Risk of septic arthritis in patients with rheumatoid arthritis and the effect of anti-TNF therapy: results from the British Society for Rheumatology Biologics Register

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ABSTRACT

Objectives To evaluate the risk of septic arthritis (SA) in patients with rheumatoid arthritis (RA) treated with anti-tumour necrosis factor (TNF) therapy.

Methods Using data from the British Society for Rheumatology Biologics Register, a prospective observational study, the authors compared the risk of SA between 11 881 anti-TNF-treated and 3673 non-biological disease-modifying antirheumatic drug (nbDMARD)-treated patients.

Results 199 patients had at least one episode of SA (anti-TNF: 179, nbDMARD: 20). Incidence rates were: anti-TNF 4.2/1000 patient years (pyrs) follow-up (95% CI 3.6 to 4.8), nbDMARD 1.8/1000 pyrs (95% CI 1.1 to 2.7). The adjusted HR for SA in the anti-TNF cohort was 2.3 (95% CI 1.2 to 4.4). The risk did not differ significantly between the three agents: adalimumab, etanercept and infliximab. The risk was highest in the early months of therapy. The patterns of reported organisms differed in the anti-TNF cohort. Prior joint replacement surgery was a risk factor for SA in all patients. The rate of postoperative joint infection (within 90 days of surgery) was 0.7%. This risk was not significantly influenced by anti-TNF therapy.

Conclusions Anti-TNF therapy use in RA is associated with a doubling in the risk of SA. Physicians and surgeons assessing the RA patient should be aware of this potentially life-threatening complication.

INTRODUCTION

Septic arthritis (SA) is a serious medical condition that, even with prompt treatment, can lead to irreversible joint damage and has a death rate of around 10%.1 The incidence of SA in the general population is around 4–10 per 100 000 patient years (pyrs) and seems to be rising,2 3 probably due to the combination of an ageing population and larger numbers of orthopaedic interventions.

Important risk factors for SA include increasing age, joint prosthesis, skin infection and pre-existing joint damage.4 5 Patients with rheumatoid arthritis (RA) may have many of these risks combined with the use of immunosuppressive medications. The risk of SA in an RA patient, irrespective of therapy, is increased by 4–15-fold.5 6 Although one might expect immunosuppressive therapy to increase the risk of SA, this has not been well studied. This question has been of increasing interest over the last decade since the emergence of biological therapies. Anti-tumour necrosis factor (anti-TNF) therapies were the first class of biological agents to become established in routine RA care. Data have emerged suggesting that these drugs confer a small but significant risk of serious infections, especially during the first months of treatment.7–9 It is also apparent that this risk differs by anatomical site and that there is increased susceptibility to certain pathogens.9–11

There is very limited information regarding the effect of anti-TNF therapy on the risk of SA. Case reports have described patients on anti-TNF therapy developing SA as a multifocal disease or with unusual causative organisms.12–14 Although case reports are a useful tool for raising questions, they cannot provide information regarding disease incidence or relative risk. An additional important question relates to the risk of SA following joint replacement surgery in anti-TNF-treated patients.

In 2001, the British Society for Rheumatology (BSR) established a national prospective cohort study of patients starting anti-TNF therapy for RA, the BSR Biologics Register (BSRBR). This is the largest register of its kind worldwide and includes detailed records of serious adverse events including SA occurring in patients receiving anti-TNF therapy as well as in a cohort of RA patients not exposed to anti-TNF therapy.

Our primary aim was to test the hypothesis that anti-TNF therapy increases the risk of SA compared with non-biological disease-modifying anti-rheumatic drug (nbDMARD) therapy. Secondary analysis considered whether anti-TNF therapy confers additional risk to patients who have joint replacement surgery either prior to starting therapy or during follow-up.

METHODS

The study commenced in 2001 alongside national recommendations within the UK that all RA patients prescribed anti-TNF therapy should be enrolled with the register.15 Patients were recruited to the anti-TNF cohort from 2001 onwards. Three anti-TNF agents were licensed for use in the UK during this period, with infliximab (INF) and etanercept (ETN) being available from the start of the study, while the third drug, adalimumab (ADA), came into clinical practice in 2003. Recruitment targets of 4000 patients for the ETN cohort were met in 2005, for INF in 2007 and for ADA in 2008. Before recruitment targets were met, it was estimated that over 80% of anti-TNF-treated patients with RA in the UK were registered on the BSRBR.15 A comparison
cohort of patients with active RA (defined as having a 28-joint count disease activity score (DAS28) >4.2) was recruited in parallel. These patients were receiving an nbDMARD and were biologically naive. Patients prescribed biologics were recruited from across the UK (over 250 hospitals) whereas controls were recruited from 29 centres. These control centres reflect a combination of secondary and tertiary care rheumatology centres distributed across the UK and are listed in full in the BSRBR control centre consortium supplementary data file.

Baseline assessment
All patients in this study had a physician diagnosis of RA. Baseline information included demographics, disease duration, a measure of self-reported physical function (the Health Assessment Questionnaire (HAQ))\(^{15}\), DAS28 score,\(^{18}\) baseline steroid use, smoking history, baseline comorbidity and surgery, including prior joint replacement. For the purpose of this analysis, we have considered only large joint replacements (shoulder, elbow, hip and knee) because when reviewing reports of small joint surgery, it was difficult to distinguish between soft-tissue procedures and arthroplasty.

Follow-up
Follow-up information was collected from three sources. First, six monthly questionnaires were sent to the treating rheumatologist for 3 years and annually thereafter. Second, data were collected directly from the patients six monthly for their first 3 years of the study. Patients were provided with a handheld diary card to record details of all hospital attendances as well as new prescriptions. Details of joint replacements during follow-up were collected from these two sources. Third, all patients were flagged with the UK National Health Service Information Centre, which informs the register of any deaths and the causes of deaths. All patients had to have at least one returned consultant follow-up questionnaire prior to 31 December 2009.

Case definition and verification
This analysis was confined to serious cases of SA. ‘Serious’ infections were defined as those requiring intravenous antibiotics or hospitalisation, or those resulting in death. All serious infections reported to the BSRBR were followed up with requests for additional information from the treating clinician to gather information on the site of infection and microbiology results. All reported cases of SA were then verified by a BSRBR clinician (JBG).

Events were ascribed to anti-TNF if they occurred while the patient was receiving anti-TNF therapy or within 90 days of the first missed dose. Events were attributed to the most recent drug exposure in patients who switched anti-TNF therapy. Patients were censored from further follow-up after their first episode of SA. Only one case of articular Mycobacterium tuberculosis was reported to the BSRBR, which has been described in a previous publication and is not included in this analysis.\(^{19}\)

Statistical methods
Crude incidence rates were calculated as the number of episodes of SA per 1000 pyrs of follow-up. Cumulative hazards were compared between the different cohorts using a Nelson–Aalen plot. A Cox model was used to calculate HR between the groups. Changes to the incidence rate over time in the anti-TNF cohort were analysed using a spline model. Potential confounders were identified prior to the analysis as variables that were either unbalanced between the nbDMARD and anti-TNF cohorts (age, gender, disease duration, DAS28, HAQ, steroid exposure, prior joint replacement, calendar year of entry into the study) or significant predictors of infection (chronic obstructive pulmonary disease, diabetes). Adjustment for these potential confounders was made using propensity scores. The use of this method in observational studies has been described in detail previously.\(^{20}\)

In a secondary analysis, the influence of joint replacement was examined. Patients were entered into this analysis when they were at risk of a prosthetic infection (ie, from baseline if they had a prior joint replacement or at the date of surgery if they had a joint replacement during follow-up). An adjusted Cox proportional hazards model was used to compare rates between groups. The risk of postoperative infection following joint replacement surgery during active follow-up was analysed separately using adjusted logistic regression to compare the odds of developing an infection within 90 days of joint replacement surgery between anti-TNF and nbDMARD cohorts. Adjustment was made for the same confounders as identified in the primary analysis. Missing baseline data were replaced using multiple imputations. All analysis was done using Stata 10.1 (StataCorp, College Station, Texas, USA).

Further details of the statistical methodology are presented in a data supplement online.

Ethics approval for this study was obtained in December 2000 from the Multicentre Research Ethics Committee for the Northwest of England.

RESULTS
The baseline characteristics of the 15 554 patients included in the analysis are shown in table 1. Although both anti-TNF and nbDMARD groups had active RA, there were significant differences at baseline. The patients receiving anti-TNF therapy were younger and proportionally more were women. The anti-TNF group had significantly longer disease duration as well as higher disease activity. History of joint replacement was higher in the anti-TNF cohort (24% vs 14% (p<0.001)).

Incident SA was reported in 199 patients during the follow-up period (179 anti-TNF; 20 nbDMARD). In univariate analysis, increasing age, longer disease duration, higher HAQ, higher DAS28 score, baseline steroid exposure, prior orthopaedic surgery and diabetes were associated with SA irrespective of anti-TNF exposure. The strongest association was with a history of large joint orthopaedic surgery at baseline irrespective of whether the SA developed in a prosthetic joint: HR 2.45 (1.90–3.17). The incident rate for SA was significantly higher in the anti-TNF cohort (4.2/1000 follow-up pyrs) than in the nbDMARD group (1.8/1000 pyrs) (table 2). The cumulative incidence over time is presented in figure 1. The unadjusted HR for SA in the anti-TNF cohort was 2.5 (95% CI 1.6 to 4.0). After full adjustment, patients on anti-TNF therapy were more than twice as likely to develop SA as the nbDMARD controls: adjusted HR 2.3 (95% CI 1.2 to 4.4).

Hazard estimates for the first year of follow-up were increased in all three anti-TNF agents; however, beyond 1 year, the hazard in the ETN cohort increased more than in the ADA and INF cohorts (figure 1). The adjusted HR was highest within the ETN cohort and lowest for ADA (table 2) although there were no differences between the three individual drugs or between the monoclonal antibody class and ETN that achieved statistical significance (results not shown).

The pattern of joint involvement was similar in the two cohorts with the knee being the most frequent site of infection.
Extended report

Table 1  Baseline patient characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>nbDMARD (n=3673)</th>
<th>All TNF (n=11 881)</th>
<th>p Value*</th>
<th>Etanercept (n=4139)</th>
<th>Infliximab (n=3475)</th>
<th>Adalimumab (n=4267)</th>
<th>p Value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (SD)</td>
<td>60 (12)</td>
<td>56 (12)</td>
<td>&lt;0.001</td>
<td>56 (12)</td>
<td>56 (12)</td>
<td>57 (12)</td>
<td>0.018</td>
</tr>
<tr>
<td>Sex (% female)</td>
<td>2652 (72)</td>
<td>9053 (76)</td>
<td>&lt;0.001</td>
<td>3193 (77)</td>
<td>2626 (76)</td>
<td>3234 (76)</td>
<td>0.203</td>
</tr>
<tr>
<td>DAS28 (mean (SD))</td>
<td>3.6 (1.3)</td>
<td>6.8 (1.0)</td>
<td>&lt;0.001</td>
<td>6.6 (1.0)</td>
<td>6.6 (1.0)</td>
<td>6.5 (1.0)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>HAQ score (mean (SD))</td>
<td>0.5 (0.6)</td>
<td>2.0 (0.6)</td>
<td>&lt;0.001</td>
<td>2.1 (0.6)</td>
<td>2.1 (0.5)</td>
<td>1.9 (0.6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Disease duration (years (median (IQR)))</td>
<td>6 (1, 15)</td>
<td>11 (6–19)</td>
<td>&lt;0.001</td>
<td>12 (6–19)</td>
<td>12 (6–19)</td>
<td>10 (5–18)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Baseline steroid use (n (%))</td>
<td>945 (23)</td>
<td>5228 (44)</td>
<td>&lt;0.001</td>
<td>1979 (48)</td>
<td>1609 (46)</td>
<td>1664 (39)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Diabetes (n (%))</td>
<td>234 (6.7)</td>
<td>675 (5.8)</td>
<td>0.033</td>
<td>255 (6)</td>
<td>169 (4)</td>
<td>261 (6)</td>
<td>0.026</td>
</tr>
<tr>
<td>COPD (n (%))</td>
<td>304 (8)</td>
<td>570 (5)</td>
<td>&lt;0.001</td>
<td>222 (5)</td>
<td>165 (5)</td>
<td>183 (4)</td>
<td>0.070</td>
</tr>
</tbody>
</table>

*Represents the significance of differences between the DMARD and anti-TNF cohorts using χ² tests for categorical outcomes and Wilcoxon rank sum tests for continuous variables.
†Represents the significance of differences between the three anti-TNF drugs using χ² tests for categorical outcomes and Kruskal–Wallis rank tests for continuous variables.

Table 2  Risk of septic arthritis in patients with rheumatoid arthritis

<table>
<thead>
<tr>
<th>Exposure time (years)</th>
<th>fbDMARD</th>
<th>All anti-TNF</th>
<th>Etanercept</th>
<th>Infliximab</th>
<th>Adalimumab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Events (n)</td>
<td>11 426</td>
<td>42 671</td>
<td>18 554</td>
<td>10 827</td>
<td>13 289</td>
</tr>
<tr>
<td>Incident rate/1000 pyrs</td>
<td>20</td>
<td>179</td>
<td>86</td>
<td>41</td>
<td>52</td>
</tr>
<tr>
<td>(95% CI)</td>
<td>1.8 (1.1 to 2.7)</td>
<td>4.2 (3.6 to 4.8)</td>
<td>4.6 (3.7 to 5.7)</td>
<td>3.8 (2.7 to 5.1)</td>
<td>3.9 (2.9 to 5.1)</td>
</tr>
<tr>
<td>Unadjusted HR (95% CI)</td>
<td>Ref</td>
<td>2.5 (1.6 to 4.0)</td>
<td>3.0 (1.8 to 4.8)</td>
<td>2.2 (1.3 to 3.8)</td>
<td>2.3 (1.4 to 3.8)</td>
</tr>
<tr>
<td>Adjusted HR (95% CI)</td>
<td>Ref</td>
<td>2.3 (1.2 to 4.4)</td>
<td>2.5 (1.3 to 4.9)</td>
<td>2.4 (1.0 to 5.8)</td>
<td>1.9 (0.9 to 4.0)</td>
</tr>
</tbody>
</table>

*Anti-TNF, anti-tumour necrosis factor; nbDMARD, non-biological disease-modifying antirheumatic drug; pyrs, patient years.

DISCUSSION

Key questions remain unanswered from clinical trial data regarding the safety of anti-TNF therapy, particularly with respect to rare outcomes including SA. This is the first study to specifically examine the rate of SA in patients treated with anti-TNF therapy. We confirmed our primary hypothesis that SA is increased in patients on anti-TNF therapy. However, these results must be interpreted with some important caveats. First, the absolute risk of SA remains very small, with an incidence rate of only 4.2 per 1000 pyrs of...
follow-up in the anti-TNF cohort. Second, given the observational nature of this study, it is important to recognise that differences exist between the nbDMARD and anti-TNF cohorts. Any difference in outcome between the two cohorts reflects both the effect of the drug and the inherent differences in patient populations. The biologically treated cohort had, as expected, more severe disease at baseline and this is an important risk factor for the development of SA. We addressed this issue through adjustment using a propensity model. The adjusted analysis continued to show a significant increase in the risk of SA in the anti-TNF cohort (HR 2.3), which reflects the risk attributable to the therapy itself. It is possible that there are unmeasured differences between the cohorts that we are unable to adjust for.

The point estimates showed the highest risk for SA in patients on ETN. However, this was not a hypothesis that we had set out to test. After adjusting for confounders, the 95% CIs for all three agents overlapped. Year of entry into the study is an important confounder when comparing between agents. We were not able to adjust for.

There were a number of potential explanations for this early increased risk. First, it may reflect a depletion of susceptible individuals from the exposed cohort. Second, it may reflect a true reduction in risk of joint infection in patients who achieve better control of their RA. However, the data presented here do not allow definite conclusions to be made about these possibilities.

We also addressed the influence of orthopaedic surgery on the risk of SA. The BSR recommends that patients on anti-TNF therapy stop their treatment temporarily for 2–4 weeks prior to major surgical procedures and do not recommence therapy until wound healing is satisfactory. Although it is not known with certainty how strictly physicians and patients adhere to these guidelines, in the context of this practice, it is very reassuring to see no evidence of an increased risk of prosthetic joint SA.

In summary, exposure to TNF inhibitor therapy is associated with an increased risk of SA in patients with RA. This risk seen in this study was greatest in the first year of treatment. Joint prosthesis is a key risk factor for infection. Careful vigilance for joint infections in anti-TNF patients remains important, especially in the early months of therapy, with awareness of the potential range of pathogens that may be responsible. Antibiotic guidelines should incorporate this information and consider giving specific advice for patients being actively treated with anti-TNF agents. Current evidence does not support any other anti-TNF agent having a safer profile with regard to SA.

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Funding The BSR commissioned the BSRBR as a UK-wide national project to investigate the safety of biological agents in routine medical practice. BSR receives unrestricted income from UK pharmaceutical companies, presently AbbVie Laboratories, Amgen, Schering Plough (now MSD) and Wyeth Pharmaceuticals (now Pfizer). This income finances a wholly separate contract between the BSR and the University of Manchester, who provide and run the BSR data collection, management and analysis services. The principal investigators and their team have full academic freedom and are able to work independently of pharmaceutical industry influence. All decisions concerning analyses, interpretation and publication are made autonomously of any industrial contribution.

Competing interests MH has received payment for advisory work for Shire, CSL and GSK, as well as receiving financial support for attending conferences from GSK, OctoPharma, Biorad, BPL, Phadia, GSK and CSL. All other authors have declared no conflicts of interest.

Ethics approval Ethics approval for this study was obtained in December 2000 from the Multicentre Research Ethics Committee (MREC) for the Northwest of England.

Provenance and peer review Not commissioned; externally peer reviewed.

Figure 2 Spline model showing changing risk of septic arthritis over time in the anti-TNF cohort. Anti-TNF, anti-tumour necrosis factor.

Table 3 Baseline patient characteristics

<table>
<thead>
<tr>
<th>Number of patients with prosthetic joints</th>
<th>nbDMARD (n=659)</th>
<th>Anti-TNF (n=2689)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exposure time (years)*</td>
<td>1954</td>
<td>12 959</td>
</tr>
<tr>
<td>Events</td>
<td>6</td>
<td>41</td>
</tr>
<tr>
<td>Incidence prosthetic joint/1000</td>
<td>2.1 (1.1 to 6.7)</td>
<td>3.2 (2.3 to 4.3)</td>
</tr>
<tr>
<td>Adjusted HR (95% CI)</td>
<td>Ref</td>
<td>1.2 (0.4 to 3.4)</td>
</tr>
</tbody>
</table>

*Patients were included in this analysis only if they had a prosthetic joint in situ. Anti-TNF, anti-tumour necrosis factor; nbDMARD, non-biological disease-modifying antirheumatic drug; pyrs, patient years.
REFERENCES
*BSRBR Control Centre Consortium*

The BSRBR Control Centre Consortium consists of the following institutions (all in the UK): Antrim Area Hospital, Antrim (Dr Nicola Maiden), Cannock Chase Hospital, Cannock Chase (Dr Tom Price), Christchurch Hospital, Christchurch (Dr Neil Hopkinson), Royal Derby Hospital, Derby (Dr Sheila O’Reilly), Dewsbury and District Hospital, Dewsbury (Dr Lesley Hordon), Freeman Hospital, Newcastle-upon-Tyne (Dr Ian Griffiths), Gartnavel General Hospital, Glasgow (Dr Duncan Porter), Glasgow Royal Infirmary, Glasgow (Prof Hilary Capell), Haywood Hospital, Stoke-on-Trent (Dr Andy Hassell), Hope Hospital, Salford (Dr Romela Benitha), King's College Hospital, London (Dr Ernest Choy), Kings Mill Centre, Sutton-In Ashfield (Dr David Walsh), Leeds General Infirmary, Leeds (Prof Paul Emery), Macclesfield District General Hospital, Macclesfield (Dr Susan Knight), Manchester Royal Infirmary, Manchester (Prof Ian Bruce), Musgrave Park Hospital, Belfast (Dr Allister Taggart), Norfolk and Norwich University Hospital, Norwich (Prof David Scott), Poole General Hospital, Poole (Dr Paul Thompson), Queen Alexandra Hospital, Portsmouth (Dr Fiona McCrae), Royal Glamorgan Hospital, Glamorgan (Dr Rhian Goodfellow), Russells Hall Hospital, Dudley (Prof George Kitas), Selly Oak Hospital, Selly Oak (Dr Ronald Jubb), St Helens Hospital, St Helens (Dr Rikki Abernethy), Weston General Hospital, Weston-super-Mare (Dr Shane Clarke/Dr Sandra Green), Wythenshawe Hospital, Manchester (Dr Paul Sanders), Withybush General Hospital, Haverfordwest (Dr Amanda Coulson), North Manchester General Hospital (Dr Bev Harrison), Royal Lancaster Infirmary (Dr Marwan Bukhari) and The Royal Oldham Hospital (Dr Peter Klimiuk).
Methodology supplementary information

Modelling of the hazard function over time was done using a flexible parametric spline model with 2 degrees of freedom (equivalent to 1 knot). Model selection was chosen by minimising the Akaike Information Criterion. The use of larger numbers of degrees of freedom did not improve the model. This method of spline modelling has been described in detail previously (Royston P, Stata journal 2001: 1; 1, 1-28).

Multiple imputations were performed in Stata using the ICE command. Missing data were present in the following variables: age, disease duration, baseline HAQ, and baseline DAS28 score. The imputation model was constructed separately for the DMARD and anti-TNF cohorts. Age, gender, disease duration, baseline HAQ, baseline DAS28 score, co-morbidity, smoking status, entry year, baseline steroid exposure and prior orthopaedic surgery were all included as predictors within the imputation model. Twenty imputation cycles were performed and the resulting data were analysed using Rubin’s rules with the MIM command.

Adjusting for confounders was performed using an inverse probability of treatment weighting score. A probability of treatment (propensity) score was generated using logistic regression. As several covariates were associated with treatment likelihood in a non-linear pattern, or demonstrated an interaction with other covariates, sequential analyses were performed to identify the components of the model. If a non-linear relationship was identified, orthogonal polynomial transformations of the covariates were added to the model until a suitable fit was obtained. The final propensity score included polynomials for age, DAS28 score and HAQ score; interactions were identified between age and DAS28 score, age and prior orthopaedic surgery, entry year and DAS28 score, co-morbidity and DAS28 score, steroid exposure and disease duration, entry year and disease duration, and entry year and steroid exposure. The inverse of the probability (or 1 minus the inverse of the probability in the DMARD cohort) was then used as the treatment weight in the analysis. Weights greater than 20 that would destabilise the model were replaced with the value of 20.