Comparison of disease activity measures in early psoriatic arthritis in usual care

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Abstract

Objectives. To compare responsiveness and longitudinal validity of Disease Activity Score 28 (DAS28), Disease Activity index for Psoriatic Arthritis (DAPSA), Composite Psoriatic Disease Activity Index (CPDAI), Psoriatic Arthritis Disease Activity Score (PASDAS), GRAppa Composite scorE (GRACE) and Minimal Disease Activity (MDA) in usual care PsA patients, within 1 year after diagnosis.

Methods. Data collected in the Dutch southwest early PsA cohort (DEPAR) were used. Responsiveness was assessed using effect size (ES), standardized response mean (SRM), and discrimination between different general health states. Longitudinal validity was tested using mixed models with outcomes health-related quality of life (HRQOL), productivity and disability.

Results. Responsiveness was highest for PASDAS, with ES 1.00 and SRM 0.95, lowest for DAPSA, with ES 0.73 and SRM 0.71, and in between for DAS28, CPDAI and GRACE. Differences in general health were best discriminated with PASDAS and GRACE. Patients reporting stable or worsening general health could not be distinguished by DAS28 or CPDAI. Discrimination was better using DAPSA, but worse than when using PASDAS and GRACE. Longitudinal evolution of HRQOL and productivity had the highest association with low disease activity according to GRACE, followed by PASDAS, MDA, DAPSA, DAS28, with the lowest association for CPDAI.

Conclusion. PASDAS and GRACE were superior with respect to responsiveness, and together with MDA best related to longitudinal evolution of HRQOL, productivity and disability. Responsiveness and longitudinal validity of most outcomes were inferior for DAS28, DAPSA and CPDAI. As alternatives to the continuous measure DAPSA, use of PASDAS or GRACE should be considered.

Key words: psoriatic arthritis, disease activity, responsiveness, longitudinal validity

Rheumatology key messages

- Responsiveness and longitudinal validity of Psoriatic Arthritis Disease Activity Score, GRAppa Composite scorE and Minimal Disease Activity were superior
- Psoriatic Arthritis Disease Activity Score and GRAppa Composite scorE reflect the spectrum of disease activity better than Disease Activity index for PSoriatic Arthritis.

Introduction

PsA is a heterogeneous disease, with manifestations arthritis, enthesitis, spondylitis, dactylitis, and psoriasis [1, 2]. The goal of treatment of PsA is to optimize function and health-related quality of life (HRQOL), and to prevent structural damage. This can be done by aiming at remission or, if this cannot be achieved, low or minimal disease activity [3, 4]. Disease activity is assessed using composite measures, in which multiple aspects of disease are combined in a total score of level of disease activity. In rheumatoid arthritis, guiding treatment based on Remission and ACR50/66/70 response criteria. As alternatives to the continuous measure DAPSA, use of PASDAS or GRACE should be considered.
measuring disease activity with the Disease Activity Score 28 (DAS28) has improved care and long term outcomes [5]. Though multiple disease activity measures are available and used in research in PsA [6], no consensus has been reached on which measure should be used [7].

The DAS28 has often been used as a disease activity measure [6], although it was not originally developed for use in patients with PsA. As a more PsA-specific measure, the Disease Activity index for Psoriatic Arthritis (DAPSA) was developed using the 66/68 joint count instead of the 28 joint count [8]. Both DAPSA and DAS28 are mainly articular measures. Some have argued that the target for PsA should take into account more than joint involvement alone, for example by using the Composite Psoriatic Disease Activity Index (CPDAI) [9], Psoriatic Arthritis Disease Activity Score (PASDAS) [10], or Minimal Disease Activity (MDA) [12]. The latter is a dichotomous measure, while the others are continuous measures similar to the DAPSA and DAS28. For clinical practice, use of either DAPSA or MDA has been advised by an international task force [7]. An overview of the components needed to calculate each measure is given in Table 1.

All measures have been shown to be related to disease burden. The continuous measures were all able to discriminate between placebo and active treatment groups in trials [13, 14], and were related to treatment change [10, 15] and a patient-acceptable symptom state [16]. Though all measures are able to distinguish two groups with different levels of disease activity, other cross-sectional studies have shown that agreement on a patient level is often moderate [16, 17]. Also, the ReFlap study has shown that agreement between patient opinion and low disease activity according to MDA or DAPSA is limited [18]. For use in clinical practice, we need to know which measure is best at measuring change of disease activity (i.e. responsiveness) and has the strongest relation with patient outcomes (i.e. longitudinal validity), which has not been studied in a usual care population of patients with early disease. Responsiveness has been tested in an analysis of trial data [14], but not in a longitudinal study of usual care patients. Also, little is known on how these measures perform in the early course of disease. We therefore aimed to compare the responsiveness and longitudinal validity of the currently available composite disease activity measure (DAS28, DAPSA, CPDAI, PASDAS, GRACE and MDA) in PsA patients within 1 year after diagnosis.

Responsiveness was evaluated using both a distribution-based and an anchor-based approach, and longitudinal validity using associations with HRQOL, productivity and disability as no reference standard for disease activity exists.

**Patients and methods**

**Patients and setting**

We used data collected in the Dutch southwest Early Psoriatic Arthritis cohoRt (DEPAR) study, of which details

| Component                  | DAS28 | DAPSA | CPDAI | PASDAS | GRACE | MDA |
|---------------------------|-------|-------|-------|--------|-------|-----|
| **Clinical assessment**    |       |       |       |        |       |     |
| Tender joint count         | 28    | 68    | 68    | 68     | 68    | 68  |
| Swollen joint count        | 28    | 66    | 66    | 66     | 66    | 66  |
| PASI                       |       |       | x     | x      | x     | x   |
| LEI                        |       | x     | x     | x      |       |     |
| Dactylitis count           |       | x     |       | x      |       |     |
| VAS physician              |       |       |       | x      |       |     |
| **Patient questionnaire**  |       |       | x     | x      | x     |     |
| VAS global                 |       |       |       |       | x     |     |
| VAS skin                   |       |       |       |       |       | x   |
| VAS joints                 |       |       |       |       |       | x   |
| VAS pain                   |       |       |       |       |       | x   |
| HAQ                        |       |       |       |       |       | x   |
| DLQI                       |       | x     |       |       |       |     |
| BASDAI                     |       | x     |       |       |       |     |
| ASQoL                      |       | x     |       |       |       |     |
| SF-36 PCS                  |       |       |       |       |       | x   |
| PsAQoL                     |       |       |       |       |       |     |
| **Laboratory assessment**  |       |       |       |       |       | x   |
| CRP                        | x     | x     |       |       |       |     |

**Table 1** Components in calculation of disease activity measures

ASQoL: Ankylosing Spondylitis Quality of Life; BASDAI: Bath Ankylosing Spondylitis Disease Activity Index; CPDAI: Composite Psoriatic Disease Activity Index; DAPSA: Disease Activity index for Psoriatic Arthritis; DAS28: Disease Activity Score 28; DLQI: Dermatology Life Quality Index; GRACE: GRAppa Composite ScoreE; LEI: Leeds Enthesitis Index; MDA: Minimal Disease Activity; PASDAS: Psoriatic Arthritis Disease Activity Score; PASI: Psoriasis Area and Severity Index; PsAQoL: Psoriatic Arthritis-specific Quality of Life; SF-36 PCS: Short Form 36 Physical Component Scale; VAS: visual analogue scale.
are described elsewhere [19]. In short, DEPAR collects data with the aim of investigating daily clinical practice of PsA patients. Patients with a new diagnosis of PsA are eligible to participate if they had not yet received treatment with disease-modifying antirheumatic drugs (DMARDs) for PsA before the first study visit. Written informed consent was obtained from all participants according to the Declaration of Helsinki. The study was approved by the local medical research ethics committee of Erasmus Medical Centre Rotterdam, the Netherlands (MEC-2012-549). For this analysis, we used data collected between August 2013 and April 2018.

Data collection

In the first year after diagnosis and inclusion in the study, data of patients were collected every 3 months in a study visit. Trained research nurses collected clinical data, including swollen joint count (SJC; 66 joints) and tender joint count (TJC; 68 joints), enthesis at clinical examination (Leeds Enthesitis Index, LEI [20], dactylitis count, physician global visual analogue scale (VAS), and psoriasis (Psoriasis Area and Severity Index, PASI [21]). Patients filled out questionnaires shortly before or after their visit to the research nurse. In DEPAR, multiple questionnaires are collected to measure patient-reported activity of disease and different outcomes. For this analysis we used the Short Form 36 (SF-36 [22]), HAQ [23], patient global, pain and skin VAS, Productivity Cost Questionnaire (PCQ [24]), Dermatology Life Quality Index (DLQI [25]), Ankylosing Spondylitis Quality of Life Questionnaire (ASQoL [26]), PsA-specific quality of life (PsAQoL [27]), and Bath Ankylosing Spondylitis Disease Activity Index (BASDAI [28]).

Disease activity

At each visit within the first year we calculated scores of DAS28-CRP [29], DAPSA [8], CPDAI [9], PASDAS [10], GRACE [11] and MDA [12]. DAS28-CRP is calculated, using the 28-joint tender and swollen counts, patient global VAS, and CRP level, as follows: DAS28-CRP = 0.56√TJC + 0.28√SJC + 0.36 ln(CRP + 1) + 0.014VAS + 0.96. DAPSA adds the SJC66, TJC68, VAS global (0–10 scale), VAS pain (0–10 scale) and CRP (mg/dl) in a total score. CPDAI assesses grades severity of involvement of five domains (joints, skin, entheses, dactylitis and spine) with a score of 0–3. Spinal disease activity (using BASDAI and ASQoL) was only assessed in patients with axial involvement according to their rheumatologist. Other domains were assessed using SJC, TJC, LEI, PASI, dactylitis count, HAQ, and DLQI. PASDAS is calculated as: PASDAS=(0.18√physician global VAS + 0.159√patient global VAS – 0.253√SF-36 PCS + 0.101 ln(SJC + 1) + 0.48 ln(TJC + 1) + 0.23 ln(LEI + 1) + 0.377 ln(dactylitis count + 1) + 0.102 ln(CRP + 1) + 2) × 1.5. GRACE score was calculated as: GRACE = (1 – AMDF) × 10, in which AMDF is the arithmetic mean of desirability function. In the AMDF, SJC, TJC, HAQ, patient global, pain and skin VAS, PASI and PsAQoL are transformed to a 0–1 score where 0 is completely unacceptable and 1 is normal. The AMDF is a weighted average of these eight scales. MDA is defined as meeting at least 5 out of 7 remission criteria: SJC ≤ 1, TJC ≤ 1, LEI ≤ 1, PASI ≤ 1, patient global VAS ≤ 20 mm, patient pain VAS ≤ 15 mm and HAQ ≤ 0.5.

In addition, at each visit patients were classified as having low disease activity according to the five continuous composite measures (DAS28 ≤ 3.2 [30], DAPSA ≤ 14 [31], CPDAI ≤ 4 [11], PASDAS ≤ 3.2 [11] and GRACE ≤ 2.3 [11]) and MDA (5/7 remission criteria). In cases where not all disease activity scores could be calculated, the visit was excluded from this analysis. DLQI and ASQoL were not collected in all patients at all visits, resulting in some exclusion by design.

Outcomes

The anchor-question of the SF-36 was used to distinguish categories of change in general health, which was reported by patients as either much improved, somewhat improved, stable, somewhat worsened or much worsened. HRQOL was determined using the SF-36 Physical Component Scale (PCS) and Mental Component Scale (MCS), which were calculated using the Dutch norm scores [32]. Work productivity was assessed using the PCQ every visit, in which patients are asked about work and productivity in the past 4 weeks. We determined employment status, absenteeism, working hours, and productivity loss at work (presenteeism) and productivity loss of unpaid work throughout the first year. Total productivity hours worked was calculated by subtracting hours of absence and productivity loss at work in hours from the total working hours. Productivity loss at work was calculated by multiplying the total hours of productivity loss by the percentage of productivity loss. Disability was assessed using the HAQ.

Statistical analysis

Responsiveness of each measure was compared in a distribution-based approach and an anchor-based approach. The former was done by comparing the effect size (ES, i.e. the difference between baseline and 1 year, divided by the s.d. of the baseline), standardized response mean (SRM, i.e. the difference divided by the s.d. of the difference). Change in disease activity from baseline to 1 year was calculated for all measures. We hypothesized that within the first year after diagnosis, the disease activity would decrease, which justifies the distribution-based approach. In the anchor-based approach we compared change of each measure over 3 months in patients reporting improvement, worsening or stable general health (as determined in the anchor question of the SF-36). The relation between low disease activity according to different disease measures and patient-reported outcomes each 3 months in the first year was assessed using mixed effects models. The variables time and disease activity measure were included in the fixed-effects part, and random intercepts and random slopes were included in the random-effects part. Outcomes were SF-36 PCS, SF-36 MCS, productivity (linear mixed effects models).
and HAQ > 0.5 (mixed-effects logistic regression). Relative model fits were compared using the Akaike information criterion, using the fit of disease activity measure DAS28 as reference. Analyses were performed in STATA 15.1 (StataCorp, College Station, TX, USA) and R 3.4.2 (R Foundation for Statistical Computing, Vienna, Austria).

Results

Of the 549 patients included until April 2018, 331 had all disease activity scores available at baseline and were included in this analysis. Excluded patients (n = 218) had missing questionnaires (21%, Supplementary Fig. 1, available at Rheumatology online). Mean age was 50.7 (S.D. 13) years, median symptom duration was 0.9 (interquartile range (IQR) 0.3–2.8) years and 171 (52%) were male. At baseline, median swollen joint count was 2 (IQR 1–4), median tender joint count was 3 (IQR 1–7), and median PASI score was 2 (IQR 0.5–4.0, Table 2).

Responsiveness

Figure 1 shows average disease activity scores throughout the first year of patients with complete data. Though the average DAS28 (green), CPDAI (grey) and GRACE (blue) scores were similar at twelve months, their initial scores and evolvement differed over the first year. Disease activity was high at baseline and low at 1 year according to DAS28 in 65 (34%), DAPSA in 76 (40%), GRACE in 63 (33%) and MDA in 66 (35%).

In the distribution-based assessment of responsiveness (i.e. ES and SRM in the first year, Table 3), the PASDAS was the most responsive, as shown with the highest ES (1.00, s.d. 1.05) and highest SRM (0.95, s.d. 1.00). The DAPSA was the least responsive, with an ES of 0.73 (s.d. 1.04) and SRM 0.71 (s.d. 1.00). The responsiveness of DAS28 (ES 0.88, s.d. 1.05; SRM 0.83, s.d. 1.00), CPDAI (ES 0.88, s.d. 1.08; SRM 0.82, s.d. 1.00) and GRACE (ES 0.75, s.d. 0.90; SRM 0.83, s.d. 1.00) were similar and all better than that of DAPSA, but worse than that of PASDAS.

In the anchor-based assessment of responsiveness, improvement in disease activity score was related to change in general health as assessed with the anchor-question of the SF-36 of the first 3 months. Of the 265 patients with complete baseline and 3 months’ assessments, general health as compared with 3 months ago was much improved in 39 patients (15%), somewhat improved in 78 (29%), stable in 100 (38%), somewhat worsened in 41 (15%) and much worsened in 7 patients (3%, Table 4). The improvement in disease activity scores in these five categories of change in general health are shown in Table 4 and Fig. 2 (much improved on the left to much worsened on the right). DAS28 and DAPSA differentiated much improved from other states [mean change 1.42 (95% CI: 1.12, 1.73) for DAS28 and 13 (95% CI: 10.3, 15.6) for DAPSA], but did not differentiate between somewhat improved, stable, somewhat worsened and much worsened. CPDAI did not differentiate between any of the general health states. Both PASDAS and GRACE differentiated between much improved [mean change 1.86 (95% CI: 1.51, 2.21) for PASDAS and 1.78 (95% CI: 1.42, 2.14) for GRACE], somewhat improved [mean change 1.06 (95% CI: 0.86, 1.26) for PASDAS and 0.97 (95% CI: 0.74, 1.20) for GRACE], and stable [mean change 0.36 (95% CI: 0.20, 0.52) for PASDAS and 0.34 (95% CI: 0.19, 0.48) for GRACE], but not between somewhat worsened and much worsened.

Longitudinal associations with outcomes

The associations between the longitudinal evolvement of disease activity measure and longitudinal evolvement of outcomes of SF-36 PCS, SF-36 MCS, productivity and HAQ were assessed using mixed effects models. To compare longitudinal validity of measures relative to each other, model fits of the mixed models were compared with the Akaike information criterion relative to DAS28 (a lower Akaike information criterion corresponds to a better fit, Supplementary Table 1, available at Rheumatology online). For SF-36 PCS, the longitudinal evolution had the highest association with low disease activity according to the GRACE, followed by PASDAS, MDA, DAPSA, DAS28, with the lowest association for CPDAI. Regarding productivity, the association was highest for GRACE and PASDAS as well, followed by DAPSA, MDA, CPDAI and DAS28. The evolution of SF-36 MCS was poorly related to any disease measure and model fit was comparable.

Table 2 Clinical characteristics at baseline (n = 331)

| Characteristic | Value |
|---------------|-------|
| Age, mean (s.d.), years | 50.7 (13) |
| Male, n (%) | 171 (52) |
| Symptom duration, median (IQR), years | 0.9 (0.3–2.8) |
| Swollen joint count (66), median (IQR) | 2 (1–4) |
| Tender joint count (68), median (IQR) | 3 (1–7) |
| LEI > 0, n (%) | 130 (39) |
| LEI if positive, median (IQR) | 2 (1–3) |
| LDI > 0, n (%) | 50 (15) |
| PASI, median (IQR) | 2 (0.5–4.0) |
| VAS Global, mean (s.d.) | 46 (26) |
| VAS Pain, mean (s.d.) | 46 (26) |
| HAQ, median (IQR) | 0.6 (0.4–1.0) |
| DAS28, mean (s.d.) | 3.1 (1.1) |
| DAPSA, mean (s.d.) | 18 (11) |
| CPDAI, mean (s.d.) | 3.9 (1.9) |
| PASDAS, mean (s.d.) | 4.1 (1.2) |
| GRACE, mean (s.d.) | 3.4 (1.5) |
| MDA, mean (s.d.) | 51 (15) |

CPDAI: Composite Psoriatic Arthritis Disease Activity Index; DAPSA: Disease Activity Index for Psoriatic Arthritis; DAS28: Disease Activity Score 28; GRACE: GRAppa Composite ScorE; IQR: interquartile range; LEI: Leeds Enthesitis Index; LDI: Leeds Dactylitis Index; MDA: Minimal Disease Activity; PASDAS: Psoriatic Arthritis Disease Activity Score; PASI: Psoriasis Area and Severity Index; VAS: visual analogue scale.
Regarding the outcome of disability (HAQ > 0.5), the order of association (high to low) was MDA, CPDAI, GRACE, DAPSA, PASDAS and DAS28. A summary of the relative performance for each aspect of responsiveness and longitudinal validity as discussed before is shown in Supplementary Table 2, available at Rheumatology online.

**Discussion**

In this study, we compared responsiveness and longitudinal validity of DAS28, DAPSA, CPDAI, PASDAS, GRACE and MDA in current usual care of newly diagnosed PsA patients. Using the change of disease activity over the first year as expressed in the ES and SRM, responsiveness of PASDAS was highest and that of DAPSA was lowest. Using change in general health status as anchor, all measures except CPDAI were able to discriminate patients that reported improved or stable general health. Stable and worsened general health, however, could only be discriminated by GRACE and PASDAS. Longitudinal evolution of HRQOL and productivity was best captured with GRACE, PASDAS and MDA and less by DAPSA.
A similar relation was seen with the outcome of disability using the HAQ, but with the difference that CDPAI had the second-best relation with HAQ.

This study is the first to assess responsiveness and longitudinal validity of all composite measures in usual care of early disease. Responsiveness has been studied using data from a trial by Helliwell and Kavanaugh; they reported that PASDAS, AMDF (i.e. the GRACE score in a different form) and DAS28 had the highest and similar responsiveness, while a modified CDPAI and DAPSA showed lower responsiveness [14]. This confirms part of our findings. They, however, analysed data from a trial including patients with active disease and a predominantly polyarticular phenotype. This patient selection probably explains why the articular measure DAS28 had a better responsiveness with higher ES and SRM than we
observed. Regarding validity, the relation of a disease measure and HRQOL has been studied in cross-sectional studies for MDA: patients in MDA report better HRQOL than patients not in MDA [33, 34]. This study is the first to test the validity longitudinally, by assessing the evolvement of outcomes in relation to disease activity over time.

The superiority of PASDAS, GRACE and MDA in terms of longitudinal validity could be attributed to their multidimensionality, but also to their use of more extensive questionnaires than for VAS scores alone. These questionnaires of HRQOL and disability used in PASDAS, GRACE and MDA are already closely related to the outcomes used in our analysis, resulting in a better performance. Analysis of relative contribution of each component was outside the scope of this work, but we will discuss the relation between use of questionnaires in disease measures here. On the one hand, the composites measures using questionnaires of HRQOL are a better representation of impact of disease to guide treatment decisions. HRQOL also belongs to the core set of domains for the assessment of patients with PsA according to the OMERACT [35]. On the other hand, generic questionnaires of HRQOL are more likely to be affected by factors other than disease activity and its burden that they aim to measure. Comorbidities for example are known to influence both disease activity measures and outcomes [36–38]. Also, some have argued that the HAQ is influenced by structural damage as well, which would make it impossible for some patients to be in remission despite absence of active inflammation [7]. In our analysis of early disease, however, we suspect disability is mostly determined by inflammation. Moreover, not only are questionnaires influenced by other factors than active disease: an increase in acute phase reactants—considered to be an objective measure of disease activity—can have other causes than an increase in PsA activity. Regardless, a composite measure needs the interpretation of a physician, who can choose not to change treatment if a higher disease activity score has other causes than PsA activity. A higher specificity for low disease activity is in that case of greater importance than a higher sensitivity.

In this analysis we assessed responsiveness and longitudinal association with outcomes of different measures, but for adaptation in clinical practice feasibility needs to be considered as well. A less feasible measure will only be accepted for use in clinical practice if it has a sufficiently better performance. All measures have a joint assessment and some form of general assessment using a VAS score. The measures differ in terms of joint count (i.e. 28 or 66/68), use of acute phase reactants, assessment of other PsA manifestations and use of questionnaires other than VAS. In clinical practice, assessment of musculoskeletal disease activity should include all joints, presence of enthesis and presence of dactylitis. Psoriasis could be assessed with a body surface area instead of a PASI score. Acute phase reactants are often already measured, along with toxicity screening for DMARDs. The biggest feasibility problem will most likely be the questionnaires: CPDAI needs four questionnaires, GRACE needs two questionnaires and three VAS scores, and PASDAS needs one questionnaire and one VAS score. With increasing use and possibilities of electronic health records, and focus on value based healthcare, collecting patient-reported outcomes in clinical care will become more feasible. Regarding feasibility in our cohort, complete data of all measures were available in 65% of visits in our cohort, but it will probably be different in clinical practice. More data on feasibility of use of these measures in clinical practice are needed.

A strength of this study is that it tested the validity of composite measures in a usual care population, including patients with monoarthitis, oligoarthitis and other phenotypes besides polyarthitis alone, which is the population of interest when composite measures are to be used in clinical practice. Also, it is the first study testing these composite measures early in the course of disease, within the first year after diagnosis. We showed that the relative performance differed from responsiveness in a clinical trial, probably owing to a difference in disease history, disease activity and phenotype. The patients eligible to participate in clinical trials often have high disease activity, so the responsiveness is expected to be higher. Further, we tested all disease measures using clinical assessments and questionnaires as instructed by developers of the disease measure.

Our study has some limitations as well. With our choice of excluding patients with incomplete baseline data and follow-up visits with incomplete data (some by design), our sample size and power was reduced. Also, the estimate of the longitudinal relation between disease activity and outcomes itself might be biased, but not the performance relative to each other. Last, as discussed before, with the lack of a gold standard for disease activity we chose anchors and outcomes of general health as reference, while the SF-36 has not been established as a disease activity anchor. We hypothesized that within the first year of disease in which treatment for PsA is initiated, the majority of change in general health is the result of change in disease activity. We cannot rule out that in some patients the associations between general health and some disease measures is increased owing to factors other than disease activity.

In conclusion, the disease activity measures PASDAS and GRACE had the highest responsiveness, were best able to discriminate between different health states, and together with MDA were best related to longitudinal evolvement of HRQOL, productivity and disability. Responsiveness and longitudinal validity was inferior for the disease activity measures DAS28, DAPSA and CPDAI. Though MDA and DAPSA are recommended for use in clinical practice by an international task force, as an alternative to DAPSA we suggest use of the continuous measures PASDAS or GRACE.

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Supplementary data

Supplementary data are available at Rheumatology online.

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