Short-term risk of major adverse cardiovascular events or congestive heart failure in patients with psoriatic arthritis or psoriasis initiating biological therapy: a meta-analysis of randomised controlled trials

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ABSTRACT

Objective The objective was to investigate the short-term risk of major adverse cardiovascular events (MACEs) or congestive heart failure (CHF) in patients with psoriatic arthritis (PsA) or psoriasis initiating a biological therapy.

Methods Screening for the study was carried out using MEDLINE, Cochrane and Embase, from the inception of the database to December 2017. Randomised controlled trials (RCTs) of anti-tumour necrosis factor (TNF), anti-interleukin (IL)12/23, anti-IL23 and anti-IL17 agents for the treatment of PsA or psoriasis were included. Two investigators independently extracted MACEs or CHF data reported during the placebo-controlled phase. The primary outcome measures were the incidence of MACEs or CHF.

Results Of 753 references screened, 62 articles were selected, and 12 articles were added by manual searches. Accordingly 77 RCTs were included in the meta-analysis (MA) (10 174 patient-years (P-Y)). No significant difference was observed in MACE incidences in patients receiving anti-TNF, anti-IL12/23, anti-IL23 or anti-IL17 agents in comparison to the placebo. However, 10 MACEs were observed in the anti-IL12/23 group (1150 P-Y) compared with 1 in the placebo group (652 P-Y), with 0.01–0.00 to 0.02 event/P-Y risk difference, which is not statistically significant. This trend was not observed in the anti-IL23 group. No significant difference was observed in CHF incidence in patients receiving biological agents in comparison to placebo.

Conclusion This MA of 77 RCTs did not reveal any significant change in the short-term risk of MACE or CHF in patients with PsA or psoriasis initiating a biological therapy.

INTRODUCTION

Both psoriatic arthritis (PsA) and psoriasis have been associated with an increased prevalence of systemic and vascular inflammation and clinical atherosclerosis. A recent meta-analysis (MA) of observational studies showed a 43% increased risk of cardiovascular (CV) diseases in patients with PsA, while morbidity risks for myocardial infarction, cerebrovascular diseases and heart failure were increased by 68%, 22% and 31%, respectively, compared with the general population.
Notwithstanding, a MA of observational studies revealed that morbidity risks for stroke and myocardial infarction were increased by 10% and 20%, respectively in patients with mild psoriasis, while the risks of stroke, myocardial infarction and CV death were increased by 38%, 70% and 37%, respectively in patients with severe psoriasis, compared with the general population. 

It is a matter of debate whether the increased risk of CV morbidity and mortality observed in patients with PsA or psoriasis represents a causal association or a predisposition due to the underlying standard CV risk factors exhibited by these patients, but one hypothesis is that the inflammatory cascade activated in patients with severe PsA or psoriasis may contribute to the development of atherosclerosis. 

On the one hand, various experimental studies demonstrated that inhibition of proinflammatory cytokines, such as tumour necrosis factor (TNF)-α, interleukin (IL)-1 and IL-6, had beneficial effects on cardiac function and outcome. 

On the other hand, several experimental studies showed that IL-12 family cytokines, including IL-12, IL-23, IL-27 and IL-35, were involved in the crosstalk between major immune cell types that drive the proinflammatory and anti-inflammatory responses in atherosclerosis. Such pleiotropic role in atherosclerosis was also reported for IL-17, with some experimental studies suggesting a proatherogenic effect, while the others proposed an atheroprotective role.

Several biological therapies are currently approved for moderate-to-severe PsA or psoriasis, when a conventional systemic therapy fails to achieve disease control or when a patient is unable to tolerate the conventional systemic therapy because of adverse effects. These biological therapies include anti-TNF agents, approved for PsA and psoriasis (adalimumab, etanercept and infliximab) or only for PsA (certolizumab and golimumab); anti-IL12/23 agents, approved for PsA and psoriasis (ustekinumab) or previously assessed in psoriasis (biakinumab); anti-IL23 agents approved for psoriasis (guselkumab) with promising results in PsA (guselkumab and risankizumab) and psoriasis (tildrakizumab) and anti-IL17A agents, approved for PsA and psoriasis (ixekizumab and secukinumab) or only for psoriasis, with promising results in PsA (broladumab).

These biological therapies, which have demonstrated anti-inflammatory effects in inflammatory rheumatic and skin diseases, could theoretically prevent atherosclerosis, and therefore decrease the long-term risk of CV diseases. However, the short-term CV safety profile of these medicines is still a matter of debate, both in PsA and psoriasis. This is especially the case since clinicians are cognizant of the withdrawal of the application for approval of biakinumab in 2011, because of major adverse cardiovascular events (MACEs) reported during phase II and phase III clinical trials in subjects with psoriasis.

The objective of this systematic review and MA of randomised controlled trials (RCTs) was to investigate the short-term risk of MACEs or congestive heart failure (CHF) in patients with PsA or psoriasis initiating a biological therapy.

### MATERIALS AND METHODS

This MA was performed on RCTs from MEDLINE (PubMed), Cochrane and EMBASE data bases. We restricted the searches to the English language. All RCTs on anti-tumor necrosis factor (anti-TNF), anti-interleukin IL12/23, anti-IL23 and anti-IL17 agents used in the treatment of PsA, chronic plaque and pustuloplastic psoriasis were reviewed. Manual searches in American college of Rheumatology (ACR), European League Against Rheumatism (EULAR), American Academy of Dermatology (AAD) and Psoriasis Gene to Clinic (PGC) abstract archives were included in the review.

The searches were conducted using the following key words: (“Psoriasis” OR “Arthritis, Psoriatic”) AND (“Adalimumab” OR “Certolizumab” OR “Etanercept” OR “Golimumab” OR “Infliximab” OR “Biakinumab” OR “Ustekinumab” OR “Guselkumab” OR “Risankizumab” OR “Tildrakizumab” OR “Broladumab” OR “Ixekizumab” OR “Secukinumab”).

On the EMBASE and Cochrane data bases, the searches were restricted to “trials”. On PubMed, we restricted the research to “clinical trials”. On Cochrane data base, the selection of studies was restricted to the same key words, using (“Stroke” OR “Cardiovascular diseases”) in addition.

We included studies from the inception of the data base to December 2017. All the randomised, placebo-controlled, double-blind studies, that assess the efficacy and safety of anti-TNF, anti-IL12/23, anti-IL23 or anti-IL17 agents in adult patients with PsA and/or psoriasis were included if safety data concerning MACEs (defined as myocardial infarction, stroke or CV death) or CHF (defined as global cardiac failure with signs of right and left cardiac decompensation) were available.

Two authors (BC and ArC) separately screened titles and abstract of articles and reviewed all the safety data on MACEs or CHF in selected articles.

For each study selected, the following data were extracted and listed:

- Study design/characteristics: study design, trial duration.
- Patient characteristics: number of patients in each arm, age, gender ratio, disease duration, proportion of patients with PsA, baseline Psoriasis Area and Severity Index score and baseline body surface area score.
- Intervention: treatment arm (biological therapy or placebo).
- Primary outcomes: number of MACEs or CHF in each arm during the placebo-controlled phase.

To take into consideration the duration of the placebo-controlled phase and the number of patients in each arm, the results were expressed in patient-year (PY) in...
the biological group and the control group to enable the comparison between RCTs.

The Jadad Scale was used for quality assessment for RCTs.

Primary outcome measures were the incidence of MACEs or CHF during the placebo-controlled phase of biological treatment. We included all the events that occurred in patients who received at least 1 dose of study agent or placebo. MACEs or CHF that occurred during the follow-up open-label period were not considered.

The results of each selected study were presented as risk differences with a 95% CI. This MA was carried out using the inverse variance method whereby estimates of each study were pooling using a fixed or random effects model according to the level and significance of heterogeneity. Heterogeneity was tested with Cochran’s Q-test and evaluated by I² statistic with the following classification: 0% to 30% indicated negligible heterogeneity, 30% to 50% moderate heterogeneity, 50% to 75% substantial heterogeneity and high heterogeneity for values above 75%. For the Q-test, a p value less than 0.10 was considered as significant, and a random-effects model was used.

Analysis involved use of RevMan V.5.3 (The Nordic Cochrane Centre, The Cochrane Collaboration, 2011) for the MA calculations. P values less than 0.05 were considered statistically significant.

RESULTS

About 753 references met the selection criteria for this MA, which was carried out until December 2017.

Finally, 62 articles met the inclusion criteria, and 12 articles were added after manual searches. The details of the selection process are available in the flow chart (eg, figure 1).

The CV safety data were extracted from 74 articles, corresponding to 77 RCTs (3 articles that investigated safety data in 2 RCTs). The summary of the RCTs included in the MA is presented in (eg, (online supplementary table 1)).

The main baseline characteristics of patients included in the RCTs selected for this MA are summarised in (eg, (online supplementary table 2)).

The JADAD Scale for each RCT is presented in (eg, (online supplementary table 3)).

Primary outcome measure: MACEs

► In the anti-TNF group: 6 of the 2461 P-Y in the anti-TNF group and 5 of the 1396 P-Y in the placebo group had a MACE. The risk difference was 0.00; 95% CI −0.01 to 0.01 event/P-Y; p=0.96 (figure 2).

► In the anti-IL12/23 group: 10 of the 1150 P-Y in the anti-IL12/23 and 1 of the 652 P-Y in the placebo group had a MACE. The risk difference was 0.01; 95% CI −0.00 to 0.02 event/P-Y; p=0.21 (figure 3).

► In the anti-IL23 group: 4 of the 1188 P-Y in the anti-IL23 and 1 of the 374 P-Y in the placebo group had a MACE. The risk difference was 0.00; 95% CI −0.01 to 0.01 event/P-Y; p=0.77 (figure 4).

► In the anti-IL17 group: 8 of the 2169 P-Y of the anti-TNF group and 5 of the 784 P-Y in the placebo group had a MACE. The risk difference was −0.00; 95% CI −0.01 to 0.01 event/P-Y; p=0.84 (eg, figure 5).

Co-primary outcome measure: CHF

► In the anti-TNF group: 0 of the 2461 P-Y in anti-TNF group and 0 of the 1396 P-Y in the placebo group had a CHF. The risk difference was 0.00; 95% CI −0.01 to 0.01 event/P-Y; p>0.99 (eg, (online supplementary figure 6)).

► In the anti-IL12/23 group: 0 of the 1150 P-Y in the anti-IL12/23 and 1 of 652 P-Y in the placebo group had a CHF. The risk difference was −0.00; 95% CI −0.01 to 0.01 event/P-Y; p=0.98 (eg, (online supplementary figure 7)).
In the anti-IL23 group: 0 of the 1188 P-Y in the anti-IL23 and 0 of 374 P-Y in the placebo group had a CHF. The risk difference was 0.00; 95% CI −0.01 to 0.01 event/P-Y; p>0.99 (eg, (online supplementary figure 8)).

In the anti-IL17 group: 1 of the 2169 P-Y of the anti-TNF group and 0 of the 784 P-Y in the placebo group had a CHF. The risk difference was 0.00; 95% CI −0.01 to 0.01 event/P-Y; p=0.99 (eg, (online supplementary figure 9)).

Discussion
In this MA of RCTs, there was no significant change in the short-term risk of MACEs or CHF in patients with PsA or psoriasis initiating a biological therapy.

The first MA published in 2011, which focused on psoriasis, included 15 RCTs on anti-TNF (adalimumab, etanercept and infliximab) and 9 RCTs on anti-IL12/23 (ustekinumab and briakinumab) agents and showed no significant difference in the rate of MACEs observed in patients receiving anti-TNF or anti-IL12/23 agents, in comparison to the placebo.12

A second MA published in 2012, which focused on psoriasis, included 9 RCTs on anti-IL12/23 (ustekinumab and briakinumab) agents and showed an increased rate...
Psoriatic arthritis

of MACEs in patients who received anti-IL12/23 agents in comparison to the placebo.87

A third MA published in 2015, which focused on PsA and psoriasis, included six observational studies on systemic therapies (mixed biological therapies, non-biological therapies, corticosteroids and NSAIDs), and concluded that there was a decreased risk of CV events with systemic therapies in comparison to no systemic therapy or topical treatment.13

A fourth MA, published in 2016, which included 35 RCTs on anti-TNF (infliximab, etanercept and adalimumab), anti-IL12/23 (ustekinumab) and anti-IL17 (secukinumab and ixekizumab) agents, showed no statistically significant difference in the rate of MACEs observed in patients receiving biological therapies in comparison to the placebo.4

Our results are consistent with those of the first and fourth MA, but differ from those of the second MA, which showed a statistically significant increase in the rate of MACEs in patients who received anti-IL12/23 therapies. Differences in methods of statistical analysis used to calculate the risk of MACE could explain these discordant results. Our MA was carried out using the inverse variance method while the MA by Tzellos et al87 used the Peto OR method which indicates that trials with no events in both arms were automatically given zero

Figure 3 Difference in the risk of MACEs in patients treated with anti-IL12/23 agents compared with the placebo in RCTs. IL, interleukin; MACEs, major adverse cardiovascular events; RCTs, randomised controlled trials.

Figure 4 Difference in the risk of MACEs in patients treated with anti-IL23 agents compared with the placebo in RCTs. IL, interleukin; MACEs, major adverse cardiovascular events; RCTs, randomised controlled trials.
### Study or Subgroup

|                      | Anti IL17 | Placebo | Risk Difference |
|----------------------|-----------|---------|-----------------|
|                      | Total     | Events  | Total Weight |
| Brodalumab           | No.       | P Y     | No. P Y        |
| Papp et al. 2012     | 0         | 37      | 0               |
| Mease et al. 2014    | 0         | 26      | 0               |
| Lebawah et al. 2015 (AMAGINE2) | 1     | 281     | 0               |
| Lebawah et al. 2015 (AMAGINE3) | 2     | 289      | 0               |
| Nakagawa et al. 2015 | 1         | 26      | 0               |
| **Subtotal (95% CI)** | **659**   | **175** | **26.1% 0.01 - 0.02** |

- Total events: 4
- Heterogeneity: $\chi^2 = 0.23$, df = 4 ($P = 0.99$); $I^2 = 0$
- Test for overall effect: $Z = 0.71$ ($P = 0.48$)

### Ixekizumab

|                      | Anti IL17 | Placebo | Risk Difference |
|----------------------|-----------|---------|-----------------|
|                      | Total     | Events  | Total Weight |
| Leonard et al. 2012  | 0         | 35      | 0               |
| Griffiths et al. 2015 (UNCOVER 2) | 0     | 161     | 0               |
| Griffiths et al. 2015 (UNCOVER 3) | 1     | 178     | 1               |
| Mease et al. 2015 (SPIRIT-P1)   | 0         | 96      | 0               |
| Nash et al. 2017 (SPIRIT-P2)    | 0         | 113     | 2               |
| **Subtotal (95% CI)** | **583**   | **193** | **15.0% -0.01 - 0.03** |

- Total events: 1
- Heterogeneity: $\chi^2 = 1.62$, df = 4 ($P = 0.80$); $I^2 = 0$
- Test for overall effect: $Z = 0.76$ ($P = 0.45$)

### Secukinumab

|                      | Anti IL17 | Placebo | Risk Difference |
|----------------------|-----------|---------|-----------------|
|                      | Total     | Events  | Total Weight |
| Papp et al. 2012     | 0         | 51      | 2               |
| Rich et al. 2012     | 0         | 78      | 0               |
| McInnes et al. 2013  | 0         | 13      | 0               |
| Blauvelt et al. 2014 (FEASURE) | 2     | 27      | 0               |
| Langley et al. 2014 (ERASURE) | 0     | 113     | 0               |
| Langley et al. 2014 (FIXTURE) | 1     | 155     | 0               |
| Mease et al. 2015 (FUTURE 1)  | 1         | 124     | 0               |
| McInnes et al. 2015 (FUTURE 2) | 0         | 92      | 0               |
| Paul et al. 2015 (JUNCTURE) | 0     | 28      | 0               |
| Bagel et al. 2017    | 0         | 11      | 0               |
| Gottlieb et al. 2017 (GESTURE) | 0     | 31      | 0               |
| Mease et al. 2017 (FUTURE S) | 0         | 204     | 0               |
| **Subtotal (95% CI)** | **927**   | **416** | **58.5% 0.00 - 0.01** |

- Total events: 3
- Heterogeneity: $\chi^2 = 3.46$, df = 11 ($P = 0.98$); $I^2 = 0$
- Test for overall effect: $Z = 0.18$ ($P = 0.86$)

### Figure 5

- Difference in the risk of MACEs in patients treated with anti–IL17 agents compared with the placebo in RCTs.
- IL, interleukin; MACEs, major adverse cardiovascular events; RCTs, randomised controlled trials.

**Discussion**

Several methodological issues explain the discordant results between our MA and the MA by Roubille et al. which included a few observational studies and both conventional synthetic and biological therapies.

However, even though our MA shows no statistically significant difference in terms of MACE incidence in patients receiving anti-IL12/23 biological Disease Modifying AntiRheumatic Drug (DMARDs) in comparison to the placebo, we observed a trend of increased risk of MACEs in patients initiating a treatment with briakinumab and to a lesser extent with ustekinumab in comparison to the placebo. It is interesting to note that briakinumab was withdrawn from further clinical development in 2011 because MACE were reported during phase II and phase III clinical trials in subjects with psoriasis. Further randomised trials involving bDMARDs targeting IL12/23 pathway could increase the number of events in treated patients compared with placebo and then improve the power of the statistical test. If this trend was confirmed in further studies, the observed difference might become statistically significant in future MA.

No increase in the risk of MACEs was observed in the anti-IL23 (guselkumab, risankizumab and tildrakizumab) group compared with the placebo group in our MA.

Our MA shows no statistically significant difference in the short-term risk of CHF in patients with PsA or psoriasis initiating a biological therapy. No previous MA has assessed the short-term risk of CHF at the initiation of biological therapies in patients with PsA or psoriasis.

Several limitations should be considered when interpreting the findings of our MA. First, MACEs were not clearly defined in all the RCTs. Second, the short duration of the placebo-controlled phase of the RCTs only enabled a short-term evaluation of the impact of biological therapies on the risk of MACEs or CHF. Third, the non-inclusion criteria of RCTs usually excluded patients at high risk of MACEs and CHF.

Our MA has certain strengths in comparison to previous MAs. First, it assessed the risk of MACEs in both PsA and...
Psoriatic arthritis, based on the assumption that in both of these chronic inflammatory diseases, there is an increase in the risk of MACEs and they both require several biological therapies. This resulted in a higher number of patients being included in this MA in comparison to the previous MAs. Second, it investigated the risk of CHF, which represents another aspect of CV events that affects patients with chronic inflammatory diseases. Third, it showed that patients with chronic inflammatory diseases, there is an increase in the short-term risk of MACEs or CHF in patients with psoriasis, based on the assumption that in both of these chronic inflammatory diseases, there is an increase in the risk of MACEs or CHF in patients with PsA or psoriasis.

While both PsA and psoriasis have been associated with an increased risk of CV morbidity and mortality, the potential impact of DMARDs on the risk of CV events should be taken into consideration by rheumatologists and dermatologists in terms of treatment decisions in clinical practice. In a short-term perspective, the results of our MA, which did not reveal any significant change in the risk of MACE or CHF in patients with PsA or psoriasis initiating a bDMARD, should reassure the physicians about the CV safety of these therapeutic agents. In a long-term perspective, the use of anti-TNF agents was associated with a lower CV event risk in patients with psoriasis in comparison with methotrexate or phototherapy and with a lower risk of myocardial infarction in patients with rheumatoid arthritis in comparison with csDMARDs.

Studies involving a larger number of patients as well as a longer duration of treatment exposure are needed to evaluate the impact of anti-IL12/23, anti-IL23 or anti-IL17 agents on the risk of MACEs in patients with PsA or psoriasis in clinical practice.

Our MA, which is focused on the placebo-controlled phase of RCTs, did not reveal any significant change in the short-term risk of MACEs or CHF in patients with PsA or psoriasis initiating an anti-TNF, anti-IL12/23, anti-IL23 or anti-IL17 agent in comparison to the placebo. Data from the long-term extension phases of these RCTs and from the long-term follow-up of patients with PsA and psoriasis included in biological therapy registries are required to further characterise the long-term impact of biological therapies on the risk of MACEs or CHF.

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