific markers of psychological distress.

Patients with RA commonly present a higher level of psychological distress compared with the general population. In agreement with this, we observed a robust association between fears and non-specific measures of psychological distress, such as the AHI, the HADS anxiety score and, to a lesser extent, the HADS depression score. Moreover, the fears expressed by our patients are likely to represent specific expressions of psychological distress in inflammatory rheumatic diseases. This would suggest that the FAIR questionnaire could be employed to measure psychological distress in a disease-specific way in patients with RA or axSpA. To this end, it might be beneficial to compare the FAIR questionnaire with existing generic scales, such as the mental component score of the SF-36 or SF-12 (36-Item and 12-Item Short Form Health Survey), or the anxiety and depression items of the Arthritis Impact Measurement Scales, in future studies. The FAIR instrument will also need to be tested in independent populations to verify its robustness and psychometric validity.

An association, although less marked, was also observed between FAIR scores and disease activity as rated by the patient. Four sociodemographic variables were also associated with high fear scores, namely low education level, living alone, being born outside France and either being in or seeking employment. Low education levels may be associated with lower access to, or more limited understanding of, information about the disease; this may also be the case for immigrants. Patients living alone may lack adequate social support for coping with stressful situations, and patients in employment or seeking employment may be particularly worried about the impact of their disease on their future career and income. On the other hand, age, gender, diagnosis (RA or axSpA) and treatment were not independently associated with high fear scores. Previous studies have identified female gender, lack of social support and a lower educational level as being associated with anxiety and depression (or both) in patients with RA.

The FAIR questionnaire may be a useful PRO in several contexts. First, it may be helpful for physicians taking care of patients with RA and axSpA to evaluate the levels of fear and psychological distress in their patients, in order to provide an appropriate level of psychological support and to initiate a physician–patient dialogue to dispel unwarranted fears and facilitate adaptive coping. In clinical research, the questionnaire may be useful for investigating differences in psychological distress between patient groups, and to provide a basis for explaining such differences. Finally, the FAIR could be included in clinical trial protocols to measure the impact of specific interventions on psychological distress; however, this would first require an assessment of the instrument’s sensitivity to change. In this context, a disease-specific PRO might be more sensitive than a non-specific tool such as the HADS.

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Contributors

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Competing interests

LG, PC, AS, CH, CD, ST, FB: no competing interests. GC and J-MJ are employees of UCB Pharma; TdC was an employee of UCB Pharma at the time of the study; VS and FR-M are employees of Fondation Arthritis.

Ethics approval

The Ethics Committee of the St Antoine Hospital, Paris (session of 7 October 2014), the National Advisory Committee on Medical Research Information (CCTIRS) and the French national data protection agency (CNIL).

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