EXTENDED REPORT

The comparative effectiveness of abatacept versus anti-tumour necrosis factor switching for rheumatoid arthritis patients previously treated with an anti-tumour necrosis factor

Leslie R Harrold, George W Reed, Joel M Kremer, Jeffrey R Curtis, Daniel H Solomon, Marc C Hochberg, Jeffrey D Greenberg

ABSTRACT

Objective We compared the effectiveness of abatacept (ABA) versus a subsequent anti-tumour necrosis factor inhibitor (anti-TNF) in rheumatoid arthritis (RA) patients with prior anti-TNF use.

Methods We identified RA patients from a large observational US cohort (2/1/2000–8/7/2011) who had discontinued at least one anti-TNF and initiated either ABA or a subsequent anti-TNF. Using propensity score (PS) matching (1:1 match), effectiveness was measured at 6 and 12 months after initiation based on mean change in Clinical Disease Activity Index (CDAI), modified American College of Rheumatology (mACR) 20, 50 and 70 responses, modified Health Assessment Questionnaire (mHAQ) and CDAI remission in adjusted regression models.

Results The PS-matched groups included 431 ABA and 746 anti-TNF users at 6 months and 311 ABA and 493 anti-TNF users at 12 months. In adjusted analyses comparing response following treatment with ABA and anti-TNF, the difference in weighted mean change in CDAI (range 6–8) at 6 months (0.46, 95% CI −0.82 to 1.73) and 12 months was similar (−1.64, 95% CI −3.47 to 0.19). The mACR20 responses were similar at 6 (28–32%, p=0.73) and 12 months (35–37%, p=0.48) as were the mACR50 and mACR70 (12 months: 20–22%, p=0.25 and 10–12%, p=0.49, respectively). Meaningful change in mHAQ was similar at 6 and 12 months (30–33%, p=0.41 and 29–30%, p=0.39, respectively) as was CDAI remission rates (9–10%, p=0.42 and 12–13%, p=0.91, respectively).

Conclusions RA patients with prior anti-TNF exposures had similar outcomes if they switched to a new anti-TNF as compared with initiation of ABA.

INTRODUCTION

Rheumatoid arthritis (RA) affects an estimated 1.3 million Americans and is associated with substantial morbidity and mortality.1 To prevent the development of permanent joint damage and deformity as well as functional impairment, the current treatment paradigm is to treat patients with active disease with disease modifying antirheumatic drugs (DMARDs).2 The typical approach is to introduce a conventional non-biologic DMARD first, followed by combination non-biologic DMARDs or biologics in non-responders.3 The initial biologic used in the vast majority of patients is the anti-tumour necrosis factor (anti-TNF). These agents were the first class of biologics approved by the Food and Drug Administration (FDA) for the treatment of RA.

However, little is known regarding what to do when patients have an inadequate response to the first anti-TNF agent. Changing mechanism of action has been shown to be beneficial in comparisons of rituximab versus a subsequent anti-TNF as demonstrated by improvements in disease activity as well as persistence of therapy. But it is unclear whether any change in mechanism of action will result in improvement or if the results seen were related specifically to B cell depletion. Further exploration is needed to assess whether switching to abatacept (ABA), as a different example of changing mechanism of action, also provides a benefit among those with inadequate response to an anti-TNF. To date, there have been no randomised controlled trials (RCTs) comparing the effectiveness of switching to ABA as compared with a subsequent anti-TNF, and a recent meta-analysis using indirect methods suggested that use of an anti-TNF was associated with a higher probably of achieving an American College of Rheumatology (ACR) 50 response than ABA, although not all indirect comparisons have shown a difference between ABA and anti-TNF agents.4–10 In biologic-naïve patients, a few studies have compared responses to ABA versus an anti-TNF.11–14 A similar efficacy was found when comparing responses to ABA and infliximab, although there have been concerns that the results may not be generalisable to the US population of RA patients due to the lower doses of infliximab used (3 mg/kg every 8 weeks) and lack of the usual infliximab loading dose. Additionally, ABA (in combination with methotrexate (MTX)) has been compared with adalimumab with MTX among biologic-naïve RA patients with similar ACR response rates.14

Given the absence of head-to-head RCTs addressing best treatment approaches in patients with inadequate response to an anti-TNF agent, comparative effectiveness studies using observational data from registries can be employed.15 Therefore, the aim of the present study was to compare the clinical effectiveness of ABA versus a subsequent anti-TNF agent among RA patients with previous anti-TNF exposure in a large US cohort of RA patients using the
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Correspondence of Rheumatology Researchers of North America (CORRONA) registry. Specifically, we sought to compare change in disease activity, composite response rates, meaningful improvement in function and remission outcomes over a 1-year period.

**METHODS**

**Data sources and data collection**

CORRONA is a US observational cohort of patients with arthritides who are enrolled by participating rheumatologists in 132 practices both academic and private practice with prospective data collection; the details have been previously described.16 17 Data are collected from both patients and their treating rheumatologists, who gather information on disease duration, prognosis, disease severity and activity, medical comorbidities, use of medications including DMARDs and adverse events.18 Laboratory data such as inflammatory markers are not available at all visits as they are not mandated due to the observational nature of the registry and some rheumatologists do not obtain this information routinely. Follow-up assessments are requested at least as often as every 4 months and completed during routine clinical encounters. Approvals for participation in the CORRONA registry are obtained from respective Institutional Review Boards (IRBs) of participating academic sites and a central IRB for private practice sites, and patients provide informed consent before enrolment.

**Study population**

There were 27 412 patients with a diagnosis of RA in the registry. We identified patients with exposure to one or more anti-TNF agents but no prior use of non-anti-TNF biologics using CORRONA data between 2/1/2000 and 8/1/2011. For the main analyses, we included all patients regardless of reason for the switch to ABA or a subsequent anti-TNF following prior exposure to an anti-TNF. However, in the sensitivity analyses, we included those patients who were inadequate responders to the prior anti-TNF. Inclusion criteria into the study cohort included receipt of either a subsequent anti-TNF or ABA, a follow-up appointment at 6 and/or 12 months following initiation and the presence of all the elements necessary to calculate response based on the Clinical Disease Activity Index (CDAI), a well-recognised instrument to measure disease activity, and ACR20, 50 and 70 responses.19–21 For the purpose of this study, RA patients in remission at baseline, meaning at the time the new agent was initiated, defined by CDAI ≤2.8, were excluded from the study population (switching while in remission is seen when cost or tolerability influences medication use).

**Measures and data collection**

Data were collected during the study period from physician assessment and patient questionnaires completed during clinical encounters. Non-biologic and biologic DMARDs, including anti-TNF agents, are recorded at the time of the clinical encounter. Among the data elements collected in the registry relevant to this study are a 28 tender and swollen joint count, physician’s and patient’s global assessments of disease activity, patient’s assessment of pain, the modified Health Assessment Questionnaire (mHAQ) assessing physical function and erythrocyte sedimentation rate (ESR). When available, acute phase reactant data are recorded from laboratory tests obtained within 10 days of the clinical encounter.

**Study outcomes**

Responsiveness to medication treatment was defined based on improvement in CDAI as our primary outcome (eg, baseline CDAI—follow-up CDAI). As a secondary outcome, we evaluated modified ACR20 (mACR20), modified ACR50 (mACR50) and modified ACR70 (mACR70) responses, which do not include acute phase reactants. These measures have been previously defined and validated and shown to highly correlate with ACR responses that include acute phase reactants.20–22 An mACR20 response required ≥20% improvement in tender joint count and swollen joint count, as well as two or more of the four remaining ACR response components, including physician’s global assessment, patient’s global assessment, patient’s global pain and mHAQ. The mACR50 and mACR70 responses were calculated using the same criteria requiring 50% and 70% improvement, respectively. In addition, we evaluated achievement of a clinically meaningful change in the mHAQ (≥0.25).23 Disease remission outcomes were defined using CDAI.

As sensitivity analyses, we examined improvement in the modified Disease Activity Score 28 (mDAS28), which is calculated without results of acute phase reactants as well as achievement of remission using the mDAS28 cut-off of <2.6 (eg, baseline mDAS—follow-up mDAS).24 25 Specifically, the predictors for the ESR include tender and swollen 28 joint counts, patient’s and physician’s global assessments, patient’s pain assessment and physical function (mHAQ). The calculation of the mDAS28 is shown as follows:

\[
\text{mDAS28} = 0.53 \times (28TJC) + 0.31 \times (28SJC) + 0.25 \times \text{mHAQ} + 0.001 \times \text{PAIN} + 0.005 \times \text{EGA} + 0.014 \times \text{PGA} + 1.694
\]

It has been previously shown that the mDAS28 performs as well as the DAS28 as a measure of response and disease activity.23

**Potential confounders**

Given concerns regarding treatment-selection bias, we used propensity score (PS) matching. A PS is the predictive probability of an outcome; in this case receiving ABA versus another anti-TNF. PS matching has been used in numerous studies to reduce selection bias in observational studies.26 In this study, the PS was estimated using a multivariable logistic regression model predicting the use of ABA versus an anti-TNF using the following key variables: age, number of prior anti-TNF agents (one prior vs two or more priors), duration of RA, swollen joint count, patient’s global assessment, physician’s global assessment and insurance type. We then matched on the PS (up to 1:4 match using a caliper of 0.01) using a greedy matching routine without replacement.27 We matched separately patients who had a follow-up at 6 months as well as those with a follow-up at 12 months. Patients could have contributed data to both groups. In addition, matching was done based on the number of available patients. Thus, for some analyses, up to four anti-TNF users were matched to one ABA user while for other analyses it was four ABA users to one anti-TNF users. Initially we performed the analyses based on prior anti-TNF use, specifically comparing PS-matched ABA and anti-TNF users exposed to only one prior anti-TNF separately from PS-matched ABA and anti-TNF users exposed to two or more anti-TNFs. However, the results were not different; so we pooled the groups together.

**Statistical analysis**

Patient clinical and demographic characteristics were compared within the two matched cohorts (6 and 12 months). For continuous measures, means and SDs were estimated and t tests were used to test statistical differences among the groups. For dichotomous measures, percentages were estimated and Fisher’s
exact test was used to test statistical differences among the groups. Response outcomes were defined based on the change in CDAI in mDAS28 as well as mACR20, mACR50 and mACR70 responses and meaningful change in mHAQ as defined above. Remission outcomes were based on CDAI and mDAS28. For patients who discontinued ABA or the subsequent anti-TNF agent, the last observation prior to the discontinuation was carried forward.

Comparisons of response rates were made at 6 and 12 months postinitiation using 3-month time windows for capturing study visits. For patients with two or more visits within the window, we used the visits closest to the date of interest (6 or 12 months). Weighted mean improvement and response rates for both time points were estimated. Unadjusted and adjusted estimates comparing differences (for CDAI and mDAS28), response rates (mACR20, mACR50 and mACR70), meaningful improvement in function (mHAQ) and remission rates (CDAI and mDAS28) were performed separately using generalized linear latent and mixed models (GLLAMM). We used GLLAMM because we needed two levels of weights due to the n:1 matching and hierarchical nature of the data (eg, patients nested by physician) as these factors have been shown to influence the evaluations of treatment effectiveness. Specifically, we included multilevel weights and clustering on physician. The first level is for the patient with individual weights based on the PS matching ratio, and the second level is for the physician. We computed the second-level weights by averaging the individual probability weights within each physician. These results were reported with estimated 95% CIs. Residual differences in baseline characteristics after matching were included in the outcome models. Patients with missing data for any of the variables in the final models were excluded from the models. Additionally, we explored potential interactions between medication selection (ABA vs anti-TNF) and number of prior anti-TNF agents in the models.

Additional sensitivity analyses were carried out, which included re-running the analyses with 1:1 matching. To assess whether responses differed based on previous experience with an anti-TNF, we re-ran the analyses (both 1:n and 1:1 matching approaches) restricting it to patients who had an inadequate clinical response to the prior anti-TNF; specifically we excluded patients who switched for safety, tolerability or other reasons from this sensitivity analysis. All analyses were conducted using STATA V10 and 11.

RESULTS

There were 3076 RA patients who met inclusion and exclusion criteria including prior exposure to one or more anti-TNFs, initiation of either of ABA (n=968) or a subsequent anti-TNF (n=2108) without prior use of a non-TNF biologic (figure 1). Among those with a follow-up appointment at 6 months following initiation, there were 440 ABA users and 958 anti-TNF users. Propensity matching was performed for those with a 6-month follow-up appointment, resulting in 431 ABA and 746 anti-TNF users of which 66% had been exposed to only one prior anti-TNF (table 1). After matching there was better balance across most baseline patient characteristics; however, there was still imbalance across several covariates, including medicare insurance, rheumatoid factor seropositivity, disease duration, mHAQ and concomitant prednisone use, which were added as individual covariates to the outcome models. For follow-up at 12 months, there were 327 ABA users and 637 anti-TNF users. Those with follow-up at 12 months were PS matched, resulting in 311 ABA users and 493 anti-TNF users of whom 68% had exposure to only one prior anti-TNF and there was good balance across most baseline characteristics except for disease duration that was added to as an individual covariate to the outcome models (see online supplementary table for the cohort with follow-up at 12 months). There were 272 ABA users and 531 anti-TNF users who contributed to both the 6-month and 12-month cohorts.
Among the ABA users, 21.5% discontinued the medication by 6 months, and by 12 months it was 27.6%. In anti-TNF users, 36.6% discontinued the medication by 6 months and 38.2% by 12 months. Among those who discontinued, toxicity was reported by 5.6% of ABA users by 6 months and 3.0% by 12 months. For anti-TNF users, it was 13.0% by 6 months and 9.0% by 12 months.

At 6 months, response rates based on mean change in CDAI were similar in ABA users as compared with anti-TNF users (–6.1 vs –5.6) in both unadjusted and adjusted analyses (figure 2). Achievement of a mACR20 occurred in 31.7% of ABA users and 28.2% of anti-TNF users, which was not significant in unadjusted or adjusted analyses. Achievement of a mACR30 was more likely in ABA users as compared with anti-TNF users (19.7% vs 14.5%; adjusted OR of 1.40; 95% CI 1.05 to 1.85). The mACR70 response rates were similar, occurring in 8.6% of ABA users and 6.8% of anti-TNF users, which were not significantly different in both unadjusted and adjusted analyses. Achievement of a clinically meaningful improvement in mHAQ was similar, occurring in 33.1% of ABA users and 30.2% of anti-TNF users. At 12 months, response rates based on change in CDAI seemed to favour anti-TNF users with borderline significance (difference in mean change of CDAI of –1.64; 95% CI –3.47 to 0.19). Between 35% and 37% of both groups achieved a mACR20, 20–22% a mACR50 and 10–12% a mACR70; none of these differences were significant in unadjusted or adjusted analyses. Achievement of a meaningful clinical improvement in function occurred similarly in the two groups (28.9% of ABA users and 29.8% of anti-TNF users).

Remission rates based on CDAI were similar at 6 months, occurring in 9% of ABA users and 10% of anti-TNF users (figure 3). At 12 months, remission rates were 12–14% among both groups based on CDAI.

Results were similar when using the change in mDAS28 (0.81 vs 0.68; p=0.13) at 6 and 12 months (1.03 vs 1.03; p=0.93). Achievement of mDAS28 remission occurred more often in ABA users (20.2% vs 17.3%) at 6 months, which was significant in adjusted analyses (OR 1.55; 95% CI 1.01 to 2.36). To further explore whether outcomes in the two groups differed based on response to the prior anti-TNF, we re-ran the analyses in those patients who discontinued their prior anti-TNF due to lack of efficacy and similar results were found (see online supplementary figures). Additionally, we explored the robustness of our results using two strategies to ensure our matching protocol did not influence the findings. First, we reid the analyses using 1:1 matching as well as evaluating the 6-month and 12-month outcomes in a single cohort of patients who were seen both at 6 and 12 months. Again, the results were similar to those found with the 1:n matched cohort presented here.

**DISCUSSION**

Using data from CORRONA, a large US-based RA registry, we compared the clinical effectiveness of ABA versus a subsequent anti-TNF agent among RA patients with previous anti-TNF exposure. Specifically, we compared change in disease activity and remission rates based on CDAI and mDAS28 as well as ACR response rates and meaningful improvement in function. Both treatments were effective for the primary outcome of

### Table 1 Baseline characteristics of unmatched and matched cohorts using abatacept or an anti-TNF with follow-up at 6 months

| Demographics | ABA (N=440) | Anti-TNF (N=958) | p Value | ABA (N=431) | Anti-TNF (N=746) | p Value |
|--------------|-------------|-----------------|---------|-------------|-----------------|---------|
| Age (mean age, SD) | 57.7 (12.4) | 55.6 (12.5) | 0.003 | 57.6 (12.4) | 57.2 (11.7) | 0.578 |
| Gender (% female) | 82.7 | 80.0 | 0.242 | 82.4 | 79.1 | 0.196 |
| Race (% white) | 81.8 | 85.4 | 0.097 | 82.4 | 85.1 | 0.216 |
| Insurance (%) | | | | | | |
| Private | 78.6 | 79.3 | 0.830 | 79.4 | 81.0 | 0.542 |
| Medicare | 36.8 | 29.8 | 0.012 | 36.4 | 30.8 | 0.053 |
| Medicaid | 5.0 | 6.9 | 0.188 | 5.1 | 6.4 | 0.374 |
| Clinical | | | | | | |
| Rheumatoid factor seropositivity (%) | 71.3 | 78.0 | 0.032 | 70.7 | 80.0 | 0.006 |
| Disease duration (mean years, SD) | 13.4 (10.3) | 11.5 (9.4) | 0.000 | 13.3 (10.0) | 12.1 (9.8) | 0.045 |
| Tender joints, N (SD) | 7.4 (7.1) | 7.3 (7.1) | 0.717 | 7.4 (7.1) | 7.3 (7.1) | 0.811 |
| Swollen joints, N (SD) | 6.8 (6.0) | 5.9 (5.3) | 0.005 | 6.7 (5.9) | 6.1 (5.2) | 0.087 |
| Patient pain score (mean, SD) | 52.7 (25.3) | 49.3 (25.8) | 0.023 | 52.7 (25.5) | 50.2 (25.8) | 0.112 |
| mHAQ (mean, SD) | 0.7 (0.5) | 0.6 (0.5) | 0.012 | 0.7 (0.5) | 0.6 (0.5) | 0.047 |
| CDAI (mean, SD) | 22.9 (14.1) | 21.4 (13.2) | 0.057 | 22.8 (14.1) | 21.7 (12.9) | 0.182 |
| Disease activity* (%) | | | | | | |
| Low | 19.8 | 20.6 | 0.210 | 20.2 | 19.0 | 0.354 |
| Moderate | 35.7 | 39.8 | 35.7 | 40.0 |
| High | 44.6 | 39.7 | 44.1 | 41.0 |
| Concomitant medications | | | | | | |
| Corticosteroids, (%) | 40.0 | 34.0 | 0.036 | 39.4 | 33.0 | 0.027 |
| Leflunomide, (%) | 12.5 | 12.7 | 0.931 | 12.5 | 13.3 | 0.787 |
| MTX, (%) | 54.8 | 57.1 | 0.417 | 55.2 | 55.5 | 0.951 |
| Dose, mg/week (SD) | 16.3 (6.2) | 16.4 (6.0) | 0.843 | 16.3 (6.2) | 16.2 (5.9) | 0.906 |

ABA, abatacept; anti-TNF, anti-tumour necrosis factor; CDAI, Clinical Disease Activity Index; mHAQ, modified Health Assessment Questionnaire; MTX, methotrexate. *Disease activity based on the CDAI.
interest as the minimally clinically important difference for change in CDAI is 4.3. We examined effectiveness based on several different outcome measures (change in disease activity, mACR response, meaningful improvement in mHAQ and remission) and timeframe. For the vast majority of the analyses, effectiveness was similar in the two treatment arms. This is further supported by our sensitivity analysis in which we limited the sample to those who discontinued their prior anti-TNF due to lack of efficacy and found similar results.

While our results stem from an observational cohort, they are similar to two recent systematic reviews based on clinical trials that compared the efficacy of ABA with other biologic agents using Bayesian network meta-analysis methods in inadequate responders to MTX. In the first review, ABA was compared with TNF and non-TNF biologics and found to have similar ACR50 and remission rates at 24 and 52 weeks. In the second review, ABA was compared with anti-TNF agents with similar responses to therapy found based on change in HAQ and ACR response rates at 6 months. A recently published systematic review of randomised clinical trials in patients with inadequate response to nbDMARDs found that use of ABA as compared with anti-TNFs was associated with equivalent ACR20 and ACR50 responses at 6 and 12 months, but there was a higher likelihood of achieving an ACR70 response and DAS28 remission at 12 months with ABA. In contrast, a meta-analysis of randomised clinical trials of patients with inadequate response to MTX suggested a higher probability of achieving an ACR50 response with anti-TNFs as compared with ABA. These differences may in part be related to different time periods for the selection of relevant studies and analysis techniques. In a recent head-to-head comparison of the addition of ABA versus adalimumab in biologic-naïve patients on background MTX, no meaningful differences in efficacy were found. Additionally, ABA has been compared with anti-TNF agents in the MTX-naïve early RA population with no differences in time to response and response achieved over a 1-year time period. Our findings complement these studies as according to our knowledge it is the first study that examines comparative effectiveness in those with prior exposure to anti-TNF.

Currently, it is not possible to predict which patients will respond to which particular biologic agent. However, this study suggests that just changing the mechanism of action may not be enough to improve disease activity. When discussing next therapeutic interventions in those not responding or unable to take their current anti-TNF agent, these results can potentially contribute to the discussion as patient weigh other factors such potential adverse events associated with the different agents.

Figure 2 Adjusted* response rates at 6 and 12 months among abatacept and anti-tumour necrosis factor (anti-TNF) users. *Adjusted for number of prior anti-TNF medications, baseline disease activity, rheumatoid arthritis disease severity and concomitant medications. Adjusted mean difference in Clinical Disease Activity Index (CDAI) at 6 months was 0.46 (95% CI –0.82 to 1.73) and at 12 months –1.64 (95% CI –3.47 to 0.19). Adjusted OR for modified American College of Rheumatology (mACR20 response at 6 months was 1.05 (95% CI 0.81 to 1.36) and at 12 months 0.87 (95% CI 0.59 to 1.29). Adjusted OR for mACR50 response at 6 months was 1.40 (95% CI 1.05 to 1.85) and at 12 months 0.86 (95% CI 0.58 to 1.27). Adjusted OR for mACR70 response at 6 months was 1.72 (95% CI 0.84 to 3.53) and at 12 months 1.12 (95% CI 0.56 to 2.24). Adjusted OR for meaningful change in modified Health Assessment Questionnaire (mHAQ) at 6 months was 1.15 (95% CI 0.86 to 1.54) and at 12 months 0.74 (95% CI 0.48 to 1.15).
out-of-pocket costs and medication route of administration that impacts convenience. The current availability of ABA as an intravenous infusion and subcutaneous injection does allow more flexibility for the patient’s preference.

Our study has several strengths. We utilised a national US-based registry of RA patients with physician-derived outcome measures to examine comparative effectiveness in patients with prior anti-TNF exposure. We examined drug response (based on change in disease activity, achievement of a modified ACR response and achievement of a meaningful improvement in mHAQ) and remission. In addition, we used both CDI and mDAS28 to assess response and remission. This work focused on the real-world effectiveness of agents in US patients, who are markedly different than RCT subjects in terms of comorbidities and RA disease activity, and more representative of patients found in clinical practice. Finally, we were able to use advanced epidemiologic methods (eg, PSs) to compare responses to the different agents when selection bias exists.

However, there are limitations to our study. The timing of the study visits was based upon clinic visits and not mandated as occurs in RCTs that influenced which patients were included based on follow-up visits. Of note, the mean study interval between study visits was approximately 4.5 months. Most of the patients were white, which limits generalisability to other racial groups. In addition, acute phase reactant data were not available for all patients in the study. As a result, we applied previously validated outcome measures not requiring acute phase reactants such as CDAI and mDAS for response and remission outcomes as well as modified ACR response rates and mHAQ scores. While PS matching improved the balance between the cohorts to most measure covariates, it is possible that there is still residual confounding for unmeasured covariates.

In summary, the results of this US-based observational study indicate similar rates of drug response and remission with use of ABA and anti-TNF agents among those with prior anti-TNF exposure. These findings were unchanged when we limited the analyses to those who failed their prior anti-TNF due to lack of efficacy. While we saw no difference when comparing ABA with anti-TNF agents, other non-TNF biologics with different mechanisms of action may be associated with superior outcomes as compared with anti-TNFs. Further comparative effectiveness studies are necessary to evaluate the benefits of switching to a different non-TNF biologic versus anti-TNF intraclax switching among patients with an inadequate response to an anti-TNF.

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REFERENCES

1. Helmick CG, Felson DT, Lawrence RC, et al. Estimates of the prevalence of arthritis and other rheumatic conditions in the United States. Part I. Arthritis Rheum 2008;58:15–25.
2. Smolen JS, Aletaha D, Machold KP. Therapeutic strategies in early rheumatoid arthritis. Best Pract Res Clin Rheumatol 2005;19:163–77.
3. van Vollenhoven RF, Ernestam G, Gebekop P, et al. Addition of infliximab compared with addition of sulfasalazine and hydroxychloroquine to methotrexate in patients with early rheumatoid arthritis (Swefot trial): 1-year results of a randomised trial. Lancet 2009;374:549–56.
4. Finch A, Ciurea A, Bruhlart L, et al. B cell depletion may be more effective than switching to an anti-tumour necrosis factor agent in rheumatoid arthritis patients with inadequate response to anti-tumour necrosis factor agents. Arthritis Rheum 2007;56:1417–23.
5. Gomez-Reino JJ, Maneiro JR, Ruiz J, et al. Comparative effectiveness of switching to alternative tumour necrosis factor (TNF) antagonists versus switching to rituximab in patients with rheumatoid arthritis who failed previous TNF antagonists: the MIRAR Study. Ann Rheum Dis 2012;71:1861–4.
6. Kekow J, Mueller-Ladner U, Schube-Koops H. Rituximab is more effective than second anti-TNF therapy in rheumatoid arthritis patients and previous TNAfalfa blocker failure. Biologics: Targets Ther 2012;6:191–9.
7. Blom M, Kleiv W, Donders AR, et al. Effectiveness of a third tumour necrosis factor-alpha-blocking agent compared with rituximab after failure of 2 TNF-blocking agents in rheumatoid arthritis. J Rheumatol 2011;38:2355–61.
8. Salliet C, Finck A, Katchamart W, et al. Indirect comparisons of the efficacy of biological antirheumatic agents in rheumatoid arthritis in patients with an inadequate response to conventional disease-modifying antirheumatic drugs or to an anti-tumour necrosis factor agent: a meta-analysis. Ann Rheum Dis 2011;70:266–71.
9. Guyot P, Taylor P, Christensen R, et al. Abatacept with methotrexate versus other biological agents in treatment of patients with active rheumatoid arthritis despite methotrexate: a network meta-analysis. Arthritis Res Ther 2011;13:R204.
10. Guyot P, Taylor P, Christensen R, et al. Indirect treatment comparison of abatacept with methotrexate versus other biologic agents for active rheumatoid arthritis despite methotrexate therapy in the United kingdom. J Rheumatol 2012;39:1198–206.
