Effect of non-steroidal anti-inflammatory drugs on radiographic spinal progression in patients with axial spondyloarthritis: results from the German Spondyloarthritis Inception Cohort

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ABSTRACT

Objective To investigate the influence of non-steroidal anti-inflammatory drugs (NSAIDs) intake on radiographic spinal progression over 2 years in patients with ankylosing spondylitis (AS) and non-radiographic axial spondyloarthritis (SpA).

Methods 164 patients with axial SpA (88 with AS and 76 with non-radiographic axial SpA) were selected for this analysis based on availability of spinal radiographs at baseline and after 2 years of follow-up and the data on NSAIDs intake. Spinal radiographs were scored by two trained readers in a concealed randomly selected order according to the modified Stoke Ankylosing Spondylitis Spine Score (mSASSS) system. An index of the NSAID intake counting both dose and duration of drug intake was calculated.

Results High NSAIDs intake (NSAID index≥50) in AS was associated with lower likelihood of significant radiographic progression defined as an mSASSS worsening by ≥2 units: OR=0.15, 95% CI 0.02 to 0.96, p=0.045 (adjusted for baseline structural damage, elevated C reactive protein (CRP) and smoking status) in comparison with patients with low NSAIDs intake (NSAID index<50). This effect was most pronounced in patients with baseline syndesmophytes plus elevated CRP: mean mSASSS progression was 4.36±4.53 in patients with low NSAIDs intake versus 0.14±1.80 with high intake, p=0.02. In non-radiographic axial SpA, no significant differences regarding radiographic progression between patients with high and low NSAIDs intake were found.

Conclusion A high NSAIDs intake over 2 years is associated with retarded radiographic spinal progression in AS. In non-radiographic axial SpA this effect is less evident, probably due to a low grade of new bone formation in the spine at this stage.

Keywords: Axial spondyloarthritis | Non-steroidal anti-inflammatory drugs | Radiographic progression | Ankylosing spondylitis | Spondyloarthritis Inception Cohort (GESPIC)

Non-steroidal anti-inflammatory drugs (NSAIDs) are considered as a preferred therapy in patients with axial spondyloarthropathy (SpA), including ankylosing spondylitis (AS). However, it has been suggested that NSAIDs might have a good symptomatic and a disease-modifying effect. It had been shown in a small retrospective study by Boersma some time ago that a continuous use of phenylbutazone was associated with retardation of spinal ossification in AS. In a more recent study by Wanders et al, continuous (daily) use of NSAIDs was also associated with an inhibition of radiographic progression in the spine over 2 years as compared with on-demand use. However, these reports have not been confirmed until now. Furthermore, NSAIDs influence on radiographic progression in early axial SpA (especially in a non-radiographic form) was not investigated so far. This analysis of the 2-year data from the German Spondyloarthritis Inception Cohort (GESPIC) was aimed at investigating the influence of NSAIDs intake on radiographic progression of the spine in patients with AS and non-radiographic axial SpA with short disease duration.

METHODS

Patient selection

Patients included in GESPIC were required to have a definite clinical diagnosis of axial SpA according to the local rheumatologist. Patients were further classified based on radiographic findings as AS or as non-radiographic axial SpA. Patients with AS ought to fulfil the modified New York criteria and the duration of symptoms was restricted to ≤10 years at the time of inclusion. Patients with non-radiographic axial SpA ought to fulfil European Spondyloarthropathy Study Group criteria with minor modifications and had to have duration of symptoms of ≤5 years. The baseline data of this cohort have been recently reported elsewhere. Radiographs of the spine (lumbar and cervical spine) and sacroiliac joints were obtained at baseline and after 2 years of follow-up. The full sets of radiographs were available for 210 GESPIC patients (115 with AS and 95 with non-radiographic axial SpA) as reported elsewhere. Of these, information on NSAIDs intake over 2 years was available for 164 patients (88 with AS and 76 with non-radiographic axial SpA) who were naïve to antitumour necrosis factor (TNF) therapy and did not receive this therapy during 2 years of follow-up.

Radiographs and scoring

X-rays of sacroiliac joints and spine (cervical and lumbar spine at baseline and after 2 years of follow-up) were performed locally. Images were centrally collected, digitised, anonymised and subsequently scored independently by two trained readers (DP, HH). The readers scored radiographs in a concealed and randomly selected order and...
were blinded for all clinical data. Grading of sacroilitis was performed according to the established scoring system. Spinal radiographs were scored according to the modified Stoke Ankylosing Spondylitis Spine Score (mSASSS) scoring system. In addition to lateral views, anteroposterior views of the lumbar spine (left and right corners of the vertebral bodies from Th12 to S1) were scored for the presence of syndesmophytes.

**NSAIDs intake**

Data on NSAIDs intake (dose and frequency of intake) were collected at baseline and every 6 months thereafter during 2 years of follow-up. An index of the NSAID intake, as recommended by ASAS (Assessment of SpondyloArthritis international Society), was calculated. High NSAIDs intake was defined as a mean NSAIDs intake index ≥50, and low NSAIDs intake as a mean NSAIDs intake index <50.

**Statistics**

Status scores for baseline and follow-up were calculated by the mean of the scores of both readers. Radiographic progression was calculated by the difference between follow-up and baseline mean values. Agreement between readers was estimated by the intraclass correlation coefficient (ICC). Significant radiographic progression was defined as a worsening of the mSASSS score by ≥2 units over 2 years. Syndesmophytes were considered to be absent or present if both readers agreed on it.

**RESULTS**

The mean NSAID intake index over 2 years was 33.7±28.0 (range 0–100) in the AS and 52.2±26.7 (range 0–100) in the non-radiographic axial SpA group. Overall, 24 with AS (27.3%) and 19 patients with non-radiographic axial SpA (25.0%) had a high NSAIDs (NSAID intake index ≥50) intake during 2 years of follow-up. Comparative characteristics of patients with low and high NSAIDs intake are presented in table 1. As shown in the table, patients with a high NSAIDs intake had at baseline higher clinical disease activity (as measured by the Bath Ankylosing Spondylitis Disease Activity Index) and higher level of functional impairment (as measured by the Bath Ankylosing Spondylitis Functional Index) as compared with patients with low NSAIDs intake. Other parameters were similar in the subgroups. Selective inhibitors of cyclo-oxygenase-2 (COX-2 inhibitors: celecoxib, etoricoxib, rofecoxib, ibuprofen).

**Ethical approval**

The study protocol was approved by the central ethical committee of the coordinating centre (Charité Universitätsmedizin Berlin, Berlin, Germany) and by all local ethical committees of the participating centres. Written informed consent was obtained from all patients.
Clinical and epidemiological research

valdecoxib), as opposed to non-selective COX inhibitors, were used in similar proportions of patients with high and low NSAIDs intake: 29.2% and 32.1%, respectively, in AS; 42.1% and 33.3%, respectively, in non-radiographic axial SpA. Among non-selective COX inhibitors, diclofenac, ibuprofen, indomethacin and piroxicam were used most frequently and accounted for more than 50% of all NSAIDs used.

There was a very good agreement between the two readers regarding status scores (mSASSS) of spinal radiographic damage: the ICC was 0.92 (95% CI 0.90 to 0.94) at baseline and 0.92 (95% CI 0.89 to 0.94) at year 2. The agreement regarding the mSASSS change score was moderate with an ICC coefficient of 0.33 (95% CI 0.19 to 0.49).

The cumulative probability plot presented in figure 1A demonstrates the divergence of mSASSS change on the patient level with less radiographic progression in the group of AS patients with high NSAIDs intake. The mean mSASSS change over 2 years in AS was 0.02±1.39 in patients with high NSAIDs intake versus 0.96±2.78 in the subgroup with low NSAIDs intake, respectively; p=0.142 (after adjustment for radiographic status at baseline p=0.22) (figure 2A). The trend for reduced radiographic spinal progression in patients with high NSAIDs intake was also present if other thresholds for the index of NSAIDs intake were chosen (eg, tertiles). In AS patients with an NSAID intake index values of <33, 33–65 and ≥66, the mSASSS changes over 2 years were 0.86±2.93, 0.75±1.84 and 0.10±1.70 units, respectively.
Similarly, less AS patients in the group with high NSAIDs intake showed radiographic spinal progression defined as a worsening of the mSASSS score by 2 units and more over 2 years in comparison with the patients with low NSAIDs intake: 8.5% (n=2) versus 21.9% (n=14), respectively, p=0.142; unadjusted OR=0.33 (95% CI 0.07 to 1.55), p=0.159. After adjustment of the OR for factors shown to be independently associated with radiographic spinal progression (mSASSS worsening by ≥2 units) in GESPIC previously (presence of radiographic damage at baseline, elevated acute phase reactants (C reactive protein (CRP)), smoking), the association of retarded radiographic spinal progression with high NSAIDs intake became significant: OR=0.15 (95% CI 0.02 to 0.96), p=0.045 (table 2). Similar results were obtained after adjustment for the elevated (>20 mm/h) erythrocyte sedimentation rate at baseline instead of CRP. OR for an association of the spinal progression with NSAIDs intake of 0.15 (95% CI 0.02 to 0.94), p=0.042.

The positive effect of the NSAIDs intake was mostly evident in patients who were at high risk for radiographic progression due to the presence of syndesmophytes and elevated acute phase reactants. mSASSS progression over 2 years in AS patients with normal (≤6 mg/l) time-averaged CRP over 2 years was 0.23±1.58 and 0.05±1.41 in the groups with low (n=32) and high (n=11) NSAIDs intake, respectively (p=0.54). In contrast, in AS patients with elevated time-averaged CRP (>6 mg/l), the mSASSS progression was 1.69±3.48 versus 0.0±1.41 in patients with low (n=32) and high (n=13) NSAIDs intake, respectively (p=0.11). Similarly, in AS patients without syndesmophytes at baseline mSASSS progression was 0.32±1.31 versus −0.04±0.97 with low (n=47) and high (n=13) NSAIDs intake, respectively (p=0.52), while in patients with syndesmophytes at baseline the progression was 2.74±4.58 (n=17) versus 0.09±1.80 (n=11), respectively, p=0.076. The difference was even clearer (and statistically significant) in patients with both syndesmophytes and elevated time-averaged CRP: 4.36±4.53 versus 0.14±1.80 in patients with low and high NSAIDs intake, respectively (p=0.02) (figure 3A), while there was nearly no progression in patients without both risk factors (0.55±1.07 vs 0.07±1.06, respectively; p=0.48) (figure 3B) or if only one risk factor was present: 0.17±1.95 versus −0.10±1.39 in patients with low (n=27) and high (n=10) NSAIDs intake, respectively (p=0.60).

There was also a statistically non-significant trend towards lower frequency of new syndesmophytes in AS patients with high NSAIDs intake in comparison with the group with low NSAIDs intake: 12.5% (n=3) versus 15.6% (n=10), respectively.

No clear association was found between radiographic spinal progression and NSAIDs intake if the NSAID dose only or duration of the NSAID intake only were taken into account (data not shown). COX selectivity was not identified as a factor influencing radiographic progression (data not shown). No differences in the proportion of patients who underwent physiotherapy could be found while comparing high and low NSAIDs users, as well as patients with and without radiographic progression (data not shown).

There was no clear and consistent difference between low and high NSAIDs intake patients in non-radiographic axial SpA: observed mSASSS change score was 0.51±1.72 versus 0.74±1.95 in patients with low and high NSAIDs intake, respectively; p=0.72 (after adjustment for radiographic status at baseline p=0.58) (figure 2B). Similar results were seen with the NSAIDs intake index values of <33, 33–65 and ≥66: mSASSS change was 0.60±1.70, 0.59±2.26 and 0.41±0.58, respectively; p=0.952. Only 5.3% (n=1) of the patients with high NSAIDs intake and 10.5% (n=6) with low NSAIDs intake showed an mSASSS worsening by ≥2 units over 2 years, p=0.492; OR=0.47 (95% CI 0.05 to 4.20), p=0.501. After adjustment for baseline radiographic damage, elevated acute phase reactants and smoking, this association remained statistically non-significant: OR=0.60 (95% CI 0.06 to 6.61), p=0.679 (table 2). New syndesmophytes occurred in 5.5% (n=1) versus 3.5% (n=2) of the patients with high and low NSAIDs intake, respectively.

Regarding progression of radiographic sacroiliitis the following results were obtained: in the AS group 0 patients with high NSAIDs intake and 14.1% (n=9) with low NSAIDs intake showed progression of sacroiliitis by at least one grade over 2 years in the opinion of both readers (p=0.053). In the non-radiographic axial SpA group, 15.8% (n=3) of the patients with high NSAIDs intake and 17.5% (n=10) demonstrated such a progression (p=0.86).

**DISCUSSION**

In the current study, we present data showing that NSAIDs retard radiographic spinal progression as assessed by an mSASSS change over 2 years in patients with AS confirming, therefore, earlier data. Although the clear difference in the absolute mSASSS change between subgroups with high and low NSAIDs intake was statistically non-significant (probably due to a relatively low sample size and high variation of the radiographic progression), an analysis of significant radiographic progression (ie, mSASSS worsening in 2 units and more over 2 years) adjusted for factors predictive for radiographic spinal progression, especially for the presence of structural damage at baseline, revealed a significant association between high NSAIDs intake and radiographic progression in AS (OR=0.15, 95%
CI 0.02 to 0.96, p=0.045: table 2). Adjustment for baseline factors such as baseline damage, smoking and CRP was important because there was heterogeneity among the groups at baseline and because these factors have been shown to be predictors of progression. It was remarkable to see a retarding effect of NSAIDs particularly in the group of AS patients with high NSAID intake because these patients had a higher proportion of syndesmophytes (table 1) – the strongest predictor of progression as compared with smoking and CRP – and, therefore, were more likely to progress compared with patients with low intake of NSAIDs who had less frequently syndesmophytes.

In AS patients with normal CRP and/or no syndesmophytes at baseline there was nearly no radiographic progression independently from the NSAIDs intake. In contrast, the effect of high NSAIDs intake was most pronounced in patients with both syndesmophytes at baseline and elevated CRP, suggesting that the observed NSAID effect on radiographic progression is mostly driven by this subgroup, which constitutes 25% of the whole AS group in the current study. These data indicate that continuous NSAIDs intake might especially be of benefit in these patients.

In non-radiographic axial SpA, however, this effect could not be shown which is most likely related to a relatively low radiographic spinal progression rate in this subgroup in general and nearly no new bone formation (syndesmophytes) in particular. Furthermore, in this group the mSASSS might be more driven by erosive changes than by new bone formation.

Interestingly, there was also a trend towards reduced radiographic sacroiliitis progression in patients with AS and high NSAIDs intake, but this was not the case in non-radiographic axial SpA. We speculate that this could be related to the interference of the NSAIDs with the new bone formation, which is mostly responsible for radiographic progression in advanced sacroiliitis, as opposed to the destructive/erosive change at the early disease.

In this analysis we used the recently introduced index of NSAIDs intake.9 This index accounts for dose and duration of NSAIDs intake which seem to be both relevant for retardation of radiographic spinal progression, since no clear differences in radiographic progression could be found if dose or duration of intake only were taken into account. The analysis with continuous variables (mean NSAIDs index over 2 years and mSASSS change score) demonstrated only a weak and statistically non-significant negative correlation between NSAIDs index and radiographic progression indicating a non-linear character of the relationship. Therefore, a clinically relevant threshold (NSAIDs intake index of 50) was chosen for this analysis to differentiate between low and high NSAID intake. Other thresholds (eg, tertiles) provided comparable results with the clearest effect of NSAIDs seen in the highest tertile, but the resulting patient numbers per group were too small to show significant differences.

Earlier investigations indeed already indicated that NSAIDs might have an inhibitory effect on new bone formation. There are several observational studies indicating a retardation of fracture healing15–16 or loosening of the hip endoprosthesis17 related to NSAIDs use. Furthermore, NSAIDs have been used for the prevention of heterotopic ossification after orthopaedic surgery, for example, total hip arthroplasty,18–25 hip resurfacing16 or fractures (eg, acetabular fractures).27 28

The observed inhibition of new bone formation by NSAIDs can probably best be explained by the inhibition of prostaglandins (especially prostaglandin E2) synthesis mediated by COX-2.29 Prostaglandin E2 is able to stimulate new bone formation by causing vasodilatation and by promoting angiogenesis.30 Prostaglandins also support blood supply to the site of new bone formation by increasing the replication and differentiation of osteoblasts.30

In experiments with COX-2 knockout mice, healing of the stabilised tibia fracture was delayed in comparison with wild-type animals and to COX-1 knockouts.31 Similarly, NSAIDs were able to retard a bone morphogenetic protein 7 induced ectopic
bone formation in an experimental mouse model indicating an important role of COX-mediated prostaglandin synthesis in new bone formation.\textsuperscript{34} Since retardation of the new bone formation is related to COX-2 inhibition, no substantial differences are expected between different NSAIDs, because in therapeutic concentrations all NSAIDs, independently from their COX-selectivity, inhibit COX-2 to nearly the same extent.\textsuperscript{35} NSAIDs are considered as a preferred therapy in patients with axial SpA due to their high symptomatic activity.\textsuperscript{3} In addition, small but significant decrease of CRP has been seen in AS patients treated with NSAIDs.\textsuperscript{36,37} If the structure-modifying effect of NSAIDs in AS is confirmed, it would be an additional argument to administer NSAIDs more, especially in patients with clinical indications for NSAIDs and in patients who are at risk for radiographic spinal progression such as presence of syndesmophytes and/or an elevated level of acute phase reactants. The potential cardiovascular, gastrointestinal and other side effects of continuous NSAIDs intake have been investigated in the greatest detail and we have argued recently that the benefit of such a treatment normally outweighs the risk in AS.\textsuperscript{38} The data presented here add further evidence to this.

Due to the character of our study there are some limitations which might have influenced the final results. GESPIC is an observational cohort and includes therefore a relative heterogeneous (in comparison with ‘standard’ drug investigating clinical trials) patient population, which was included based on diagnosis and symptom duration without further limitations. On the other hand, this comes closer to daily clinical practice. GESPIC was not specially designed and powered for the investigation of the influence of NSAIDs on radiographic progression. The NSAIDs doses and duration of intake were not strictly counted, but relied on the information given by the patient every 6 months, and represent therefore the best possible approximation. Finally, the treatment with NSAIDs was heterogeneous with a variety of NSAIDs used, but again this is closer to daily clinical practice.

In conclusion, we add further evidence here that NSAIDs seem to retard new bone formation in AS patients. Given the good anti-inflammatory capacity of TNF blockers in AS but their failure in stopping new bone formation, a trial combining TNF blockers with a variety of NSAIDs used, but again this is closer to daily clinical practice.

Contributors All the authors fulfil the authorship criteria. Contributors not fulfilling the authorship criteria are listed in the acknowledgement part.

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