EXTENDED REPORT

A patient-derived and patient-reported outcome measure for assessing psoriatic arthritis: elaboration and preliminary validation of the Psoriatic Arthritis Impact of Disease (PsAID) questionnaire, a 13-country EULAR initiative

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ABSTRACT

Introduction The objective was to develop a questionnaire that can be used to calculate a score reflecting the impact of psoriatic arthritis (PsA) from the patients’ perspective: the PsA Impact of Disease (PsAID) questionnaire.

Methods Twelve patient research partners identified important domains (areas of health); 139 patients prioritised them according to importance. Numeric rating scale (NRS) questions were developed, one for each domain. To combine the domains into a single score, relative weights were determined based on the relative importance given by 474 patients with PsA. An international cross-sectional and longitudinal validation study was performed in 13 countries to examine correlations of the PsAID score with other PsA or generic disease measures. Test–retest reliability and responsiveness (3 months after a treatment change) were examined in two subsets of patients.

Results Two PsAID questionnaires were developed with both physical and psychological domains: one for clinical practice (12 domains of health) and one for clinical trials (nine domains). Pain, fatigue and skin problems had the highest relative importance. The PsAID scores correlated well with patient global assessment (N=474, Spearman r=0.82–0.84), reliability was high in stable patients (N=88, intraclass correlation coefficient=0.94–0.95), and sensitivity to change was also acceptable (N=71, standardised response mean=0.90–0.91).

Conclusions A questionnaire to assess the impact of PsA on patients’ lives has been developed and validated. Two versions of the questionnaire are available, one for clinical practice (PsAID-12) and one for clinical trials (PsAID-9). The PsAID questionnaires should allow better assessment of the patient’s perspective in PsA. Further validation is needed.

INTRODUCTION

Assessments based on patients’ opinion—patient-reported outcome measures (PROMs)—have received increasing recognition as being critically important end points in both clinical trials and long-term observational studies in rhematic diseases during the last decade.1–4 PROMs are elicited directly from the patient and assess how the patient feels or functions with respect to their health condition. These measures reflect the patient burden of disease.5–6

Psoriatic arthritis (PsA) is a heterogeneous disease with high impact on patients’ lives.7–8 However, studies on patient-reported outcomes in PsA have been limited.9–12 PROMs used in PsA clinical trials are mostly generic or adapted from rheumatoid arthritis (RA), and few disease-specific PROMs for PsA are currently available.13–14 The core set of domains of health proposed by physician experts to be included in randomised clinical trials and observational studies of PsA includes the following PROMs: pain, patient global assessment, physical function and health-related quality of life.15 Fatigue is considered here as a research item.15–16 And, indeed, a literature review17 of recently published articles indicated that the only patient-reported outcomes frequently reported were those in the core set, in particular physical function (in 48% of publications, usually using the Health Assessment Questionnaire (HAQ)18), but also pain (47%) and patient global assessment (40%). Fatigue was rarely reported (15%), and psychological aspects were not reported at all.17

However, other dimensions of health may be important from the patients’ perspective.19–20 3–4 PROMs for PsA were an item on the research agenda resulting from the recent development of the European League Against Rheumatism (EULAR) recommendations for the management of PsA.21 Subsequently, EULAR agreed to support the development of a new questionnaire and score to better assess patient-reported outcomes in this disease, in line with the previous development of a similarly focused tool for patients with RA.22–23
The objectives of this study were to elaborate and validate a disease impact questionnaire for PsA based on patients’ experience of the impact of the disease on domains or dimensions of health.

MATERIALS AND METHODS
Elaboration and validation of the PsA Impact of Disease (PsAID) questionnaire were performed in 2011–2012 in three steps (as summarised in online supplementary figure S1), adapting the methodology developed for a similar questionnaire in RA, the RA Impact of Disease (RAID) score. The entire process was very much driven by the patient perspective: the group included 12 patient research partners from 12 European countries, two of whom (MdW, MM) were also part of the steering committee. Many of the patient research partners are coauthors of this paper. The patient research partners all had personal experience of PsA and were fluent in English, but had varying experience in research partnership. Their input was key at all stages of the project (for more information, see online supplementary text).

Step 1: identification and selection of candidate domains for the PsAID score
Initial choice of domains
A literature review summarising published criteria, measures and questionnaires used in trials of PsA was presented to the 12 patient research partners. During a subsequent 2 h ‘focus group’ type meeting (where group discussions took place and notes were taken), the participants identified important domains of health in PsA in terms of impact on life, based on their personal experience. For each domain, a brief explanation was elaborated by the group.

Prioritisation of the domains
A priority exercise was performed during February to April 2011 to obtain an order of importance of the domains of health identified by the 12 research partners. The objective of this exercise was to improve external validity and to possibly reduce the number of domains. This part of the elaboration was designed as an international cross-sectional study in 140 patients with definite PsA (10 per investigator/centre), but without any other selection criteria. The names of the domains obtained in the previous step were translated by the investigators and patient partners into each language with a brief explanation and presented as a list in random order. Participants were asked to give an order of decreasing importance to the domains of health and to give a priority rating to those domains they found to be of priority (priority could be attributed to any number of domains by the patients). Additional domains of health could also be commented on. The 12 highest-rated domains were retained (see Results section for more information) for the next step after extensive discussions within the group (ie, with the health professionals and the 12 patient research partners).

Step 2: elaboration and translation of questions to measure the candidate domains
The experts selected or elaborated one question to assess each of these 12 domains of health (January to June 2011). These experts were 14 physicians (including one dermatologist) and two health professionals from 13 European countries and the 12 patient research partners.

Through a data-driven process, after an extensive literature review of published questionnaires, a numeric rating scale (NRS) was selected and modified for our purposes for each domain. When no question in the literature was found satisfactory, a question was developed de novo by the group, with much thought devoted to the wording in English. A translation/validation process was performed subsequently under the responsibility of the national principal investigator. This process followed published recommendations and included two separate translations, simple consensus, back-translation and cross-cultural validation by a multidisciplinary consensus committee and pretesting on five patients.

Step 3: weighting and validation study
Overall organisation
A cross-sectional international observational study with a longitudinal component for reliability and sensitivity to change was performed in 13 countries. Applicable general and local regulations were respected, and the project was endorsed or approved by ethics committees in each participating country. The inclusion criteria and data collected are described below. During a meeting in 2012, the results were discussed, and decisions were taken within the group (11 health professionals and nine patient research partners were present at this meeting). Thus, final decisions regarding the items of the PsAID score were driven by both data and expert opinion, with important input from the patient research partners.

Patients
Consecutive adult patients with definite PsA examined in rheumatology outpatient clinics in the participating secondary or tertiary care centres (Austria, Belgium, Estonia, Germany, France, Hungary, Ireland, Italy, Norway, Romania, Spain, Turkey and the UK) were included in 2011–2012. It was planned to include at least 400 patients (30–40 from each country) based on experience from validation of the RAID score. Selection criteria were as follows: definite PsA according to the rheumatologist, ability to fill in a questionnaire, and signed informed consent. We aimed to include patients with a range of disease severities and treatments. Patients with other concomitant inflammatory disease(s) and/or severe comorbidities (eg, recent stroke, severe cardiac failure, severe neurological disease) that could potentially influence results of assessments were excluded.

Relative importance of each domain of health
We developed a patient-derived weighting system to be able to combine the results of each question into a single score. Thus, the weights given to each question reflect its importance and relevance to the patients.

The participating patients were given the list of domains (translated as needed) and asked to assign a relative weight to each; they were asked to ‘distribute’ 100 points between the different domains. After the initial selection of 12 domains of health, it was decided also to explore the validation study scores with either nine or 12 domains of health. Thus, the patients had to give weights twice—first 100 points distributed across nine domains and then 100 points across 12 domains.

Assessment of psychometric properties
Psychometric properties were examined according to the OMERACT filter.
pain visual analogue scale (0–100 mm), and patient global assessment (0–100 mm) assessed by four questions, one for global health and one each for joints, skin and axial symptoms. Demographic data were collected. In parallel, health professionals recorded clinical features of PsA, CASPAR classification criteria, medications, recent erythrocyte sedimentation rate (ESR), and the physical and laboratory examination elements required to calculate the Disease Activity Score: DAS28-ESR.

The patients with longitudinal assessments (reliability and sensitivity of change) were asked a global question at the second data collection about whether their condition was stable, improved or worse compared with baseline (minimal clinically important difference question).

Assessment of validity (‘truth’) of the PsAID score
Face validity was ascertained by feedback from the group and from the five patients testing the questionnaire in each country.

Construct and external validity were assessed by examining cross-sectional correlations of the PsAID score with other scores.

Assessment of reliability
Patients considered to be in a stable state by the physician and with stable treatment were included in the reliability arm of the study. For this assessment, the patients filled in the questionnaire a second time from home, 2–10 days after the baseline assessment. Only patients reporting themselves to be stable at the second assessment were analysed. The objective was to include 130 patients, 10 per centre, to obtain analysable data from 100 patients (arbitrary sample size, but based on experience from validation of the RAID score).

Assessment of sensitivity to change
Patients who required an essential therapeutic change because of unacceptable clinical disease activity were included. The therapeutic change could be initiation of a synthetic or biological disease-modifying antirheumatic drug (DMARD). Patients were reassessed in the clinic 10–16 weeks after the treatment change. Only patients reporting themselves to be improved on a global change question were analysed. It was planned to include 130 patients (10 per centre) in this part of the study, with the estimate of having data to be adequately analysed from 100 patients (arbitrary sample size, but based on experience from validation of the RAID score).

Preliminary cut-off values of the PsAID score
Cut-offs were calculated to define an acceptable symptom state with the PsAID score and the minimal clinically important improvement.

Statistical analyses
Weights of domains of health in the combined PsAID scores
Mean and median points given to each domain were computed and linearly transformed to a 0–100 range. Ranks of importance of domains (based on these points) were identified in each participating patient—for example, if a domain received 20 points and was the second most important, it was ranked 2, whereas it was ranked 4 if the points were similar but it was the fourth domain. Mean and median ranks were then also computed for the group of patients and linearly transformed to a 0–100 range. In both cases, equal points gave equal weights. These two techniques were discussed with the group, and it was decided to use the ranked analysis as the basis for the final weights, as this analysis better reflects the relative importance of each domain and gives slightly less importance to the most prioritised domain (pain). The final weights for each domain are expressed as a percentage of the total score—that is, if a domain has a weight of 10%, the result for that domain will contribute 10% of the final PsAID score.

Weights were also analysed, for subgroups of patients according to the CASPAR criteria fulfilment, as sensitivity to gender, to high versus low patient global assessment, to treatment with biological agents or not, and to current psoriasis or not.

Psychometric properties
In the cross-sectional study, the several possible PsAID scores were each assessed for psychometric properties using SAS V9.2.

Feasibility
Feasibility was assessed in the cross-sectional study using the percentage of missing data for each of the questions, and distributions of scores were examined for identification of floor and ceiling effects.

Truth
Internal consistency was evaluated using Cronbach’s α coefficient. Construct validity was determined by Spearman’s correlation between the PsAID scores and other measures of disease activity/impact (including patient and physician global assessment, SF-36 summary values (physical component summary and mental component summary), HAQ, DLQI, EQ-5D and DAS28).

Reliability
This was tested with the intraclass correlation coefficient (ICC) (two-way model, single measure) with a 95% CI. An ICC of more than 0.8 is usually considered to be indicative of excellent reliability. Pearson’s correlations were also calculated. Agreement was evaluated by the Bland and Altman approach.

Sensitivity to change
The standardised response mean (SRM)—that is, the mean change from baseline to 2–4 months after the treatment change divided by the SD of the change—was calculated. An SRM >0.8 is considered large. CIs were calculated by bootstrap.

Preliminary cut-off values to interpret the PsAID scores
The cut-off value for patient-acceptable symptom state (PASS) was estimated as the 75% centile of patients considering themselves in an acceptable state at baseline. The minimal clinically relevant improvement was estimated using receiver operating characteristic curves, which were plotted using improvement versus no improvement as the outcome, and the minimal clinically relevant improvement was computed as the change score that had maximal sensitivity while maintaining a specificity of 0.80. This measure indicates the degree of change that 80% or more of patients would indicate as important.

RESULTS

Step 1: identification and selection of candidate domains for the PsAID questionnaire

Initial choice of domains
During the initial phase, the patient research partners identified 16 domains of health reflecting the impact of PsA.
Prioritisation of the domains

The 16 identified domains were ordered by 139 patients with PsA according to their importance and priority for patients (table 1). It was found that nine domains had high importance, the next four domains (embarrassment, social participation, depression, family life) had less importance, and the last three domains (concentration, discrimination, sex life) had low priority. No additional domains were identified at this stage. The entire project group (the health professionals and the 12 patient research partners) decided to move forward with validation of the NRS in the PsAID score. Thus the NRS was retained. The clinical disability.18 However, in subsequent steps, the HAQ was used with nine and 12 domains was high (see online supplementary figure S2). Thus, the nine-item and 12-item scores will provide similar information on a group level. However, the patient partners stated that the last three domains (10–12) had strong face validity because they represent important domains that should not be excluded, in particular from the consultation room to keep two versions of the PsAID questionnaire. The shorter version (PsAID-9) is geared to clinical trials, since a shorter version (PsAID-12) is geared to clinical practice, as the responses to each question can

<table>
<thead>
<tr>
<th>Domain number (by order of importance)</th>
<th>Domain and short defining statement</th>
<th>Median order of importance (range of importance 1–16)</th>
<th>% patients ordering this domain in the top 8</th>
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<tbody>
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<td>2.56</td>
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<td>0</td>
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<td>6.43</td>
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Patients gave each domain both an order of importance (from 1 to 16) and a priority rating (yes/no). The 12 domains with highest median rank were retained, and domain 13 was merged with domain 11 for the final PsAID scores.

PsA, psoriatic arthritis; PsAID, PsA Impact of Disease.

Step 2: elaboration and translation of questions to measure the candidate domains

It was decided to use single questions (assessed by NRS) for each domain, and to assess both an NRS and the HAQ for physical disability.18 However, in subsequent steps, the HAQ was not found to perform better for psychometric properties than the NRS in the PsAID score. Thus the NRS was retained. The wording and time frame of the single questions were also discussed. Specific wordings were obtained for each question, and a time frame of 1 week was decided on (see online supplementary table S1). The 12 questions were translated into 11 languages (Estonian, Flemish, French, German, Hungarian, Italian, Norwegian, Romanian, Russian, Spanish, Turkish).

Step 3: weighting and validation study

In total, 499 patients participated in this part of the study; 474 had analysable data (table 2). Mean±SD age was 50.4±12.6 years, mean disease duration was 9.6±9.4 years; 50.2% were female. The population had, on average, moderate disease activity, and half were treated with biological DMARDs; 75.5% satisfied the CASP AR criteria.29

Final decisions taken for the PsAID questionnaire

Comparison of 12 and nine domains in terms of psychometric properties showed no improvement with the three additional domains, and the correlation and agreement between results with nine and 12 domains was high (see online supplementary figure S2). Thus, the nine-item and 12-item scores will provide similar information on a group level. However, the patient partners stated that the last three domains (10–12) had strong face validity because they represent important domains that should not be excluded, in particular from the consultation room where these three domains provide relevant additional information to the individual health professional. Thus, it was decided to keep two versions of the PsAID questionnaire. The shorter version (PsAID-9) is geared to clinical trials, since a shorter questionnaire is more feasible. The longer version (PsAID-12) is geared to clinical practice, as the responses to each question can

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**Table 1** Sixteen domains of health identified as important by 12 patients with PsA and their order of importance and priority ratings based on information from 139 patients with PsA

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provide important information to the healthcare provider, which will help in making shared decisions with the patient on a management plan.

Weights of domains of health

The relative importance of the individual domains was decided for both PsAID questionnaires (PsAID-9 and PsAID-12) (table 3): pain, skin problems and fatigue had the highest relative importance in both weighting exercises. Of note, these weights were very stable in the subgroups of patients (sensitivity analyses, data not shown), although, as expected, weights of questions related to skin were higher in patients with current psoriasis. For PsAID-12 (developed for clinical practice), a simplified weighting system was chosen so that calculation of the final aggregate score does not entail the use of a calculator (tables 3 and 4 and see online supplementary table S1).

Psychometric properties

Feasibility

The percentage of missing data was very low (1%), and floor and ceiling effects were also very low (respectively, 1% and 0%).

‘Truth’

The scores using nine and 12 domains had very similar internal consistency (Cronbach’s $\alpha$ 0.93–0.94) and similar correlations with other scores, and correlations were, as expected, higher with other PROMs, particularly with patient global assessment, than with physician-derived scores (table 5).

Test–retest reliability

A total of 107 patients had a second assessment for reliability, but only 88 were analysed who all estimated themselves to be in a stable state. Reliability was high and similar for the two possible PsAID scores. The ICCs for the PsAID-12 and PsAID-9 questionnaires were 0.95 (95% CI 0.92 to 0.96) and 0.94 (95% CI 0.91 to 0.96), respectively. These results were consistent with results for other widely used measures (eg, ICC of HAQ was 0.97 (0.95 to 0.98) in the same population). Corresponding results for Pearson correlation were 0.91 (95% CI 0.86 to 0.93) for PsAID-9, 0.90 (0.86 to 0.93) for PsAID-12, and 0.96 (0.95 to 0.97) for HAQ.

Sensitivity to change

This was assessed in 105 patients, but only 71 estimated themselves to be improved and were analysed. Half were started on a biological DMARD and half on a conventional synthetic

### Table 2 Description of the 474 patients with PsAID participating in the weighting and validation study

<table>
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<tr>
<th>Characteristic</th>
<th>Mean (SD) (range) or N (%)</th>
<th>Range across countries of mean values or proportions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex</td>
<td>235 (50.2%)</td>
<td>32.5–81.8%</td>
</tr>
<tr>
<td>Age, years</td>
<td>50.4 (12.6) (20.8–80.1)</td>
<td>42.6–55.1%</td>
</tr>
<tr>
<td>Disease duration, years</td>
<td>9.6 (9.4) (0.0–41.9)</td>
<td>5.4–16.2%</td>
</tr>
<tr>
<td>Formal education, years</td>
<td>12.5 (3.8) (6–20)</td>
<td>9.9–14.9%</td>
</tr>
<tr>
<td>Proportion of patients fulfilling CASPAR criteria</td>
<td>351 (75.5%)</td>
<td>23.8–94.6%</td>
</tr>
<tr>
<td>Current disease-modifying drug</td>
<td>306 (66.8%)</td>
<td>36.5–85.0%</td>
</tr>
<tr>
<td>Current biological treatment</td>
<td>202 (46.8%)</td>
<td>23.5–79.1%</td>
</tr>
<tr>
<td>Swollen joint count (0–66)</td>
<td>2.4 (4.1) (0–36)</td>
<td>0.6–5.1</td>
</tr>
<tr>
<td>Tender joint count (0–68)</td>
<td>5.4 (8.0) (0–52)</td>
<td>1.8–14.0</td>
</tr>
<tr>
<td>Current skin psoriasis</td>
<td>286 (65.2%)</td>
<td>10.5–97.3%</td>
</tr>
<tr>
<td>DAS28-ESR</td>
<td>2.8 (1.4) (0.0–7.1)</td>
<td>2.2–4.1</td>
</tr>
<tr>
<td>HAQ (0–3)</td>
<td>0.81 (0.70) (0–2.75)</td>
<td>0.63–1.01</td>
</tr>
<tr>
<td>Patient global assessment (0–10)</td>
<td>4.1 (2.8) (0–10)</td>
<td>2.9–5.1</td>
</tr>
<tr>
<td>Pain (0–10)</td>
<td>4.7 (2.9) (0–10)</td>
<td>3.6–6.2</td>
</tr>
<tr>
<td>DLOI (0–30)</td>
<td>4.3 (5.7) (0–30)</td>
<td>1.7–8.8</td>
</tr>
<tr>
<td>EQ-5D</td>
<td>0.60 (0.30) (−0.59 to 1)</td>
<td>0.47–0.72</td>
</tr>
<tr>
<td>SF-36 physical component</td>
<td>37.7 (10.5) (13.5–58.3)</td>
<td>32.1–40.3</td>
</tr>
<tr>
<td>SF-36 mental component</td>
<td>47.0 (11.5) (15.1–68.9)</td>
<td>42.9–54.8</td>
</tr>
</tbody>
</table>

Fever than 10% of the data were missing for all elements except swollen joint count (10.2% missing data), DAS28 (27.0%) and SF-36 (21.9%).

*Percentages are % of available data.

†Range of means or percentages: minimum and maximum values for means or percentages observed in participating countries.

DAS28, Disease Activity Score (28 joints); DLOI, Dermatology Life Quality Index; EQ-5D, EuroQol-5D; ESR, erythrocyte sedimentation rate; HAQ, Health Assessment Questionnaire; SF-36, short form (36) generic quality of life scale.

### Table 3 Domains of health in the PsAID questionnaires with their relative importance as a percentage: results from the weighting exercise

<table>
<thead>
<tr>
<th>Domain of health</th>
<th>Relative importance in the PsAID-12 score for clinical practice</th>
<th>Relative importance in the PsAID-9 score for clinical trials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain</td>
<td>15</td>
<td>17.4</td>
</tr>
<tr>
<td>Fatigue</td>
<td>10</td>
<td>13.1</td>
</tr>
<tr>
<td>Skin problems</td>
<td>10</td>
<td>12.1</td>
</tr>
<tr>
<td>Work and/or leisure activities</td>
<td>10</td>
<td>11.0</td>
</tr>
<tr>
<td>Functional capacity</td>
<td>10</td>
<td>10.7</td>
</tr>
<tr>
<td>Discomfort</td>
<td>10</td>
<td>9.8</td>
</tr>
<tr>
<td>Sleep disturbance</td>
<td>10</td>
<td>8.9</td>
</tr>
<tr>
<td>Coping</td>
<td>5</td>
<td>8.7</td>
</tr>
<tr>
<td>Anxiety</td>
<td>5</td>
<td>8.5</td>
</tr>
<tr>
<td>Embarrassment and/or shame</td>
<td>5</td>
<td>NA</td>
</tr>
<tr>
<td>Social participation</td>
<td>5</td>
<td>NA</td>
</tr>
<tr>
<td>Depression</td>
<td>5</td>
<td>NA</td>
</tr>
</tbody>
</table>

NA, not applicable; PsAID, Psoriatic Arthritis Impact of Disease.
The PASS cut-off was assessed in 274 patients and was found to be a PsAID-9 value of ≤4.10 and a PsAID-12 value of ≤3.95. Thus, the proposed PASS cut-off is 4 for both scores.

Preliminary cut-off values for interpreting the PsAID scores

The PASS cut-off was assessed in 274 patients and was found to be a PsAID-9 value of ≤4.10 and a PsAID-12 value of ≤3.95. Thus, the proposed PASS cut-off is 4 for both scores.

A preliminary value for the minimal clinically important improvement was found to be 3 points (calculations were 3.6 points for change in PsAID-9 and 3.0 points for change in PsAID-12). The proposed PASS cut-off is 4 for both scores.

The PsAID questionnaires are available online free of charge with their translations in 12 languages (English, Estonian, Russian, Spanish, Turkish) at http://www.eular.org/index.cfm?framePage=/st_com_clinical_tools.cfm.

**DISCUSSION**

Two patient-derived questionnaires for assessing the impact of PsA from the patients’ perspective are proposed. The longer questionnaire, developed for clinical practice, includes 12 domains of health, each assessed by a single question with response on an NRS. The calculation to obtain a single score result is simplified in order to be feasible for use in the clinic. The shorter questionnaire, developed for clinical trials, includes nine domains of health and appears to bring similar levels of information at the group level. The PsAID scores had satisfactory psychometric properties in an international validation study. These questionnaires, elaborated with the participation of patient research partners, should allow a more thorough quantified assessment of the patient perspective in PsA.

The final selection of domains seems to have good face validity, as pain, skin problems and fatigue appear to be the most important to many patients; however, qualitative studies to confirm the priority of these domains are lacking. Of these three domains, only pain is part of the PsA core set and is regularly reported in PsA studies. Skin problems as experienced by patients and fatigue are originally identified aspects of the disease found in the present study and these domains should be further explored. The other domains of health in the PsAID scores reflect physical, psychological and societal aspects, which is in keeping with the International Classification of Functioning, Disability and Health. Feelings of shame due to physical appearance seem to be more important in patients with PsA than in RA, probably because of the skin component of the disease. Of note, five of the 12 domains of the PsAID questionnaire (ie, 42%) were not previously identified as important in PsA. Patient global assessment was not selected by the people with PsA in the present study, which may indicate that ‘patient global’ is not a notion that ‘makes sense’ to patients. The international development of the PsAID scores means that the domains of health assessed in the scores are probably relevant across countries and cultures, but this assumption should also be studied further.

The weights attributed to each domain were based on the patients’ scoring of the importance of the domains. We recognise that other approaches could have been applied to weight the domains, but this approach allowed us to score each domain according to its importance for the group of 474 patients included in the study. Regarding external validity of the weights,
Clinical and epidemiological research

it should be noted that, in the validation study, a high percentage of patients were taking biological agents, which reflects the centres involved (secondary or tertiary care rheumatology centres) and the inclusion process—many patients were followed-up in day hospital units. These issues may be a limitation to keep in mind for generalisability. However, the relative importance of the different domains was similar in patients treated with biological agents or not, and in patients with a high or low patient global assessment (results not shown). These observations support the relevance and generalisability of the preliminary PsAID questionnaires.

This study has both strengths and weaknesses. Strengths include the central involvement of patients in the elaboration of the PsAID score and the inclusion of patient research partners with PsA from 12 countries with different cultures and socioeconomic backgrounds. Furthermore, the PsAID questionnaires were validated with more than 470 patients from 13 countries. The scores have good face validity and also potentially good generalisability. Finally, the methodology used to obtain patient-derived weights for combination of the results into a single score is novel for PsA and could be applied in other contexts. It has, for example, also been used in the elaboration of a Pancreatic Cancer Impact of Disease (PACADI) score.45 One weakness is that patients had a clinical diagnosis of PsA, and it turned out that 24.5% of the patients in the validation study did not formally fulfil the CASPAR classification criteria.46 However, weighting results were similar in the subpopulation that did fulfil the CASPAR criteria.

The impact of PsA is usually assessed by levels of pain and patient global assessment.10–17 Concerns have been raised that these instruments may not adequately capture all patient-relevant data, which was the basis for the development of this new questionnaire. Complex generic or arthritis-specific quality of life instruments such as SF-3631 and the Arthritis Impact Measurement Scale (AIMS2)46 capture information on many domains of health, including mental health and social functioning. However, these questionnaires are long, and interpretation of the scores is complex, especially since they do not provide a single score reflecting all domains. However, patient-reported data are strongly collinear, and the new PsAID scores correlate strongly with patient global assessment. Therefore the additional information over the existing indices obtained by adding more variables will need to be further explored at a group level. At the patient level, however, the PsAID-12 questionnaire (for clinical practice) should allow a more precise assessment of the impact of PsA, helping healthcare providers and patients to make shared treatment decisions geared to either disease activity or, for example, psychological distress.

In conclusion, this study enabled us to propose two preliminary patient-derived weighted questionnaires for assessing the impact of PsA. PsAID-9 is viewed as an additional instrument for the assessment of PsA in clinical trials, giving supplementary information on patient-relevant domains of health. The PsAID-12 score will hopefully be valuable in clinical practice, both for identification of areas that should be addressed in clinical management and by monitoring the patients longitudinally. However, further validation of the PsAID score is needed, in particular regarding sensitivity to change in comparison with other outcome measures in PsA.47

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Collaborators On behalf of the EULAR PsAID Taskforce.

Contributors All of the authors fulfill the following criteria: substantial contributions to the conception or design of the work; or the acquisition, analysis or interpretation of data for the work; and/or drafting the work or revising it critically for important intellectual content; and final approval of the version to be published; and agreement to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The guarantors who accept full responsibility for the work and/or the conduct of the study, had access to the data, and controlled the decision to publish are LG, MdW and T.KK.

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Competing interests LG has received fees for speaking and/or consulting from AbbVie, Janssen, Novartis, Pfizer/Wyeth, Roche and UCB. MdW has received fees for consulting from AbbVie and Roche. UK has received grant and research support and consultancy fees from AbbVie, MSD, Pfizer and UCB and also fees for speaking and/or consulting from AbbVie, Pfizer/Wyeth, Roche and MSD/Shering-Plough. JB has received honoraria for talks, advisory boards, paid consultancies and grants for studies from AbbVie, Amgen, BMS, Boehringer, Celgene, Celltrion, Centocor, Chugai, EBEWY Pharma, Medac, MSD (Shering-Plough), Mundipharma, Novartis, Pfizer (Wyeth), Roche, Sanofi-Aventis, UCB. RS has received fees for consulting from AbbVie. KO has received fees for research funding and consulting from MSD. JDC has received fees for speaking and/or consulting from AbbVie, Celgene, Janssen, MSD/Shering Plough, Novartis, Pfizer and UCB. AB has received fees for speaking and/or consulting from AbbVie, BMS, Janssen, MSD, Pfizer/Wyeth, Roche and UCB. KB has received fees for speaking and/or consulting from AbbVie, BMS, Janssen, MSD, Pfizer/Wyeth, Roche and UCB. PH has received fees for speaking and/or consulting from AbbVie, BMS, Janssen, Celgene, MSD, Novartis, Pfizer/Wyeth, Roche and UCB. TL has received fees for speaking and/or consulting from AbbVie, Janssen-Cilag, MSD, Novartis, Pfizer and Sandoz. T.KK has received fees for speaking and/or consulting from AbbVie, Hospira, Celltrion, Orion Pharma, MSD/Shering-Plough, Pfizer/Wyeth and UCB and received...
REFERENCES
Procedure of patient involvement in the elaboration and validation of the PsAID

Making the involvement of patients with PsA an integral part of the development and validation of the PsAID has been essential for the project. To incorporate the patient perspective people with PsA have been involved in different roles, different phases and in different numbers. They carried out the following tasks:
- Collaborating as Patient Research Partner (n=12; during the entire project)
- Acting as member of the Steering Group (n=2; during the entire project)
- Prioritizing important domains (n=139; step 1)
- Pre-testing of translated items through cognitive debriefing (n=65, 5 per country; step 2)
- Filling in questionnaires (n=499; step 3)

The 12 patient research partners were actively involved in many phases of the study. Here we will describe their role as well as that of the members of the Steering Group.

Patient Research Partners

Recruitment and selection
Proportional representation of patients as equal collaborators in the project was sought during the first as well as the second meeting of the entire research team. This means that there was an almost equal number of patient research partners (n=12) compared to the number of national principal investigators (n=13). The patients came from 12 different European countries and the recruitment was carried out through the clinics of the participating investigators. They were able to make an adequate assessment of whether a patient was competent to contribute to this research project and whether a patient fulfilled the inclusion criteria as recommended by EULAR among which personal experience with the disease under research and being fluent in English.

Identifying domains
During the breakout session on the morning of the first research team meeting (January 2011) the patient research partners met as a subgroup and identified important domains that reflected the impact of the disease on their health. They came up with a total number of 16 domains. In the afternoon a plenary session with the investigators took place discussing the 16 domains and trying to formulate brief descriptions of the domains.

Translation process
Some research partners were involved in the process of translating the English version of the PsAID to their national language, as part of a multi-disciplinary team.

Decision making process
Along the way the research partners were involved in many decisions that were taken. Sometimes this process was carried out through emails or telecalls. Some decisions were taken during short meetings at existing annual conferences of the

Online supplementary text
American College of Rheumatology (ACR) and the European League Against Rheumatology (EULAR). These meetings were attended by some of the national principal investigators that were attending the conference and only a few research partners. The most important decisions were taken during the two research team meetings. The research partners had full voting rights and received in advance of the meetings all documents written and explained in lay language (see below among ‘support’).

Co-authorship
Research partners who contributed during different phases of the study and who reviewed and commented on the draft article were offered co-authorship of the manuscript.

Members of the Steering Group

The Steering Group consisted of five investigators and two expert patients with extensive experience in international collaboration with researchers. They followed the EULAR recommendations for the inclusion of patient representatives in scientific projects. The inclusion of the patient perspective was felt a shared responsibility. Here we describe the role of the Steering Group in supporting the patient research partners and the additional task of the patient representative.

Supporting the patient research partners
Enabling patient research partners to contribute to the research process requires additional support by the researchers before, during and after meetings. The Steering Group undertook the following actions to make sure that the patient research partners were prepared for their role and were well informed about the research process:

• A personalized invitation letter and background information, including a lay version of the protocol.
• Patient sessions before the start of the first and the second research team meetings (90 min.); participation by national investigators was optional.
• Meeting reports, newsletters and regular project updates in lay language
• Pre-meeting patient guide before the second meeting
• Additional individual support was provided by the participating national investigators.

Moderation of sessions
An important challenge for the Steering Group was to preserve the characteristics of a genuine dialogue between research partners and physicians in which arguments were shared in an open and safe atmosphere and where all participants felt equally facilitated to contribute to the discussions without limitations caused by traditional doctor-patient hierarchy. Especially for the first breakout session during the first research team meeting it was thought essential that the patients would not feel restricted to speak up due to the presence of their treating physician. Therefore it was decided to organize two homogeneous subgroups led by different moderators. One patient member of the Steering Group with professional moderation skills, together with an experienced nurse researcher, facilitated the discussion in the patient group to identify the domains of interest for patients. Two
investigators of the Steering Group facilitated the discussion in the physician group to discuss the instruments to measure the potential domains.

Two weeks in advance of the second research team meeting (November 2012) patient research partners received a 12 page introduction package explaining the previous steps of the project, the validation process of the PsAID, the objectives of the second meeting and a glossary of often used terms and abbreviations. During the second meeting the facilitators were keen to make sure that all participants could follow the discussions and felt confident to contribute to the discussions at all stages. At the end of the day the meeting was evaluated and the patient research partners confirmed that they believed that they had contributed something that the physicians could not provide. They felt well supported in the process and satisfied about their role.

**Added value of involving patient research partners**

The input of patient research partners during the first meeting of the research team resulted in a concrete list of 16 relevant domains from the patient perspective. Thereafter the research partners were involved in lively discussions with the researchers on the formulation of the 16 domains (wording, phrasing). Similarities, overlap or differences between different terms used in the questions were explained. This discussion included also the length of the recall period (1 week or 1 month) and the choice of the instrument: NRS or other tools.

There was a long debate during the first meeting regarding the domain ‘coping’. The physicians argued that “coping” does not represent the impact of the disease and should be seen as another dimension. The patients experienced the extent to which one is able to cope with the disease as a clear outcome of the treatment and therefore also as an appropriate symptom or feature of the impact of the disease. Because the patients felt strongly about this domain the research team decided to keep ‘coping’ in the list of relevant domains.

During the second research team meeting the input of the research partners was less tangible because of the nature of the meeting: presenting the final data and statistical analyses from the validation process. However, patients were involved in several discussions on outstanding issues that needed to be decided on. These discussions included the decision about the number of items (9 or 12). The result was a compromise that achieved a high level of agreement, suggesting 9 items as the recommended version for clinical trials and the 12 items as the recommended version for clinical practice. Patients wanted to keep the three domains ‘embarrassment and shame’, ‘social participation’ and ‘depression’ because there was a strong believe these domains are important for people with PsA although patients often don’t want to admit or acknowledge that their disease contribute to the experience of feeling down, depressed, socially isolated or not valued as a person. The assumption was that there exists a hidden impact of PsA that it often not recognized by patients as well as physicians. And because the PsAID will hopefully also be used in clinical practice, keeping these 3 items in, would facilitate treating physicians to focus on separate items of the composite score, including the 3 items that are not contributing to the pooled result. Most of the physician present at the meeting acknowledged the opinions of patients and accepted the view of the patients as being decisive for developing a patient derived PRO. For this reason they voted in favor of the 12-item version for clinical practice. Patient representatives did acknowledge the arguments of the physicians that for
clinical trials short questionnaires are needed. Because the data showed no significant difference in performance of the 9 and 12 item versions, the research partners accepted the 9 item version as recommended for clinical trials. The research partners also agreed with the decision for different weighting systems for the 9 and 12 item versions.

Online supplementary table S1. The final PsAID questionnaires in English. Separate document
Online supplementary Figure S1 – The process used to develop the PsAID

Identification / selection of domains of health

Identification / elaboration of questions

Physician expert opinion
Patient-partner opinion on wording

Domains prioritised for importance
139 patients from 13 countries
Priority (yes/no)

PsAID finalised

First physical meeting of Task force

Validation

Cross-sectional and longitudinal study in 13 countries, define PsA patients according to the physician

To obtain weights of the different questions to combine them in a score
Patients asked to ‘spend’ 100 points across domains
Analysis by ranks of points

Validation of ‘Truth’
Construct validity: correlation with other questionnaires: OTQ8, PCQ and MOS, HAQ, Pain VAS, Patient global VAS, LQ-5D, ULQI
Other measures: UMPIA, UMPAS

Feasibility: % missing data

Test-retest reliability
Stable patients
Second questionnaire filled in one week later at home
Intraclass correlation (ICC) and Pearson correlation

Sensitivity to change
Change 10-16 weeks after treatment initiation (conventional DMARD or biologic)
Standardized response mean (SRM)

Validation process
Cross-sectional N=474 patients
Test-retest N=88
Sensitivity to change N=71

12 domains of health
12 questions

9 domains of health
9 questions

2 validated PsAID questionnaires
PsAID-9 for clinical trials
PsAID-12 for clinical practice
**Online Figure S2.** Agreement between PsAID-9 and PsAID-12 score results in 474 PsA patients by Bland and Altman technique.

Agreement was high, with a mean difference close to 0 and agreement limits of -0.7 to +0.3.
The EULAR Psoriatic Arthritis Impact of Disease: PsAID9 for clinical trials

We want you to indicate how much your psoriatic arthritis impacts your health. Please tell us how you have been feeling this last week.

1. Pain
Circle the number that best describes the pain you felt due to your psoriatic arthritis during the last week:

None 0 1 2 3 4 5 6 7 8 9 10 Extreme

2. Fatigue
Circle the number that best describes the overall level of fatigue due to your psoriatic arthritis you have experienced during the last week:

No fatigue 0 1 2 3 4 5 6 7 8 9 10 Totally exhausted

3. Skin problems
Circle the number that best describes the skin problems including itching you felt due to your psoriatic arthritis during the last week:

None 0 1 2 3 4 5 6 7 8 9 10 Extreme

4. Work and/or leisure activities
Circle the number that best describes the difficulties you had to participate fully in work and/or leisure activities due to your psoriatic arthritis during the last week:

None 0 1 2 3 4 5 6 7 8 9 10 Extreme

5. Functional capacity
Circle the number that best describes the difficulty you had in doing daily physical activities due to your psoriatic arthritis during the last week:

No difficulty 0 1 2 3 4 5 6 7 8 9 10 Extreme difficulty

6. Discomfort
Circle the number that best describes the feeling of discomfort and annoyance with everyday tasks due to your psoriatic arthritis during the last week:

None 0 1 2 3 4 5 6 7 8 9 10 Extreme

7. Sleep disturbance
Circle the number that best describes the sleep difficulties (i.e., resting at night) you felt due to your psoriatic arthritis during the last week:

No difficulty 0 1 2 3 4 5 6 7 8 9 10 Extreme difficulty
8. Coping
Considering your psoriatic arthritis overall, how well did you cope (manage, deal, make do) with your psoriatic arthritis during the last week?

<table>
<thead>
<tr>
<th>Very well</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>Very poorly</th>
</tr>
</thead>
</table>

9. Anxiety, fear and uncertainty
Circle the number that best describes the level of anxiety, fear and uncertainty (for example about the future, treatments, fear of loneliness) due to your psoriatic arthritis you have experienced during the last week:

<table>
<thead>
<tr>
<th>None</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>Extreme</th>
</tr>
</thead>
</table>

THANK YOU FOR ANSWERING THIS QUESTIONNAIRE
PsAID9 SCORING AND CALCULATION RULES

The PsAID is calculated based on 9 Numerical rating scales (NRS) questions. Each NRS is assessed as a number between 0 and 10.

1. Calculation

PsAID final value =
   (PsAID1 (pain) NRS value (range 0-10) x 0.174)
   + (PsAID2 (fatigue) NRS value (range 0-10) x 0.131)
   + (PsAID3 (skin) NRS value (range 0-10) x 0.121)
   + (PsAID4 (Work and/or leisure activities) NRS value (range 0-10) x 0.110)
   + (PsAID5 (function) NRS value (range 0-10) x 0.107)
   + (PsAID6 (discomfort) NRS value (range 0-10) x 0.098)
   + (PsAID7 (sleep) NRS value (range 0-10) x 0.089)
   + (PsAID8 (coping) NRS value (range 0-10) x 0.087)
   + (PsAID9 (anxiety) NRS value (range 0-10) x 0.085)

Thus, the range of the final PsAID value is 0-10 where higher figures indicate worse status.

2. Missing data imputation

If one of the 9 NRS values composing the PsAID is missing, the imputation is as follows:
   a. calculate the mean value of the 8 other (non-missing) NRS (range, 0-10)
   b. impute this value for the missing NRS
   c. Then, calculate the PsAID as explained above.

If 2 or more of the NRS are missing, the PsAID is considered as missing value (no imputation).
The EULAR Psoriatic Arthritis Impact of Disease: PsAID12 for clinical practice

We want you to indicate how much your psoriatic arthritis impacts your health. Please tell us how you have been feeling this last week.

1. Pain
Circle the number that best describes the pain you felt due to your psoriatic arthritis during the last week:

| None | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme |

2. Fatigue
Circle the number that best describes the overall level of fatigue due to your psoriatic arthritis you have experienced during the last week:

| No fatigue | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Totally exhausted |

3. Skin problems
Circle the number that best describes the skin problems including itching you felt due to your psoriatic arthritis during the last week:

| None | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme |

4. Work and/or leisure activities
Circle the number that best describes the difficulties you had to participate fully in work and/or leisure activities due to your psoriatic arthritis during the last week:

| None | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme |

5. Functional capacity
Circle the number that best describes the difficulty you had in doing daily physical activities due to your psoriatic arthritis during the last week:

| No difficulty | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme difficulty |

6. Discomfort
Circle the number that best describes the feeling of discomfort and annoyance with everyday tasks due to your psoriatic arthritis during the last week:

| None | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme |

7. Sleep disturbance
Circle the number that best describes the sleep difficulties (i.e., resting at night) you felt due to your psoriatic arthritis during the last week:

| No difficulty | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | Extreme difficulty |
8. Coping
Considering your psoriatic arthritis overall, how well did you cope (manage, deal, make do) with your psoriatic arthritis during the last week?

<table>
<thead>
<tr>
<th>Very well 0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Very poorly</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

9. Anxiety, fear and uncertainty
Circle the number that best describes the level of anxiety, fear and uncertainty (for example about the future, treatments, fear of loneliness) due to your psoriatic arthritis you have experienced during the last week:

<table>
<thead>
<tr>
<th>None</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Extreme</td>
<td></td>
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<td></td>
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<td></td>
</tr>
</tbody>
</table>

10. Embarrassment and/or shame
Considering your psoriatic arthritis overall, circle the number that best describes the level of embarrassment and/or shame due to your appearance experienced during the last week:

<table>
<thead>
<tr>
<th>None</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Extreme</td>
<td></td>
<td></td>
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<td></td>
<td></td>
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</tr>
</tbody>
</table>

11. Social participation
Circle the number that best describes the difficulties you had to participate fully in social activities (including relationships with family and/or people very close to you) due to your psoriatic arthritis during the last week:

<table>
<thead>
<tr>
<th>None</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Extreme</td>
<td></td>
<td></td>
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<td></td>
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</tr>
</tbody>
</table>

12. Depression
Circle the number that best describes the level of depression due to your psoriatic arthritis you have experienced during the last week:

<table>
<thead>
<tr>
<th>None</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Extreme</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

THANK YOU FOR ANSWERING THIS QUESTIONNAIRE

Final PsAID out of 10
Add up the □ and divide by 20:
PsAID12 SCORING AND CALCULATION RULES

The PsAID is calculated based on 12 Numerical rating scales (NRS) questions. Each NRS is assessed as a number between 0 and 10.

3. Calculation

PsAID final value =

(PsAID1 (pain) NRS value (range 0-10) x 3) + (PsAID2 (fatigue) NRS value (range 0-10) x 2) + (PsAID3 (skin) NRS value (range 0-10) x 2) + (PsAID4 (Work and/or leisure activities) NRS value (range 0-10) x 2) + (PsAID5 (function) NRS value (range 0-10) x 2) + (PsAID6 (discomfort) NRS value (range 0-10) x 2) + (PsAID7 (sleep) NRS value (range 0-10) x 2) + (PsAID8 (coping) NRS value (range 0-10) x 1) + (PsAID9 (anxiety) NRS value (range 0-10) x 1) + (PsAID10 (embarrassment) NRS value (range 0-10) x 1) + (PsAID11 (social life) NRS value (range 0-10) x 1) + (PsAID12 (depression) NRS value (range 0-10) x 1)

The total is divided by 20.

Thus, the range of the final PsAID value is 0-10 where higher figures indicate worse status.

4. Missing data imputation

If one of the 12 NRS values composing the PsAID is missing, the imputation is as follows:

a. calculate the mean value of the 11 other (non-missing) NRS (range, 0-10)
b. impute this value for the missing NRS
c. Then, calculate the PsAID as explained above.

If 2 or more of the NRS are missing, the PsAID is considered as missing value (no imputation).