Role of clinical and biochemical inflammation in structural progression of patients with psoriatic arthritis

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ABSTRACT

Objective To determine the contribution of clinical and biochemical inflammation to structural progression of patients with psoriatic arthritis (PsA).

Methods We analysed patients from the Infliximab Multinational Psoriatic Arthritis Controlled Trial 2 trial (infliximab vs placebo). We obtained total modified Sharp/van-der-Heijde Scores from baseline and year one images, and swollen joint counts (SJC) and levels of C reactive protein (CRP) throughout the second half of year 1 (5 measurements) from 74 placebo-treated patients. We computed radiographic progression, time-averaged SJC (taSJC) and CRP (taCRP) values and assessed their impact on structural progression by logistic regression analysis. We further categorised patients as ‘active’ (+) or ‘inactive’ (−) based on their taSJC (cut-off point: 2/66 joints) and taCRP (cut-off point: 0.5 mg/dL) and compared radiographic progression across three groups (double inactive, single active, double active).

Results ORs for progression were 1.24 (95 % CI 1.04 to 1.47; p=0.016) for taSJC and 6.08 (95 % CI 1.12 to 33.03; p=0.036) for taCRP. When predictors were dichotomised (+ vs −), differences were maintained between taSJC+ and taSJC− patients (1.05±3.21 and 0.56±2.30, respectively), as well as for taCRP+ vs taCRP− patients (1.14±3.23 and 0.05±2.37, respectively). Progression was intermediate in the presence of abnormalities of one but not the other inflammatory variable, indicating increasing radiographic progression with increasing inflammation (p=0.05).

Conclusion In patients with PsA, both clinical and biochemical inflammation have an impact on structural progression. Overall, progression is smallest in the absence of both clinical and biochemical inflammation, higher when either clinical or biochemical inflammation is present and highest with both clinical and biochemical inflammation.

Psoriatic arthritis (PsA) is a heterogeneous inflammatory musculoskeletal disease that can affect peripheral and axial joints, entheses and tendons in patients with psoriatic skin and often also nail disease; it is typically seronegative for autoantibodies. Structural impact of PsA includes damage of bone and cartilage, but also has an osteoproliferative component. Similar to rheumatoid arthritis (RA), disease activity is a driver of structural progression, and both clinical and biochemical inflammation may be good surrogates for this risk, although it is unclear how independent these effects are and how they interrelate. Here, we investigate whether clinical inflammation presenting as swollen joint counts (SJC), and biochemical inflammation, as mirrored by C reactive protein (CRP) levels, have independent roles with respect to structural progression of PsA, and whether there is an effect modification between the two.

METHODS

Patients and data

In this study, we conducted a secondary data analysis on patient data from the Infliximab Multinational Psoriatic Arthritis Controlled Trial (IMPACT) 2 assessing the efficacy of infliximab (INF) compared with placebo.
(PLC) in PsA. In this trial, the patients either received INF 5 mg/kg or PLC. Concomitant methotrexate treatment (dosage not provided to the authors) was allowed in both treatment arms, but not mandatory. All patients had active PsA (≥5 swollen joints, ≥5 tender joints and CRP levels of ≥1 mg/dL and/or morning stiffness of ≥45 min), active plaque psoriasis (≥1 plaque of ≥2 cm in diameter) at trial entry, and an inadequate response to previous treatment with disease-modifying antirheumatic drugs (DMARDs) and/or non-steroidal anti-inflammatory drugs.5

From the provided trial data, we extracted descriptive patient information (age, sex, disease duration), total modified Sharp/van-der-Heijde Scores (mTSS) based on radiographs performed at baseline and after 54 weeks, as well as CRP values and SJC throughout the study period.

To determine the 1-year radiographic progression, we subtracted the mTSS at baseline from the mTSS at week 54. Additionally, the calculated radiographic progression was dichotomised into two groups (non-progression vs progressor) using a mTSS cut-off of 1 (≤1: non-progressor; >1: progressor) to account for measurement and inter-reader error.6

We further extracted levels of SJC and CRP from multiple measurements and calculated time-averaged SJC (taSJC) and CRP (taCRP) to reflect the clinical and biochemical inflammation, respectively. Since all patients per inclusion criteria of the IMPACT 2 trial had to have five or more swollen joints and CRP levels of 1 mg/dL or higher, we calculated time-averaged levels of SJC and CRP over the second half of the first trial year (ie, including weeks 24, 30, 38, 46 and 54). In this way a separation of patients with persistent inflammation was possible from those without. Next, we divided patients depending on their taSJC and taCRP status into ‘active’ (+) or ‘inactive’ (−) using a cut-off of 2 for taSJC (≥2: active, taSJC+; <2: inactive, taSJC−), and of 0.5 mg/dL (upper limit of normal) for taCRP (≥0.5 mg/dL: active, taCRP+; <0.5 mg/dL: inactive, taCRP−). These cut-off points are different from those previously used in RA, which were 1 for both measurements, to account for the larger overall inflammatory load in PsA compared with RA.7–9 This approach also allowed for more balanced group sizes since data distribution for CRP and SJC was skewed. Based on these definitions, we for more balanced group sizes since data distribution for

**Validations**  
To validate the results from the clinical trial, we performed analyses on data of our clinical outpatient database of routinely followed patients (2005–2019), and calculated taSJC and taCRP, actives (+) and inactives (−) and structural progression (by mTSS) identically to the trial cohort used for the main analysis. The radiographs were routinely assessed by one trained reader. Statistical methods from the main analysis were applied in an analogous way.

All statistical preparations of the dataset and analyses were performed using SPSS Statistics V.27.0 (IBM).

**RESULTS**

**Patient characteristics**  
A total of 200 patients were enrolled in the IMPACT 2 trial (100 INF, 100 PLC). Due to drop-out or missing data, 55 patients were not considered for further analyses. The baseline characteristics of the excluded patients did not differ significantly from those completing the study (data not shown). The detailed characteristics of the remaining 145 patients (74 PLC, 71 INF) are shown in table 1.

**Analysis (clinical trial cohort)**  
In line with the previous findings, INF effectively suppressed radiographic progression (−1.16±3.96).5 Patients in the PLC arm progressed significantly more until week 54 (0.74±2.98; p=0.008) compared with INF. Only 2.8 % of patients randomised to INF progressed >1 in mTSS, whereas 17.6 % progressed in the PLC group (p=0.004) (figure 1). As our main aim was the evaluation of the contribution of clinical and biochemical inflammation to radiographic progression, the INF group was considered non-informative, and analyses were limited to patients treated with PLC±background methotrexate (n=74), where such progression signal was actually present.

We identified 21.6 % of taSJC+ and 19.1 % of taCRP+ patients as progressors (figure 2A and B). Despite the small overall progression signal, progression was significantly higher in patients with active compared with

**Analyses**  
We assessed progression using cumulative distribution plots (‘probability plots’) and compared progression statistically by non-parametric tests (Mann-Whitney U test). To differentiate the contribution of clinical and biochemical inflammation, a multivariable binary logistic regression was performed using progression/non-progression as the dependent variable, and taSJC, taCRP and their interaction as independent variables. OR with two-sided 95 % CIs were calculated.

Differences in 1-year radiographic progression were assessed for taSJC+ vs taSJC−, and taCRP+ vs taCRP− using the Mann-Whitney U test. To assess the effect of increasing states of inflammation (double inactive: taSJC−/taCRP−; single active: taSJC−/taCRP+, taSJC+/taCRP−; double active: taSJC+/taCRP+) on radiographic progression, we used Jonckheere’s trend test. For all analyses, a p=0.05 was considered statistically significant.
The stratified approach by 2×2 division by taSJC and taCRP active (+)/inactive (−) status resulted in relatively low numbers (taSJC+ /taCRP−: n=33, taSJC− /taCRP+: n=18, taSJC− /taCRP−: n=14, taSJC+ /taCRP: n=9), but the higher progression in active (+) compared with inactive (−) patients was numerically maintained for both taSJC (1.05±3.21 vs 0.56±2.30) and taCRP (1.14±3.23 vs 0.05±2.37).

When we assessed the importance of clinical inflammation in presence or absence of biochemical inflammation, and vice versa, the greatest progression was found in the subgroup of patients with both clinical and biochemical inflammation (1.47±3.35, n=33); the smallest progression was observed in those in whom clinical and biochemical inflammation were both absent (−0.43±0.54, n=9); if one of the two variables was active and the other inactive, progression was in between (0.32±2.84, n=32). Thus, increasing inflammatory states (no inflammation vs either clinical or biochemical inflammation vs both clinical and biochemical inflammation) resulted in increasing radiographic progression (p=0.050, figure 3A).

In the presence of clinical inflammation, biochemical inflammation had a numerical effect on radiographic progression (taSJC+ /taCRP−: 0.29±2.87, n=18; taSJC+ /taCRP+: 1.47±3.35, n=33); the same was true when evaluating the effect of clinical inflammation in the presence of biochemical inflammation (taSJC− /taCRP+: 0.37±2.92, n=14; taSJC− /taCRP+: 1.47±3.35, n=33). Further, in the absence of biochemical inflammation, numerically higher progression was seen in the presence of clinical inflammation (taSJC− /taCRP−: −0.43±0.54, n=9; taSJC+ /taCRP−: 0.29±2.87, n=18). Vice versa, in patients without clinical inflammation, higher progression was seen in...
the presence of biochemical inflammation (taSJC−/taCRP−: −0.43±0.54, n=9; taSJC−/taCRP+: 0.37±2.92, n=14). For all these comparisons, there was insufficient power to perform statistical analyses in the small stratified subgroups, but the data suggesting effects of clinical and biochemical inflammation were consistent.

Validation (real-life cohort)
A total of 105 patients were included in the validation analysis (mean age in years: 53.5±12.1; female: 47.6 %). Baseline mTSS, SJC and CRP were 26.0±47.6, 1.3±2.1 and 0.8±1.1, respectively. We included patients treated with or without biologic DMARD (bDMARD) treatment. On average, patients received bDMARDs 54.4 % of the time between two consecutive radiographs. The average duration between two consecutive radiographs was 1.8 years.

Using the same cut-off points as for trial data analysis, 37.1 % were identified as progressors (online supplemental figure 1). Division into subgroups again resulted in relatively small numbers (taSJC+/taCRP+: n=12, taSJC+/taCRP−: n=13, taSJC−/taCRP+: n=39, taSJC−/taCRP−: n=41). Nevertheless, all findings from the highly selected clinical trial cohort were reproduced in this clinical practice cohort, including increasing radiographic progression from the taSJC/taCRP double inactive group to the single active group and double active group (0.54±5.90, n=41; vs 2.52±6.49, n=52; vs 6.92±12.26, n=12, respectively; p=0.046; figure 3B). Also, all subgroup comparisons allowed deriving similar conclusions on the relevance of CRP activity in taSJC+ and taSJC− patients, and vice versa, based on the data from this real-life cohort with inclusion of bDMARDs: in the presence of clinical inflammation, patients with biochemical inflammation had higher radiographic progression than those without (taSJC+/taCRP−: 2.69±2.95, n=13; taSJC+/taCRP+: 6.92±12.26, n=12). The same was true for the impact of clinical inflammation in the presence of biochemical inflammation (taSJC−/taCRP+: 2.46±7.33, n=39; taSJC+/taCRP+: 6.92±12.26, n=12). Also, as seen in the clinical trial cohort, clinical inflammation led to a higher degree of structural damage in patients without biochemical inflammation (taSJC−/taCRP−: 0.54±5.90, n=41; taSJC+/taCRP−: 2.69±2.95, n=13) and vice versa, biochemical inflammation influenced radiographic progression in patients without clinical inflammation (taSJC−/taCRP+: 0.54±5.90, n=41; taSJC+/taCRP+:2.46±7.33, n=39).

**DISCUSSION**
PsA is a chronic inflammatory disease leading to structural damage and functional impairment. Therefore, determining predictors for structural progression is
crucial as this can lead to a better management of PsA patients. Several predictors for radiographic progression, such as CRP and SJC have been reported. However, the interplay between those factors—and in particular between clinical and biochemical inflammation—remained to be elucidated and thus has been the focus in our analyses. The link between inflammation and structural progression is evident also in PsA, although less than in RA. Here, we show that clinical inflammation and biochemical inflammation are independently related to structural deterioration in PsA. In a simple analysis classifying patients as active or inactive for SJC and CRP, we established that not only those active for both have the highest, and those inactive for both have the lowest progression, but also that those patients who are active by one and inactive by the other variable have intermediate progression. Hence, increasing states of inflammation (no inflammation vs clinical or biochemical inflammation vs clinical and biochemical inflammation) present a continuum leading to increasing radiographic progression. In an exploratory 2×2 analysis, even despite low numbers, we showed the numerical effects of SJC and CRP on progression, even if the other one is inactive. While CRP and SJC both reflect inflammation, the differences in associations of these markers with progression may lie in the use in the different cut-off points. This underpins both a similarity to and difference from RA: inflammation is closely linked to structural damage in both diseases, but clinical inflammation has been shown to be the major driver of radiographic progression RA, whereas biochemical inflammation appears to play a bigger role in PsA.

We used a clinical trial cohort to test the hypothesis and then validated the results derived from this highly selected group in a real-life population, where we observed similar trends.

One limitation of our study is the small sample size, particularly in the subgroup analyses, which have to be considered as exploratory. Given the limited sample size, lack of statistical significance does not rule out true difference (potential false negative results). Furthermore, the IMPACT 2 trial was published in 2005. However, despite their low number and the timespan since initial publication, data support the main conclusions about the relevance of both, clinical and biochemical inflammation, in PsA, and were fully validated in a second, independent real-life cohort. Thus, the data reveal the clinical inflammation, even in the absence of a biochemical response, is similarly critical for progression of joint damage in PsA as biochemical inflammation, but their combined effect is detrimental for structural joint integrity. Evaluation of joint swelling and radiographic progression on the joint level would have been an even more direct way to assess a relationship, but the joint level data were not available for us, and also the progression signal on the individual joint level is considered quite small. In summary, our data reveal that clinical inflammation as quantified by clinical examination of joints, as well as biochemical inflammation as measurable by CRP levels are both relevant targets for comprehensive management of PsA that aims at preserving structural integrity of joint, and thus also functional ability.

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