Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
The risk of respiratory tract infections and interstitial lung disease with interleukin 12/23 and interleukin 23 antagonists in patients with autoimmune diseases: A systematic review and meta-analysis

Shintaro Akiyama, MD, PhD, Akihiro Yamada, MD, PhD, Dejan Micic, MD, and Atsushi Sakuraba, MD, PhD

Chicago, Illinois and Chiba, Japan

Background: Respiratory tract infections (RTIs) and interstitial lung disease (ILD) secondary to interleukin (IL) 12/23 or IL-23 antagonists have been reported in autoimmune diseases.

Objective: To assess the risk of RTIs and noninfectious ILD with these drugs.

Methods: We conducted a systematic review and meta-analysis of randomized controlled trials. Risk of RTIs and noninfectious ILD was compared to placebo by Mantel-Haenszel risk difference. We divided RTIs into upper RTIs (URTIs), viral URTIs, and lower RTIs (LRTIs) including infectious pneumonia. Noninfectious ILD included ILD, eosinophilic pneumonia, and pneumonitis.

Results: We identified 54 randomized controlled trials including 10,907 patients with 6 IL-12/23 or IL-23 antagonists and 5175 patients with placebo. These drugs significantly increased the risk of RTIs (Mantel-Haenszel risk difference, 0.019; 95% confidence interval, 0.005-0.033; \(P = .007\)), which was attributed to URTIs, but not viral URTIs or LRTIs. There was no significant difference in infectious pneumonia and noninfectious ILD between 2 groups.

Limitations: Because of the rarity of infectious pneumonia and ILD, sensitivity analysis was required.

Conclusions: The use of IL-12/23 or IL-23 antagonists for autoimmune diseases increased the risk of URTIs, but not viral URTIs, LRTIs, and noninfectious ILD. (J Am Acad Dermatol 2021;84:676-90.)

Key words: autoimmune diseases; IL12/23 and IL23 antagonists; meta-analysis; noninfectious interstitial lung disease; respiratory tract infections.

The clinical benefit of interleukin (IL) 12 and IL-23 inhibition has been shown in psoriasis and Crohn’s disease (CD) by briakinumab or ustekinumab. Moreover, IL-23-specific antagonists, such as tildrakizumab, risankizumab, guselkumab, and brazikumab have completed phase 2 or 3 trials. Currently, IL-12/23 or IL-23 antagonists are the second most commonly prescribed category of biologics for psoriasis and CD, behind anti–tumor necrosis factor agents.
However, randomized controlled trials (RCTs) of these drugs reported respiratory tract infections (RTIs) as the most common adverse events. Furthermore, the surveillance conducted by the US Food and Drug Administration (FDA) reported the development of noninfectious interstitial lung disease (ILD) after ustekinumab. Hence, physicians need evidence to decide whether to continue or hold these drugs, particularly during the current COVID-19 pandemic.

This systematic review and meta-analysis aimed to determine the risk of RTIs and noninfectious ILD with anti–IL-12/23 or anti–IL-23 agents in autoimmune diseases.

**METHODS**

**Search strategy and study selection**

This meta-analysis was conducted according to an a priori defined protocol based on Preferred Reporting Items for Systematic Reviews and Meta-analyses (PRISMA) guidelines. The protocol of this meta-analysis has been submitted to the International Prospective Register of Systematic Reviews (PROSPERO).

We searched PubMed/MEDLINE, Google Scholar, Scopus, Embase, ClinicalTrials.gov (https://clinicaltrials.gov/), and the Cochrane database from inception to February 1, 2019, to identify studies assessing the efficacy and safety of anti–IL-12/23 and anti–IL-23 therapies in autoimmune diseases. We also searched abstracts from medical conferences and bibliographies of identified articles for additional references. For Google Scholar, only the first 1000 articles were reviewed because this is the maximum number of results provided by the database. When a study registered in ClinicalTrials.gov or presented as an abstract became later available as an article, data were updated accordingly.

As for inclusion criteria, we considered RCTs reporting the incidence of adverse events, including RTIs and noninfectious ILD, with anti–IL-12/23 and anti–IL-23 therapies. There were no restrictions regarding age, sex, or duration of the study. We imposed no geographic or language restrictions. Two authors (SA and DM) independently screened each of the potential trials, abstract, and/or full article to determine whether they were eligible for inclusion. Area of disagreement or uncertainty were resolved by consensus among the authors. Studies were identified with the terms “briakinumab,” “ustekinumab,” “tildrakizumab,” “guselkumab,” “risankizumab,” “brazikumab,” “mirikizumab,” “LY2525623,” “anti–interleukin-12/23,” “anti–interleukin-23,” “anti–IL-12/23,” “anti–IL-23,” “anti–interleukin-12,” and “anti–IL-12” (both as medical subject headings and free text terms). RCTs without placebo-controlled groups were excluded. The search strategy is described in Fig 1.

**Data extraction and quality assessment**

All data were independently abstracted in duplicate by 2 authors (SA and DM) by using a data extraction form. Data on the study characteristics, such as author name, year of publication, study design, duration, sample size, age of patients, type of medications, and incidence of events were collected. Several published studies included data from multiple RCTs with different regimens or participant characteristics. For instance, the study published by Gordon et al included a comparison between ustekinumab and placebo and another comparison between risankizumab and placebo. These comparisons were considered as separate individual RCTs in our meta-analysis to ensure proper comparison and to avoid selection bias. The Jadad score was used to assess the quality of RCTs. We also used Cochrane risk-of-bias assessment instrument to evaluate the quality of the RCTs.

The Grading of Recommendations Assessment, Development, and Evaluation (GRADE) was applied to assess the certainty of evidence obtained from this meta-analysis.

**Outcome assessment**

The primary outcome measure of interest was the risk difference (RD) of the development of RTIs and noninfectious ILD among patients on anti–IL-12/IL-23 or anti–IL-23 agents compared with placebo. Subgroup analyses with each monoclonal antibody and underlying disease were performed. Data were analyzed based on the intention-to-treat principle except where indicated. We determined the number of each adverse event from the articles and the ClinicalTrials.gov database. We sorted RTIs into the following 3 categories: (1) upper RTIs (URTIs), (2) viral URTIs, and (3) lower RTIs (LRTIs). Based on Medical Dictionary for Regulatory
Activities (MedDRA) Terminology (https://biportal.bioontology.org/ontologies/MEDDRA), URTIs included the following diagnoses: nasopharyngitis, laryngitis, pharyngitis, rhinitis, sinusitis, tonsillitis, pharyngotonsillitis, tracheobronchitis, and upper respiratory infection. Viral URTIs included influenza and viral URTI. LRTIs were bronchitis, LRTI, and pneumonia. We included the following diseases in noninfectious ILD: ILD, eosinophilic pneumonia, and pneumonitis.

Statistical analysis

We undertook a meta-analysis with a random effects model. Mantel-Haenszel (MH) RD was used as our primary method. As for rare events, including infectious pneumonia and noninfectious ILD, we performed the sensitivity analysis as described in the “Results” section because of uncertainty regarding the preferred method for rare events. First, we analyzed data by MH odds ratio (OR), Peto OR, and MH risk ratio (RR) by excluding double-zero-event studies. Second, data were pooled among each drug, and then meta-analysis was performed by MH RD, MH OR, Peto OR, and MH RR. Finally, data were assessed by MH OR, Peto OR, and MH RR by adding 0.5 continuity correction or treatment arm correction to zero-event studies.

We evaluated the presence of heterogeneity across trials by using the $I^2$ statistic. An $I^2$ value of less than 25% indicated low heterogeneity, 25% to 75% indicated moderate heterogeneity, and greater than 75% indicated considerable heterogeneity. Heterogeneity was evaluated by using the Cochran $Q$ statistic with a significance level of $P < .10$. Begg and Egger tests were performed to assess publication bias, and funnel plots were constructed to visualize possible asymmetry when 3 or more studies were available.

Statistical analyses were performed using Comprehensive Meta-Analysis Software, version 2.0 (Biosstat, Englewood, NJ). All statistical tests except for the $Q$ statistic used a 2-sided $P$ value of .05 for significance.

RESULTS

Study characteristics

We identified 21,102 citations through the literature search, excluded 21,030 titles and abstracts after the initial screening, and assessed 72 studies for eligibility. A final number of 43 full-text articles and 2 studies registered only in ClinicalTrials.gov met all eligibility criteria and included 54 RCTs with a total of 10,907 patients with anti–IL-12/IL-23 or anti–IL-23 antibodies and 5175 with placebo (Fig 1). The 54 RCTs included 1 study of brazikumab (59 patients), 8 of briakinumab (1817 patients), 9 of guselkumab (1321 patients), 5 of risankizumab (830), 5 of tildrakizumab (1596 patients), and 26 of ustekinumab (5284 patients). All of the included studies are randomized, double-blind, placebo-controlled studies and have high scores in the Jadad scoring system. The characteristics and outcomes of the included studies are summarized in Table 1. The mean ages of patients and percentages of male patients were 43.8 years and 66.6% for guselkumab, 42.8 years and 65.4% for risankizumab, 37.6 years and 62.4% for tildrakizumab, 43.3 years and 49.5% for briakinumab, and 43.3 years and 54.0% for ustekinumab. The percentages of studies that permitted use of concomitant drugs (eg, corticosteroids, budesonide, thiopurines, methotrexate, calcineurin inhibitors, or aminosalicylates) during the trials were 40.0% for risankizumab, 43.3 years and 66.6% for guselkumab, 42.8 years and 65.4% for risankizumab, 37.6 years and 62.4% for tildrakizumab, 43.3 years and 49.5% for briakinumab, and 43.3 years and 54.0% for ustekinumab. The percentages of studies that permitted use of concomitant drugs (eg, corticosteroids, budesonide, thiopurines, methotrexate, calcineurin inhibitors, or aminosalicylates) during the trials were 40.0% for risankizumab, 38.5% for ustekinumab, 37.5% for briakinumab, 0% for guselkumab, and 0% for tildrakizumab. As for brazikumab, 1 study for CD was included in this analysis and permitted concomitant drugs.

RTIs

Meta-analysis with a random effects model showed that the overall risk of RTIs with anti–IL-12/IL-23 or anti–IL-23 agents was significantly higher than that of placebo (MH RD, 0.019; 95% confidence interval [CI], 0.005-0.033; $P = .007$) (Fig 2). The number needed to harm of RTIs was 58.8. Subgroup analysis showed a significantly increased risk of RTIs with briakinumab (MH RD, 0.021; 95% CI, 0.001-0.041; $P = .036$) and risankizumab (MH RD, 0.040; 95% CI, 0.005-0.076; $P = .026$). Heterogeneity was absent ($I^2 = 0$%) in overall and subgroup analyses except for briakinumab ($I^2 = 31$%). Funnel plot showed no asymmetry, therefore suggesting there were no
small-study effects or publication biases, which was supported by Begg and Egger tests (Supplemental Fig 1; available via Mendeley at https://doi.org/10.17632/j7dfkr2s8v.1). We also assessed the differential risk of RTIs by underlying disease and showed a significantly increased risk of RTIs in psoriasis (MH RD, 0.023; 95% CI, 0.010-0.036; \( P < .001 \)) and ankylosing spondylitis (MH RD, 0.136; 95% CI, 0.006-0.265; \( P = .040 \)) (Supplemental Fig 2; available via Mendeley at https://doi.org/10.17632/j7dfkr2s8v.1).

We divided RTIs into URTIs, viral URTIs, and LRTIs and investigated each risk with IL-12/23 or IL-23 inhibitors. The overall risk of URTIs was significantly higher in the treatment group compared to placebo (MH RD, 0.017; 95% CI, 0.005-0.029; \( P = .006 \)) (Supplemental Fig 3, A; available via Mendeley at https://doi.org/10.17632/j7dfkr2s8v.1). Subgroup analysis showed an elevated risk of URTIs with risankizumab (MH RD, 0.028; 95% CI, 0.004-0.053; \( P = .024 \)). No publication bias was detected by Begg and Egger tests (Supplemental Fig 3, B).

Anti–IL-12/IL-23 or anti–IL-23 agents did not increase the overall risks of viral URTIs (MH RD, 0.001; 95% CI, -0.002 to 0.003; \( P = .60 \)) and LRTIs (MH RD, 0; 95% CI, -0.002 to 0.002; \( P = .71 \)) (Supplemental Figs 4, A and 5, A; available via Mendeley at https://doi.org/10.17632/j7dfkr2s8v.1). Heterogeneity was absent (\( I^2 = 0\% \)) in these analyses. Publication bias was indicated in the analysis of viral URTIs (Begg: \( P < .001 \); Egger: \( P = .019 \)) (Supplemental Fig 4, B) and LRTIs (Begg: \( P < .001 \); Egger: \( P = .55 \)) (Supplemental Fig 5, B), but
the funnel plots did not appear asymmetric on visual inspection.

**Infectious pneumonia and noninfectious ILD**

The total numbers of infectious pneumonia were 9 and 4 cases in the treatment group and placebo, respectively. *Mycobacterium tuberculosis* and viral pneumonia were not reported. The overall risk of infectious pneumonia was not significantly increased in the treatment group compared to placebo (MH RD, 0; 95% CI, -0.002 to 0.002; \( P = .87 \)) (Fig 3). Heterogeneity was absent (\( I^2 = 0 \%)\). The funnel plot was not asymmetric, indicating no publication bias, which was supported by Egger test (\( P = .93 \)) but not Begg test (\( P < .001 \)) (Supplemental Fig 6; available via Mendeley at https://doi.org/10.17632/j7dkfr2s8v.1).

In terms of noninfectious ILD, 2 and 1 cases were identified in the treatment and placebo groups, respectively. All 3 cases were reported in trials of ustekinumab. Other potential risk factors such as age and sex were not different among the drugs. The sensitivity analysis showed consistent results (Supplemental Tables 1-6; available via Mendeley at https://doi.org/10.17632/rdxgpw9yxk.2), except the analysis with 0.5 constant correction of zero-event studies showed a lower risk of infectious pneumonia (Supplemental Table 7; available via Mendeley at https://doi.org/10.17632/rdxgpw9yxk.2) and ILD in the treatment group (Supplemental Table 8; available via Mendeley at https://doi.org/10.17632/rdxgpw9yxk.2).

**Grading the quality of evidence**

Based on the GRADE criteria, the overall quality of evidence for this analysis was moderate because infectious pneumonia and ILD were rare events (Supplemental Tables 9 and 10; available via Mendeley at https://doi.org/10.17632/rdxgpw9yxk.2).

**DISCUSSION**

Our meta-analysis showed that IL-12/23 or IL-23 inhibitors increased the risk of RTIs, especially URTIs, but not viral URTIs and LRTIs, and noninfectious ILD in autoimmune diseases.

We found that risankizumab and briakinumab particularly enhanced the risk of RTIs and hypothesized that concomitant therapies during the trials might differentiate the risk of RTIs. In terms of anti–IL-23 agents, risankizumab showed a higher rate of RCTs that permitted concomitant therapies (40.0%) compared with guselkumab (0%) and tildrakizumab (0%). Among RCTs of risankizumab, the only study reporting an increased risk of RTIs was performed in patients with ankylosing spondylitis, who were permitted to use conventional disease-modifying antirheumatic drugs or low-dose systemic steroids. This suggests that combination therapy of anti–IL-23 agents with immunosuppressants might work synergistically to surface the risk of RTIs. As for anti–IL-12/IL-23 agents, each of briakinumab and ustekinumab has a similar percentage of RCTs that permitted concomitant drugs (37.5% for briakinumab and 38.5% for ustekinumab). Other potential risk factors such as age and sex were not different among the drugs. Given that briakinumab has been withdrawn from the application with the FDA because of severe adverse events, the difference in risk of RTIs among the 2 drugs would be explained by different properties of these drugs.

Our study might support that anti–IL-12/23 and anti–IL-23 therapies can be safely used for autoimmune diseases even during the current COVID-19 pandemic. However, given that influenza vaccination is generally recommended for patients with autoimmune diseases receiving immunosuppressive therapies, viral URTIs caused by influenza virus could be prevented in both the treatment and placebo groups, and our finding regarding the risk of viral URTIs might be affected. In addition, studies included in this analysis were conducted before the pandemic and the risk of severe acute respiratory syndrome coronavirus 2 in patients with anti–IL-12/23 or anti–IL-23 therapies could not be assessed. Further studies, particularly during the current pandemic, are necessary to provide enough evidence to ensure the safety of these drugs. Additional investigations are also needed to understand the differential risk of RTIs in psoriasis and ankylosing spondylitis and the mechanism of ILD by IL-12/23 inhibition because patients who are indicated for these drugs, namely psoriasis, rheumatoid arthritis, and inflammatory bowel disease, all carry an increased risk of lung disease.
| Drugs       | Target | Disease                  | References                          | Age, y | Sex, % male | Study duration, wk | Concomitant therapies during trials | Regimen, mg (unless noted otherwise) | Jadad score | Patients, N | RTIs, n | Infectious pneumonia, n | Non infectious ILD, n |
|-------------|--------|--------------------------|-------------------------------------|--------|-------------|------------------|-------------------------------------|-------------------------------------|-------------|--------------|---------|----------------------|---------------------|
| Brazikumab  | IL-23  | CD                       | Sands et al (2017)¹¹               | 37     | 38          | 12               | Yes                                 | 700 IV at wk 0, 4                   | 5           | 59           | 60       | 9        | 11      | 0       | 0       | 0       |
| Guselkumab  | IL-23  | Psoriasis                | Ohtsuki et al (2018)³³             | 50     | 68          | 16               | No                                  | 50, 100 SC at wk 0, 4, 12           | 5           | 128          | 64       | 22       | 7       | 0       | 0       | 0       |
|             |        |                          |                                    | 46     | 68          | 16               | No                                  | 100 SC at wk 0, 4, 12               | 4           | 62           | 16       | 13       | 1       | 0       | 0       | 0       |
|             |        |                          | Nemoto et al (2018)³⁴             | NA     | 60          | 24               | No                                  | 10, 30, 100, 300 SC (s)             | 5           | 20           | 4        | 2        | 0       | 0       | 0       | 0       |
|             |        |                          | Blauvelt et al (2017)³⁵           | 44     | 73          | 16               | No                                  | 100 SC at wk 0, 4, 12               | 5           | 329          | 174      | 55       | 26      | 0       | 0       | 0       |
|             |        |                          | Reich et al (2017)¹⁰             | 44     | 70          | 16               | No                                  | 100 SC at wk 0, 4, 12               | 5           | 494          | 248      | 51       | 26      | 0       | 0       | 0       |
|             |        |                          | Gordon et al (2015)⁹              | 44     | 72          | 16               | No                                  | 5, 50, 200 SC at wk 0, 4, q12 wk; 15, 100 SC q8 wk | 5           | 207          | 42       | 25       | 3       | 0       | 0       | 0       |
| PPP         |        |                          | Sofen et al (2014)³⁶             | 43     | 63          | 24               | No                                  | 10, 30, 100, 300 SC (s)             | 4           | 20           | 4        | 5        | 0       | 0       | 0       | 0       |
|             |        |                          | Terui et al (2018)³⁷             | 52     | 29          | 24               | No                                  | 200 SC at wk 0, 4                   | 5           | 25           | 24       | 8        | 9       | 0       | 0       | 0       |
|             |        |                          | Zhuang et al (2016)³⁸            | 27     | 96          | 16               | No                                  | 0.03, 0.1, 0.3, 1, 3, 10 mg/kg IV; 3 mg/kg SC (s) | 5           | 36           | 11       | 5        | 1       | 0       | 0       | 0       |
| Risankizumab| IL-23  | Psoriasis                | Gordon et al (2018)⁷              | 48     | 71          | 16               | No                                  | 150 SC at wk 0, 4                   | 5           | 304          | 102      | 37       | 8       | 0       | 0       | 0       |
|             |        |                          | Gordon et al (2018)⁷              | 47     | 68          | 16               | No                                  | 150 SC at wk 0, 4                   | 5           | 294          | 98       | 21       | 4       | 0       | 0       | 0       |
|             |        |                          | Krueger et al (2015)³⁹           | 42     | 81          | 24               | No                                  | 0.01, 0.05, 0.25, 1, 3, 5 mg/kg IV; 0.25, 1 mg/kg SC (s) | 5           | 31           | 8        | 11       | 2       | 0       | 0       | 0       |
| Ankylosing  |        | spondylitis              | Baeten et al (2018)⁶⁴             | 38     | 70          | 16               | Yes                                 | 18 SC (s); 90, 180 SC at wk 0, 8, 16 | 5           | 119          | 40       | 31       | 5       | 0       | 0       | 0       |

Continued
| Drugs          | Target  | Disease           | References                     | Age, y | Sex, % male | Study duration, wk | Concomitant therapies during trials | Regimen, mg (unless noted otherwise) | Jadad score | Patients, N | IL-12/23 Placebo | IL-12/23 Placebo | IL-12/23 Placebo | IL-12/23 Placebo | RTIs, n | Infectious pneumonia, n | Non infectious ILD, n |
|----------------|---------|-------------------|--------------------------------|--------|-------------|-------------------|-------------------------------------|------------------------------------|-------------|--------------|------------------|------------------|------------------|------------------|--------|---------------------|------------------|
| CD             | Feagan et al (2017) | 39 | 37 | 12 | Yes | 200, 600 IV at wk 0, 4, 8 | 5 | 82 | 39 | 10 | 5 | 1 | 1 | 0 | 0 |
| Tildrakizumab IL-23 | Psoriasis | Reich et al (2017) | 46 | 67 | 12 | No | 100, 200 SC at wk 0, 4 | 5 | 617 | 154 | 69 | 17 | 0 | 0 | 0 | 0 |
|              |          | Reich et al (2017) | 46 | 71 | 12 | No | 100, 200 SC at wk 0, 4 | 5 | 621 | 156 | 76 | 12 | 0 | 0 | 0 | 0 |
|              |          | Papp et al (2015) | 43 | 74 | 16 | No | 5, 25, 100, 200 SC at wk 0, 4, 16 | 5 | 308 | 45 | 63 | 11 | 0 | 0 | 0 | 0 |
|              |          | Khalilieh et al (2018) | 26 | 38 | 28 | No | 0.1, 0.5, 3, 10 mg/kg IV (s) | 5 | 22 | 7 | 0 | 0 | 0 | 0 | 0 | 0 |
|              |          | Khalilieh et al (2018) | 27 | 62 | 20 | No | 50, 200 SC (s) | 5 | 28 | 9 | 0 | 0 | 0 | 0 | 0 | 0 |
| Briakinumab IL-12 IL-23 | Psoriasis | Gordon et al (2012) | 46 | 69 | 12 | No | 200 SC wk 0, 4; 100 SC wk 8 | 5 | 981 | 484 | 114 | 40 | 2 | 0 | 0 | 0 |
|              |          | Gottlieb et al (2011) | 43 | 67 | 12 | No | 200 SC wk 0, 4; 100 SC wk 8 | 5 | 138 | 68 | 19 | 8 | 0 | 0 | 0 | 0 |
|              |          | Strober et al (2011) | 45 | 64 | 12 | No | 200 SC wk 0, 4; 100 SC wk 8 | 5 | 139 | 72 | 20 | 6 | 0 | 0 | 0 | 0 |
|              |          | Kimball et al (2008) | 46 | 75 | 12 | No | 200 SC (s); 100 SC q2wks for 12 wk; 200 SC q1wk for 4 wk; 200 SC q2wks or q1wk for 12 wk | 5 | 150 | 30 | 38 | 3 | 0 | 0 | 0 | 0 |
| CD | Panaccione et al (2015) | 36 | 51 | 12 | Yes | 200, 400, 700 IV at wk 0, 4, 8 | 5 | 200 | 46 | 0 | 0 | 0 | 0 | 0 | 0 |
|              | Mannon et al (2004) | 43 | 25 | 28 | Yes | 1, 3 mg/kg SC at wk 0; q1wk from wk 4 to 10 | 5 | 32 | 8 | 2 | 0 | 0 | 0 | 0 | 0 |
|              | Mannon et al (2004) | 40 | 20 | 25 | Yes | 1, 3 mg/kg SC q1wk for 7 wk | 5 | 31 | 8 | 2 | 0 | 0 | 0 | 0 | 0 |
| MS | Vollmer et al (2011) | 47 | 25 | 24 | No | 200 SC q2wk; 200 SC q1wk | 5 | 146 | 69 | 51 | 29 | 0 | 0 | 0 | 0 |
| Ustekinumab IL-12 IL-23 Psoriasis | Gordon et al (2018) | 48 | 71 | 16 | No | 45 SC (wt ≤ 100 kg); 90 SC (wt > 100 kg) at wk 0, 4 |
|-----------------------------------|---------------------|----|----|----|-----|---------------------------------|
| Gordon et al (2018)               | 47                  | 68 | 16 | No | 45 SC (wt ≤ 100 kg); 90 SC (wt > 100 kg) at wk 0, 4 |
| Landells et al (2015)             | 15                  | 49 | 12 | No | 0.75 mg/kg SC (wt ≤ 60 kg); 45 SC (60 < wt ≤ 100 kg); 90 SC (wt > 100 kg); 0.375 mg/kg SC (wt ≤ 60 kg); 22.5 SC (60 < wt ≤ 100 kg); 45 SC (>100 kg) at wk 0, 4 |
| Lebwohl et al (2015)              | 45                  | 69 | 12 | No | 45 SC (wt ≤ 100 kg); 90 SC (wt > 100 kg) at wk 0, 4 |
| Lebwohl et al (2015)              | 45                  | 68 | 12 | No | 45 SC (wt ≤ 100 kg); 90 SC (wt > 100 kg) at wk 0, 4 |
| Zhu et al (2013)                  | 40                  | 77 | 12 | No | 45 SC at wk 0, 4 |
| Igarashi et al (2012)             | 46                  | 80 | 12 | No | 45, 90 SC at wk 0, 4 |
| Tsai et al (2011)                 | 40                  | 85 | 12 | No | 45 SC at wk 0, 4 |
| Papp et al (2008)                 | 45                  | 69 | 12 | No | 45, 90 SC at wk 0, 4 |
| Leonardi et al (2008)             | 45                  | 69 | 12 | No | 45, 90 SC at wk 0, 4 |
| Drugs        | Target | Disease             | Age, y | Sex, % male | Study duration, wk | Concomitant therapies during trials | Regimen, mg (unless noted otherwise) | Jadad score | Patients, N | RTIs, n | Infectious pneumonia, n | Non infectious ILD, n |
|--------------|--------|---------------------|--------|-------------|-------------------|-------------------------------------|--------------------------------------|-------------|--------------|----------|---------------------|----------------------|
|              |        |                     |        |             |                   |                                     |                                      |             |              |          |                     |                      |
| Krueger et al (2007)51 | PsA     |                  | 46     | 59          | 20                | No                                  | 45, 90 SC at wk 0, q4wk             | 5           | 252          | 67       | 95     | 24       | 1 0 0 0 0          |
| Ritchlin et al (2014)52 |         |                   | 49     | 47          | 16                | Yes                                 | 45, 90 SC at wk 0, 4, 16             | 5           | 207          | 104      | 31     | 15       | 0 0 0 0 1          |
| McInnes et al (2013)52 |         |                   | 48     | 52          | 16                | Yes                                 | 45, 90 SC at wk 0, 4, 16             | 5           | 409          | 205      | 33     | 18       | 0 0 0 0 0          |
| Gottlieb et al (2009)53 |         |                   | 50     | 59          | 12                | Yes                                 | 90 SC q1wk for 4 wk                  | 5           | 76           | 70       | 20     | 12       | 0 0 0 0 0          |
| Feagan et al (2016)54 | CD      |                  | 37     | 40          | 8                 | Yes                                 | 130 mg, 6 mg/kg IV (s)               | 5           | 495          | 245      | 23     | 13       | 0 0 0 0 0          |
| Feagan et al (2016)54 |         |                   | 39     | 50          | 8                 | Yes                                 | 130 mg, 6 mg/kg IV (s)               | 5           | 419          | 208      | 24     | 10       | 0 0 0 0 0          |
| Sandborn et al (2012)5 |         |                   | 39     | 41          | 8                 | Yes                                 | 1, 3, 6 mg/kg IV (s)                 | 5           | 394          | 132      | 41     | 8        | 0 0 0 0 0          |
| Sandborn et al (2008)55 |         |                   | 40     | 55          | 8                 | Yes                                 | 90 SC at wk 0, 1, 2, 3; 4.5 mg/kg IV at wk 0 | 5           | 52           | 52       | 6      | 3        | 0 0 0 0 0          |
| Khattri et al (2017)56 | Atopic dermatitis |             | 37     | 63          | 16                | No                                  | 45 SC (wt ≤ 100 kg), 90 SC (wt > 100 kg) at wk 0, 4, 16 | 5           | 16           | 16       | 2      | 1        | 0 0 0 0 0          |
| Saeki et al (2017)51 | GVHD    |                  | 39     | 71          | 24                | No                                  | 45, 90 SC at wk 0, 4                | 5           | 52           | 27       | 15     | 7        | 0 0 0 0 0          |
| NCT0171340021 |         |                   | 53     | 63          | 52                | Yes                                 | 45 SC (wt ≤ 100 kg), 90 SC (wt > 100 kg) at day -1 and day 20 after transplantation | 5           | 15           | 15       | 1      | 3        | 0 0 0 0 0          |
| van Vollenhoven et al (2018)58 | SLE     |                  | 40     | 3           | 24                | Yes                                 | 260 (wt 35-55 kg), 390 (55 < wt ≤ 85), 520 (wt > 85 kg) | 5           | 60           | 42       | 21     | 12       | 1 0 0 0 0          |
| Disease   | Study Authors         | Age | Sex | Treatment | Follow-up | CD  | IL  | ILD | IV  | RTI | SLE  | NA  | PPP    | PPP  |
|-----------|-----------------------|-----|-----|-----------|-----------|-----|-----|-----|-----|-----|------|-----|--------|-------|
| Sarcoidosis | Judson et al (2014)   | 50  | 51  | 44        | Yes       | 180 SC wk 0; 90 SC wk 8, 16, 24 | 5   | 60  | 58  | 40  | 35  | 3    | 0   | 1     | 0     |
| MS        | Segal et al (2008)    | 37  | 36  | 37        | No        | 27, 90, 180 SC wk 0, 1, 2, 3, 7, 11, 15, 1990 SC q8wks | 5   | 200 | 49  | 73  | 17  | 0    | 0   | 0     | 0     |
| PPPP      | Bissonnette et al (2014) | 55  | 10  | 16        | No        | 45 SC (wt < 100 kg), 90 SC (wt ≥ 100 kg) at wk 0, 4, 16 | 5   | 10  | 10  | 0   | 0   | 0    | 0   | 0     | 0     |
| PPP       | Bissonnette et al (2014) | 50  | 0   | 16        | No        | 45 SC (wt < 100 kg), 90 SC (wt ≥ 100 kg) at wk 0, 16 | 5   | 5   | 8   | 0   | 1   | 0    | 0   | 0     | 0     |

*Regarding age and sex, data from overall patients in each trials or patients treated IL-12/23 or IL-23 antagonists were used.

This study was registered in ClinicalTrials.gov but later became available as an article.
naive T cells into interferon gamma–producing T helper type (Th) 1 cells, which contribute to viral clearance and prevent infections of *Mycobacterium* and *Salmonella* species. Meanwhile, IL-23 maintains IL-17–producing Th17 cells, and the deficiency of IL-17 immunity results in infections of *Candida* species. Our results showed that IL-12/23 or IL-23 antagonists did not increase the risk of LRTIs and infectious pneumonia, including *Mycobacteria tuberculosis* or any virus. As for viral RTIs, previous studies showed that IL-17 knockout mice had lower levels of lung inflammation by influenza virus compared with the wild type. A study of the Middle East respiratory syndrome coronavirus showed that a patient with a poor outcome had an increased level of IL-17 expression in the lung. These data suggest that IL-12/23 or IL-23 inhibitors might theoretically be preventive for severe acute respiratory syndrome coronavirus 2–induced pneumonia rather than detrimental in autoimmune diseases during the COVID-19 pandemic.

**Limitations**
First, this study did not assess the long-term effect of IL-12/23 or IL-23 antagonists on RTIs and ILD. However, 92.6% (50/54) of the included studies reported RTIs during placebo-controlled phases. The FDA reported whether the onset of ILD was acute or subacute, so our data would most likely include the incidence of these events. Second, regarding infectious pneumonia and ILD, many studies had zero events in both arms (87% [47/54] and 94% [51/54], respectively). Thus, we undertook comprehensive analyses that either included or excluded double-zero-event studies. The analysis with 0.5 constant correction showed a lower risk of

**Fig 2.** Meta-analysis of the Mantel-Haenszel (MH) risk difference of respiratory tract infections with IL-12/23 and IL-23 antagonists. CI, Confidence interval; IL, interleukin.
these events in the treatment group. We also used treatment arm correction because this method performed better than 0.5 constant correction to examine rare events. Third, our study may not reflect the risk in patients at high risk for RTIs because of the possible exclusion of patients with recent RTIs or chronic lung disease in clinical trials. Fourth, we categorized RTIs into URTIs, viral URTIs, and LRTIs based on MedDRA, which is widely used in clinical trials, but not so much in clinical research. Furthermore, the included studies were conducted before the pandemic. Hence, it does not provide evidence of whether there is an increase in RTIs or ILD during the pandemic in patients receiving IL-12/23 or IL-23 antagonists, nor whether these agents can be continued after a diagnosis of COVID-19. A meta-analysis of real-world studies of COVID-19 in patients with autoimmune diseases is needed.

CONCLUSION
This meta-analysis showed that IL-12/23 antagonists had an increased risk of RTIs, but not viral URTIs, LRTIs including infectious pneumonia, and noninfectious ILD.

---

**Table: Risk of RTIs with IL-12/23 antagonists**

| Drug              | Study name                        | Lower limit | Upper limit | Treatment effect | Control effect | Relative weight |
|-------------------|-----------------------------------|-------------|-------------|-----------------|---------------|----------------|
| Brazilkumab       | Sando et al 2017                  | 0.00         | 0.03         | 0.00            | 0.00          | 100.00         |
|                   |                                    | 0.00         | 0.02         | 0.00            | 0.00          | 100.00         |
|                   |                                    | 0.00         | 0.02         | 0.00            | 0.00          | 100.00         |

---

**Figure 3:** Meta-analysis of Mantel-Haenszel (MH) risk difference of infectious pneumonia with IL-12/23 and IL-23 antagonists. CI, Confidence interval; IL, interleukin.
REFERENCES

1. Gordon KB, Langley RG, Gottlieb AB, et al. A phase III, randomized, controlled trial of the fully human IL-12/23 mAb briakinumab in moderate-to-severe psoriasis. *J Invest Dermatol*. 2012;132:304-314.

2. Panaccione R, Sandborn WJ, Gordon GL, et al. Briakinumab for treatment of Crohn’s disease: results of a randomized trial. *Inflamm Bowel Dis*. 2015;21:1329-1340.

3. Papp KA, Langley RG, Lebwohl M, et al. Efficacy and safety of ustekinumab, a human interleukin-12/23 monoclonal antibody, in patients with psoriasis: 52-week results from a randomised, double-blind, placebo-controlled trial (PHOENIX 2). *Lancet*. 2008;371:1675-1684.

4. Sandborn WJ, Gasink C, Gao LL, et al. Ustekinumab induction and maintenance therapy in refractory Crohn’s disease. *N Engl J Med*. 2012;367:1519-1528.

5. Reich K, Papp KA, Blauvelt A, et al. Tildrakizumab versus placebo or etanercept for chronic plaque psoriasis (reSURFACE 1 and reSURFACE 2): results from two randomised, controlled, phase 3 trials. *Lancet*. 2017;390:276-288.

6. Papp K, Thaci D, Reich K, et al. Tildrakizumab (MK-3222), an anti–interleukin-23p19 monoclonal antibody, improves psoriasis in a phase IIb placebo-controlled trial. *Br J Dermatol*. 2015;173:930-939.

7. Gordon KB, Strober B, Lebwohl M, et al. Efficacy and safety of risankizumab in moderate-to-severe plaque psoriasis (UltIMMa-1 and UltIMMa-2): results from two randomised controlled trials. *Lancet*. 2017;389:276-288.

8. Feagan BG, Sandborn WJ, D’Haens G, et al. Induction therapy with the selective interleukin-23 inhibitor risankizumab in patients with moderate-to-severe Crohn’s disease: a randomised, double-blind, placebo-controlled and ustekinumab-controlled phase 3 trial. *Lancet*. 2017;390:650-661.

9. Gordon KB, Duffin KC, Bissonnelette R, et al. A phase 2 trial of guselkumab versus adalimumab for plaque psoriasis. *N Engl J Med*. 2015;373:137-147.

10. Reich K, Armstrong AW, Foley P, et al. Efficacy and safety of guselkumab, an anti–interleukin-23 monoclonal antibody, compared with adalimumab for the treatment of moderate-to-severe plaque psoriasis. *J Am Acad Dermatol*. 2021;84:1-12.

**Fig 4.** Meta-analysis of the Mantel-Haenszel (MH) risk difference of noninfectious interstitial lung disease with IL-12/23 and IL-23 antagonists. CI, Confidence interval; IL, interleukin.
of patients with moderate to severe psoriasis with randomized withdrawal and retreatment: results from the phase III, double-blind, placebo- and active comparator-controlled VOYAGE 2 trial. J Am Acad Dermatol. 2017;76:418-431.

11. Sands BE, Chen J, Feagan BG, et al. Efficacy and safety of MEDI0207, an antibody against interleukin 23, in patients with moderate to severe Crohn’s disease: a phase 2a study. Gastroenterology. 2017;153:77-86.

12. Frontinilla D, Vasiliou C, Sawalhi-Leckenby N, Kurniawan A. Crohn’s disease: disease coverage. Datamonitor Healthcare. Available at: https://pharmastore.infoforma.com/wp-content/uploads/2018/08/crohnsdisease_5415.pdf; 2018. Accessed April 2020.

13. Lopez-Ferrer A, Laiz A, Puig V. The safety of ustekinumab for the treatment of psoriatic arthritis. Expert Opin Drug Saf. 2017;16:733-742.

14. Brinker A, Cheng C, Chan V. Association of noninfectious pneumonia with ustekinumab use. JAMA Dermatol. 2019;155:221-224.

15. Zumla A, Niederman MS. Editorial: the explosive epidemic of novel coronavirus disease 2019 (COVID-19) and the persistent threat of respiratory tract infectious diseases to global health security. Curr Opin Pulm Med. 2020;26:193-196.

16. Lebwohl M, Rivera-Oyola R, Murrell DF. Should biologics for psoriasis be interrupted in the era of COVID-19? J Am Acad Dermatol. 2020;82:1217-1218.

17. Monteleone G, Ardizzone S. Are patients with inflammatory bowel disease at increased risk for Covid-19 infection? J Crohns Colitis. 2020;14:1334-1336.

18. Moher D, Liberati A, Tetzlaff J, Altman DG, Group P. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. J Clin Epidemiol. 2009;62:1006-1012.

19. Booth A. PROSPERO’s progress and activities 2012/13. Syst Rev. 2013;2:111.

20. Ferris LK, Ott E, Jiang J, et al. Efficacy and safety of guselkumab, administered with a novel patient-controlled injector (One-Press), for moderate-to-severe psoriasis: results from the phase 3 ORION study. J Dermatolog Treat. 2020;31:152-159.

21. Pidala J, Beato F, Kim J, et al. In vivo IL-12/IL-23p40 neutralization blocks Th1/Th17 response after allogeneic hematopoietic cell transplantation. Haematologica. 2018;103:531-539.

22. Jadad AR, Moore RA, Carroll D, et al. Assessing the quality of reports of randomized controlled trials: is blinding necessary? Control Clin Trials. 1996;17:1-12.

23. Higgins JP, Altman DG, Gotzsche PC, et al. The Cochrane Collaboration’s tool for assessing risk of bias in randomised trials. BMJ. 2011;343:d5928.

24. Guyatt GH, Oxman AD, Vist G, et al. GRADE guidelines: 4. rating the quality of evidence—study limitations (risk of bias). J Clin Epidemiol. 2011;64:407-415.

25. Friedrich JO, Adhikari NK, Beyene J. Inclusion of zero total event trials in meta-analyses maintains analytic consistency and incorporates all available data. BMC Med Res Methodol. 2007;7:5.

26. Brockhaus AC, Bender R, Skipka G. The Peto odds ratio viewed as a new effect measure. Stat Med. 2014;33:4861-4874.

27. Friedenreich CM. Methods for pooled analyses of epidemiologic studies. Epidemiology. 1993;4:295-302.

28. Bongartz T, Sutton AJ, Sweeting MJ, et al. Anti-TNF antibody therapy in rheumatoid arthritis and the risk of serious infections and malignancies: systematic review and meta-analysis of rare harmful effects in randomized controlled trials. JAMA. 2006;295:2275-2285.

29. Higgins JP, Thompson SG, Deeks JJ, Altman DG. Measuring inconsistency in meta-analyses. BMJ. 2003;327:557-560.

30. Higgins J, Green S. Cochrane Handbook for Systematic Reviews of Interventions, version 5.1.0. The Cochrane Collaboration; 2011.

31. Beggs CB, Mazumdar M. Operating characteristics of a rank correlation test for publication bias. Biometrics. 1994;50:1088-1101.

32. Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. BMJ. 1997;315:629-634.

33. Ohtsuki M, Kubo H, Morishima H, et al. Guselkumab, an anti-interleukin-23 monoclonal antibody, for the treatment of moderate to severe plaque-type psoriasis in Japanese patients: efficacy and safety results from a phase 3, randomized, double-blind, placebo-controlled study. J Dermatol. 2018;45:1053-1062.

34. Nemoto O, Hirose K, Shibata S, Li K, Kubo H. Safety and efficacy of guselkumab in Japanese patients with moderate-to-severe plaque psoriasis: a randomized, placebo-controlled, ascending-dose study. Br J Dermatol. 2018;178:689-696.

35. Blauvelt A, Papp KA, Griffiths CE, et al. Efficacy and safety of guselkumab, an anti-interleukin-23 monoclonal antibody, compared with adalimumab for the continuous treatment of patients with moderate to severe psoriasis: results from the phase III, double-blinded, placebo- and active comparator-controlled VOYAGE 1 trial. J Am Acad Dermatol. 2017;76:405-417.

36. Sofen H, Smith J, Matheson RT, et al. Guselkumab (an IL-23-specific mAb) demonstrates clinical and molecular response in patients with moderate-to-severe psoriasis. J Allergy Clin Immunol. 2014;133:1032-1040.

37. Terui T, Kobayashi S, Okubo Y, et al. Efficacy and safety of guselkumab, an anti-interleukin 23 monoclonal antibody, for palmoplantar pustulosis: a randomized clinical trial. JAMA Dermatol. 2018;154:309-316.

38. Zhuang Y, Calderon C, Marciniak SJ Jr, et al. First-in-human study to assess guselkumab (anti-IL-23 mAb) pharmacokinetics/safety in healthy subjects and patients with moderate-to-severe psoriasis. Eur J Clin Pharmacol. 2016;72:1303-1310.

39. Krueger JG, Ferris LK, Menter A, et al. Anti-IL-23A mAb BI 655066 for treatment of moderate-to-severe psoriatic: safety, efficacy, pharmacokinetics, and biomarker results of a single-rising-dose, randomized, double-blind, placebo-controlled trial. J Allergy Clin Immunol. 2015;136:116-124.

40. Khaliliieh S, Hodsdman P, Xu C, et al. Pharmacokinetics of tildrakizumab (MK-3322), an anti-IL-23 monoclonal antibody, after intravenous or subcutaneous administration in healthy subjects. Basic Clin Pharmacol Toxicol. 2018;123:294-300.

41. Gottlieb AB,Leonardi C, Kerdel F, et al. Efficacy and safety of briakinumab vs. etanercept and placebo in patients with moderate to severe chronic plaque psoriasis. Br J Dermatol. 2011;165:652-660.

42. Strober BE, Crowley JJ, Yamauchi PS, Olds M, Williams DA. Efficacy and safety results from a phase III, randomized controlled trial comparing the safety and efficacy of briakinumab with etanercept and placebo in patients with moderate to severe chronic plaque psoriasis. Br J Dermatol. 2011;165:661-668.

43. Kimball AB, Gordon KB, Langley RG, et al. Safety and efficacy of ABT-874, a fully human interleukin 12/23 monoclonal
antibody, in the treatment of moderate to severe chronic plaque psoriasis: results of a randomized, placebo-controlled, phase 2 trial. Arch Dermatol. 2008;144:200-207.

44. Mannon PJ, Fuss IU, Mayer L, et al. Anti-interleukin-12 antibody for active Crohn’s disease. N Engl J Med. 2004;351:2069-2079.

45. Vollmer TL, Wynn DR, Alam MS, Valdes J. A phase 2, 24-week, randomized, placebo-controlled, double-blind study examining the efficacy and safety of an anti-interleukin-12 and -23 monoclonal antibody in patients with relapsing-remitting or secondary progressive multiple sclerosis. Mult Scler. 2011;17:181-191.

46. Landells I, Marano C, Hsu MC, et al. Ustekinumab in adolescent patients age 12 to 17 years with moderate-to-severe plaque psoriasis: results of the randomized phase 3 CADMUS study. J Am Acad Dermatol. 2015;73:594-603.

47. Lebwohl M, Strober B, Menter A, et al. Phase 3 studies comparing brodalumab with ustekinumab in psoriasis. N Engl J Med. 2015;373:1318-1328.

48. Zhu X, Zheng M, Song M, et al. Efficacy and safety of ustekinumab in Chinese patients with moderate to severe plaque-type psoriasis: results from a phase 3 clinical trial (LOTUS). J Drugs Dermatol. 2013;12:166-174.

49. Tsai TF, Ho JC, Song M, et al. Efficacy and safety of ustekinumab for the treatment of moderate-to-severe psoriasis: a phase III, randomized, placebo-controlled trial in Taiwanese and Korean patients (PEARL). J Dermatol Sci. 2011;63:154-163.

50. Leonardi CL, Kimball AB, Papp KA, et al. Efficacy and safety of ustekinumab, a human interleukin-12/23 monoclonal antibody, in patients with psoriasis: 76-week results from a randomised, double-blind, placebo-controlled trial (PHOENIX 1). Lancet. 2008;371:1665-1674.

51. Krueger GG, Langley RG, Leonardi C, et al. A human interleukin-12/23 monoclonal antibody for the treatment of psoriasis. N Engl J Med. 2007;356:580-592.

52. Mclnnes IB, Kavanaugh A, Gottlieb AB, et al. Efficacy and safety of ustekinumab in patients with active psoriatic arthritis: 1 year results of the phase 3, multicentre, double-blind, placebo-controlled PSUMMIT 1 trial. Lancet. 2013;382:780-789.

53. Gottlieb A, Menter A, Mendelson A, et al. Ustekinumab, a human interleukin 12/23 monoclonal antibody, for psoriatic arthritis: randomised, double-blind, placebo-controlled, crossover trial. Lancet. 2009;373:633-640.

54. Feagan BG, Sandborn WJ, Gasink C, et al. Ustekinumab as induction and maintenance therapy for Crohn’s disease. N Engl J Med. 2016;375:1946-1960.

55. Sandborn WJ, Feagan BG, Fedorak RN, et al. A randomized trial of ustekinumab, a human interleukin-12/23 monoclonal antibody, in patients with moderate-to-severe Crohn’s disease. Gastroenterology. 2008;135:1130-1141.

56. Khattri S, Brunner PM, Gar cet S, et al. Efficacy and safety of ustekinumab treatment in adults with moderate-to-severe atopic dermatitis. Exp Dermatol. 2017;26:28-35.

57. Saeki H, Kabashima K, Tokura Y, et al. Efficacy and safety of ustekinumab in Japanese patients with severe atopic dermatitis: a randomized, double-blind, placebo-controlled, phase II study. Br J Dermatol. 2017;177:419-427.

58. van Vollenhoven RF, Hahn BH, Tsokos GC, et al. Efficacy and safety of ustekinumab, an IL-12 and IL-23 inhibitor, in patients with active systemic lupus erythematosus: results of a multicentre, double-blind, phase 2, randomised, controlled study. Lancet. 2018;392:1330-1339.

59. Segal BM, Constantinescu CS, Raychaudhuri A, et al. Repeated subcutaneous injections of IL12/23 p40 neutralising antibody, ustekinumab, in patients with relapsing-remitting multiple sclerosis: a phase II, double-blind, placebo-controlled, randomised, dose-ranging study. Lancet Neurol. 2008;7:796-804.

60. Bissonnette R, Nigen S, Langley RG, et al. Increased expression of IL-17A and limited involvement of IL-23 in patients with palmo-plantar (PP) pustular psoriasis or PP pustulosis; results from a randomised controlled trial. J Eur Acad Dermatol Venereol. 2014;28:1298-1305.

61. Igarashi A, Kato T, Kato M, et al. Efficacy and safety of ustekinumab in Japanese patients with moderate-to-severe plaque-type psoriasis: long-term results from a phase 2/3 clinical trial. J Dermatol. 2012;39:242-252.

62. Ritchlin C, Rahman P, Kavanaugh A, et al. Efficacy and safety of the anti-IL-12/23 p40 monoclonal antibody, ustekinumab, in patients with active psoriatic arthritis despite conventional non-biological and biological anti-tumour necrosis factor therapy: 6-month and 1-year results of the phase 3, multicentre, double-blind, placebo-controlled, randomised PSUMMIT 2 trial. Ann Rheum Dis. 2014;73:990-999.

63. Judson MA, Baughman RP, Costabel U, et al. Safety and efficacy of ustekinumab or golimumab in patients with chronic sarcoidosis. Eur Respir J. 2014;44:1296-1307.

64. Baeten D, Ostergaard M, Wei J, et al. Risankizumab, an IL-23 inhibitor, for ankylosing spondylitis: results of a randomised, double-blind, placebo-controlled, proof-of-concept, dose-finding phase 2 study. Ann Rheum Dis. 2018;77:1295-1302.

65. Lopez A, Mariette X, Bachelez H, et al. Vaccination recommendations for the adult immunosuppressed patient: a systematic review and comprehensive field synopsis. J Autoimmun. 2017;80:10-27.

66. Kawamoto H, Hara H, Minagawa S, et al. Interstitial pneumonia in psoriasis. Mayo Clin Proc Innov Qual Outcomes. 2018;2:370-377.

67. Mathai SC, Danoff SK. Management of interstitial lung disease associated with connective tissue disease. BMJ. 2016;352:h6819.

68. Black H, Mendoza M, Murin S. Thoracic manifestations of inflammatory bowel disease. Chest. 2007;131:524-532.

69. Faure E, Poissy J, Goffard A, et al. Distinct immune response in patients with active systemic lupus erythematosus: results of sparse data. Stat Med. 2004;23:1351-1375.