**Name of Journal:** *World Journal of Clinical Cases*

**Manuscript NO:** 74609

**Manuscript Type:** META-ANALYSIS

**Efficacy and safety of adalimumab in comparison to infliximab for Crohn's disease: A systematic review and meta-analysis**

Yang HH *et al*. ADA *vs* IFX for CD

Hua-Hua Yang, Yi Huang, Xu-Chun Zhou, Ruo-Nan Wang

**Hua-Hua Yang, Yi Huang, Xu-Chun Zhou,** Department of Gastroenterology, The First Affiliated Hospital of Chongqing Medical University, Chongqing 400016, China

**Ruo-Nan Wang,** Department of Endocrinology, Liyuan Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan 430077, Hubei Province, China

**Author contributions:** Yang HH and Zhou XC contributed to study design; Yang HH, Huang Y, and Wang RN contributed to the literature search; Yang HH, Huang Y, and Zhou XC participated in data extraction and analysis; Yang HH and Huang Y wrote the paper.

**Corresponding author: Xu-Chun Zhou, MD, Professor,** Department of Gastroenterology, The First Affiliated Hospital of Chongqing Medical University, No. 1 Youyi Road, Yuzhong District, Chongqing 400016, China. chqxchzh@163.com

**Received:** December 29, 2021

**Revised:** February 14, 2022

**Accepted:** April 30, 2022

**Published online:** June 26, 2022

**Abstract**

BACKGROUND

Adalimumab (ADA) and infliximab (IFX) are the cornerstones of the treatment of Crohn’s disease (CD). It remains controversial whether there is a difference in the effectiveness and safety between IFX and ADA for CD.

AIM

To perform a meta-analysis to compare the effectiveness and safety of ADA and IFX in CD.

METHODS

PubMed, Embase, Cochrane Library, and Web of Science databases were searched. Cohort studies were considered for inclusion. The primary outcomes were induction of response and remission, maintenance of response and remission, and secondary loss of response. Adverse events were secondary outcomes.

RESULTS

Fourteen cohort studies were included. There was no apparent difference between the two agents in the induction response [odds ratio (OR): 1.27, 95% confidence interval (CI): 0.93-1.74, *P* = 0.14] and remission (OR: 1.11, 95%CI: 0.78–1.57, *P* = 0.57), maintenance response (OR: 1.08, 95%CI: 0.76–1.53, *P* = 0.67) and remission (OR: 1.26, 95%CI: 0.87–1.82, *P* = 0.22), and secondary loss of response (OR: 1.01, 95%CI: 0.65–1.55, *P* = 0.97). Subgroup analysis revealed ADA and IFX had similar rates of response, remission, and loss of response either in anti-tumor necrosis factor-α naïve or non-naïve patients. Further, there was a similar result regardless of whether CD patients were treated with optimized therapy, including dose intensification, shortening interval, and combination immunomodulators. However, ADA had a fewer overall adverse events than IFX (OR: 0.62, 95%CI: 0.42–0.91, *P* = 0.02).

CONCLUSION

ADA and IFX have similar clinical benefits for anti-tumor necrosis factor-α naïve or non-naïve CD patients. Overall adverse events rate is higher in patients in the IFX group.

**Key Words:** Crohn disease; Adalimumab; Infliximab; Clinical efficacy; Adverse effects; Meta-analysis

**©The** **Author(s) 2022.** Published by Baishideng Publishing Group Inc. All rights reserved.

**Citation:** Yang HH, Huang Y, Zhou XC, Wang RN. Efficacy and safety of adalimumab in comparison to infliximab for Crohn's disease: A systematic review and meta-analysis. *World J Clin Cases* 2022; 10(18): 6091-6104

**URL:** https://www.wjgnet.com/2307-8960/full/v10/i18/6091.htm

**DOI:** https://dx.doi.org/10.12998/wjcc.v10.i18.6091

**Core Tip:** Differences in immunogenicity and route of administration among adalimumab (ADA) and infliximab (IFX) allow for potential variability in therapeutic properties and efficacy. However, clear recommendations have been limited due to a lack of head-to-head comparison. We conducted a meta-analysis to synthesize current results and compared the efficacy and safety of ADA and IFX. The results showed that both have similar clinical benefits for anti-tumor necrosis factor-α naïve or non-naïve Crohn’s disease patients. Overall adverse events rate is higher in patients in the IFX group. ADA and IFX can be selected based on a possible history of adverse events and patient compliance.

**INTRODUCTION**

Crohn's disease (CD) is an incurable chronic progressive condition characterized by abdominal pain, diarrhea, and weight loss. Aminosalicylic acid preparations, glucocorticoids, immunosuppressants, and biological agents have been used for treatment. Of these, biological agents are most widely used, especially anti-tumor necrosis factor-α (anti-TNF-α) blockers, including infliximab (IFX) and adalimumab (ADA). They all have been proven effective in inducing and maintaining remission and are routinely used in the treatment of CD[1,2]. We do not know, however, which treatment should be considered the priority?

IFX, a chimeric monoclonal antibody against TNF-α, is the first approved anti-TNF-α for moderate to severe CD. ADA is a humanized monoclonal antibody against TNF-α. IFX is given by intravenous infusion every 8 wk, whereas ADA is administered subcutaneously every 4 wk. Differences in immunogenicity and route of administration among them allow for potential variability in therapeutic properties and efficacy. However, clear recommendations have been limited due to a lack of head-to-head treatment comparison. A network meta-analysis published in 2014 found that ADA may be the most efficacious agent for maintenance of remission in CD in biologic-naïve patients[3], while many new clinical practice experience studies have shown their effectiveness and safety data were comparable. Furthermore, even though there has been cumulative research, few studies have focused on secondary loss of response, anti-TNF naïve or non-naïve patients, and the benefits of treatment optimization, such as dose intensification, shortening interval, and combination with immunomodulators. We performed a meta-analysis to synthesize these results and compared the efficacy and safety of ADA and IFX.

**MATERIALS AND METHODS**

Our protocol was registered with PROSPERO (CRD: 42021191655). We followed the Preferred Reporting Items for the Systemic Review and Meta-Analysis guidelines.

***Search*** ***retrieval***

We performed literature search of electronic sources, including PubMed, Cochrane Library, Web of Science, and Embase, from initiation until October 31, 2020. No language restrictions were applied. The search terms included “Crohn disease,” “adalimumab”, and “infliximab” as Medical Subject Headings terms and their entry terms (Crohn disease: Crohn\*; ileitis. Adalimumab: Humira; Exemptia. Infliximab: Remicade) to improve search outcomes. We also screened references of relevant articles to avoid omissions.

***Inclusion and exclusion criteria***

We included cohort studies comparing ADA and IFX for treating adults with CD. Comparisons of induction of remission and response rates, maintenance of remission and response rates, secondary loss of response rates, and the incidence of adverse events were among the outcomes of included studies. Excluded studies included those conducted in the pediatric population, those that did not investigate patients with inflammatory bowel disease, and those that did not report any outcomes of interest.

***Study selection***

Two investigators (Yang HH and Huang Y) independently screened the titles, abstracts, and full texts of all papers to determine trial eligibility for inclusion. Investigators used a consensual approach to determine the inclusion or exclusion of selected studies after full-text assessment. Any disagreement was resolved through discussion or with a third researcher. The study characteristics were extracted independently by two authors using a standardized datasheet.

***Data extraction***

We collected the following variables: First author’s name, year of publication, country or area, study design, number of patients, gender, median age, Montreal classification, duration of follow-up, previous treatment, and outcomes of interest. The endpoint of this meta-analysis mainly included the induction response and remission, maintenance response and remission, overall adverse events rate, severe adverse events rate, and the rate of opportunistic infections.

The outcomes of interest included: (1) Induction of clinical remission defined as Crohn’s disease activity index (CDAI) < 150, Harvey Bradshaw Index (HBI) ≤ 4, or by physician's global assessment after ≤ 14 wk; (2) Induction of clinical response was defined as ΔCDAI ≥ 70, ΔHBI ≥ 2, or by physician's global assessment after ≤ 14 wk; (3) Maintenance of remission referred to clinical remission after ≤ 54 wk; (4) Maintenance of response referred to clinical response after ≤ 54 wk; (5) Secondary loss of response was defined as a reappearance of disease activity after achieving induction response, coupled with the need to change treatment, including dose intensification, the addition of an immunomodulator, or need to discontinue treatment; and (6) Secondary outcomes included a comparison of the incidence of overall adverse events, severe adverse events, and opportunistic infections in trials of maintenance therapy.

***Quality assessment***

One author assessed the quality of included studies through the Newcastle-Ottawa Quality Assessment Scale (NOS). High-quality studies were defined by a total score of ≥ 6.

***Statistical analysis***

RevMan 5.3 and Stata 16.0 software were used for statistical analysis. Odds ratio (OR) and concomitant 95% confidence interval (CI) were evaluated for the quantitative analyses. The random-effect model was used. Heterogeneity was explored by calculating *I*2 and employing the Q test. An *I*2 estimate > 50% and a *P* < 0.05 were regarded markers of significant heterogeneity, and its causes were investigated. We performed sensitivity analyses and subgroup analyses to detect the source of heterogeneity. *P* < 0.05 was considered to indicate a significant difference. Subgroup analyses were conducted based on the following grouping criteria: (1) Studies evaluating outcomes on anti-TNF naïve patients *vs* studies on non-naïve patients; (2) Studies evaluating outcomes on more perianal diseases in IFX group *vs* equal perianal disease in IFX and ADA group; (3) Studies evaluating primary outcomes given with treatment optimization, *i.e.* shortening the administration intervals, increasing the dose, and/or combination with immunomodulator therapy; and (4) Studies evaluating secondary outcomes at ≤ 48 wk *vs* > 48 wk. Funnel plots and Egger’s test was used to test for publication bias.

**RESULTS**

***Literature search***

A preliminary search of the above database identified 2228 documents. Of these, we removed 562 duplicates, discarded 1632 studies after screening the titles and abstracts, and assessed the full text of 34 studies for eligibility. Finally, 14 cohort studies were included,and 20 were excluded. The flow diagram describes this process in detail (Figure 1).

***Study characteristics***

Study design, outcomes, the definition of outcomes, inclusion criteria, and follow-up time differed among the included studies. Our meta-analysis consisted of two prospective cohort studies and 12 retrospective cohort studies. Three pieces of research evaluated maintenance response or remission at 54 wk[4-6], five at 48 wk[7-11], and one at 26 wk[12]. Regarding the definition of outcomes, most incorporated studies evaluated clinical response or remission by CDAI or HBI except for the study by Macaluso *et al*[10]. In addition, seven studies only included anti-TNF-naïve patients[4,7-9,12,13], and no study only included patients who failed anti-TNF treatment. Follow-up intervals across studies varied, ranging from 4 to 14 wk for induction period and 26 to 168 wk for maintenance period. The high NOS scores reflected the high quality of the enrolled studies. Thirteen studies got a score of ≥ 6, except for the study by Bau *et al*[14], which scored 5. Table 1 showed the overall characteristics of the selected studies.

***Primary outcomes***

**Induction of response:** Fivestudies (1040 patients) recorded induction of response[6,7,9,10,15]. No difference was shown between groups in response rates (OR: 1.27, 95%CI: 0.93–1.74, *P* = 0.14). Of the 1040 patients in five studies, 515 received ADA therapy. The heterogeneity of those studies was insignificant (*P* = 0.58, *I*2 = 0%) (Figure 2A). Sensitivity analysis showed no significant changes to the exclusion of any one of the studies (Supplementary Table 1). Subgroup analysis revealed no remarkable difference between groups (Table 2).

**Induction of remission:** When combining all four studies[6,8,9,16] reporting induction of remission data (318 on ADA therapy and 494 on IFX therapy), we found no difference between the two groups of patients (OR: 1.11, 95%CI: 0.78–1.57, *P* = 0.57). Heterogeneity was low (*P* = 0.85, *I*2 = 0%) (Figure 2B). Subsequent subgroup analysis showed similar results (Table 2). In sensitivity analyses, excluding any one of the studies did not significantly impact the results (Supplementary Table 1).

**Maintenance of response:** Of the 14 studies, seven reported the response rate in maintenance therapy[4,6,7,9-12]. A number of 1828 patients were included: 896 IFX-treated *vs* 932 ADA-treated. Data analysis showed that ADA and IFX had a similar rate of maintenance of response (OR: 1.08, 95%CI: 0.76–1.53, *P* = 0.67). Heterogeneity was significant (*P* = 0.03, *I*2 = 56%) (Figure 3A). Cosnes *et al*[12] evaluating response at 26 wk increased heterogeneity. In the sensitivity analysis, the result remained unchanged with the exclusion of any study (Supplementary Table 1). Subgroup analyses also showed no difference between the two groups (Table 2).

**Maintenance of remission:** There were 770 patients (328 on ADA therapy) available for analysis from six studies[5-9,11]. Data analysis showed that ADA and IFX had a similar rate of maintenance of remission (OR: 1.26, 95%CI: 0.87–1.82, *P* = 0.22). Heterogeneity was low (*P* = 0.29, *I*2 = 19%) (Figure 3B). Subgroup analyses also showed no statistical differences (Table 2). Sensitivity analysis indicated that the results were stable (Supplementary Table 1).

**Secondary loss of response:** Six studies with 1307 patients were included (603 receiving ADA and 704 IFX therapy)[5-7,9,12,17].There was no statistical difference between the two treatments (OR: 1.01, 95%CI: 0.65–1.55, *P* = 0.97). Heterogeneity was notable (*P* = 0.05, *I*2 = 54%) (Figure 4). Heterogeneity was linked to the study by Narula *et al*[9], which found that IFX had more rate of loss of response than ADA. On sensitivity analyses, the results remained the same after excluding any one study (Supplementary Table 1). There was also no significant difference between ADA and IFX when subgroup analysis was done (Table 2).

***Secondary outcomes***

**Overall adverse events:** The incidence of overall adverse events was recorded in a total of eight cohort studies[4,5,7-11,14] that included 1653 patients, of which ADA was less than IFX (OR: 0.62, 95%CI: 0.42–0.91, *P* = 0.02). There was high heterogeneity (*P* = 0.04, *I*2 = 53%) (Figure 5A). Subgroup analysis revealed that ADA had fewer overall adverse events than IFX in ≤ 48 wk follow-up time (OR: 0.50, 95%CI: 0.33–0.76, *P* = 0.001); and in anti-TNF-α-naïve patients, IFX had more adverse events (OR: 0.67, 95%CI: 0.50–0.89, *P* = 0.005) (Table 2). Sensitivity analysis indicated that the results were slightly unstable (Supplementary Table 1).

**Severe adverse events:** Our analysis of seven studies[6,8,9,11,12,14,15] with a total of 1547 patients showed ADA had a similar rate of severe adverse events with IFX (OR: 0.75, 95%CI: 0.32–1.72, *P* = 0.49). Sensitivity analysis was performed due to notable heterogeneity (*P* = 0.003, *I*2 = 72%) (Figure 5B). Heterogeneity mainly originated from Zorzi *et al*[6] with more severe adverse events occurring in IFX therapy. The result remained unchanged with the exclusion of any study (Supplementary Table 1). Subgroup analysis also showed similar results (Table 2).

**Opportunistic infections:** Sixstudies[4,7,9,13-15] reported side effects, with a total number of 1910 cases (ADA: IFX = 922:988). Opportunistic infections rates in the IFX and ADA groups were similar (OR: 0.96, 95%CI: 0.66-1.40, *P* = 0.83), and no apparent heterogeneity was detected (Figure 5C). There was no significant difference when subgroup analysis was done (Table 2). Sensitivity analysis showed no significant changes when any one of the studies was excluded (Supplementary Table 1).

***Publication bias and GRADE evaluation***

The symmetry of the funnel plot indicated there was no publication bias (Figure 6). The Egger’s test showed no significant publication bias for maintenance of response (*P* = 0.7024 > 0.05), maintenance of remission (*P* = 0.1003 > 0.05), secondary loss of response (*P* = 0.0510 > 0.05), and overall adverse events (*P* = 0.6717 > 0.05). GRADE evidence of all outcomes was judged as “low”. The results are shown in Table 3.

**DISCUSSION**

The immunogenicity of anti-TNF-α agents triggered the formation of anti-drug antibodies (ADAbs) specific to the agent administered. ADAbs of IFX or ADA and reduced serum concentrations in association with ADAbs together lead to decreased clinical benefit and increased adverse events. Although the immunogenicity of IFX is usually higher than that for ADA, we found both of them have similar response characteristics in CD patients. In our meta-analyses, no significant differences in primary outcomes were found between groups treated with IFX and ADA. These results were consistent with the results of most published studies[5-12,15-17]. One unexpected finding was the extent to which the overall adverse events rate of IFX was higher than that of ADA. Our meta-analysis indicated that physicians may choose on an individual basis, according to a possible history of adverse events to either IFX or ADA and to patient compliance, to give either an intravenous infusion or a self-administered subcutaneous injection.

CD is a heterogeneous disease, and the therapeutic efficacy differs between the types of disease, *e.g.*,location of disease, the existence of stenosis and/or fistula, or perianal involvement. There was no significant difference between IFX and ADA groups in the location of disease and existence of stenosis and/or fistula of included studies. However, IFX patients had more perianal diseases in the studies of Benmassaoud *et al*[7], Varma *et al*[8], Narula *et al*[9], and Cosnes *et al*[12]. Clinicians tended to choose IFX over ADA in patients with more severe disease activity or phenotypes (perianal disease) due to its intravenous administration and weight-based dosing schedule. We attempted to adjust for these differences through subgroup analysis, which led to the same conclusions (Supplementary Tables 2 and 3). Additionally, Ji *et al*[18] found the cumulative rate of nonrecurrence or aggravation of fistula at 24 mo was not significantly different between IFX and ADA groups (62.5% *vs* 83.9%, *P* = 0.09). Current evidence suggested that IFX and ADA had similar effects in patients with perianal disease.

Biologic-naïve or non-naïve patients were important factors to influence the results. It is controversial whether ADA had similar efficacy to IFX in previous anti-TNF exposure CD patients. Macaluso *et al*[10] compared clinical benefits between IFX and ADA only in biologic non-naïve CD patients and reported that there was no difference in clinical benefits at 12 wk and after 1 year (*P* = 0.600 and *P* = 0.620, respectively). A retrospective case-control study[19] found that the risk for ADAbs to IFX was higher than ADAbs to ADA when patients had prior antibodies to anti-TNF. They did not investigate clinical efficacy. However, Sasson and Ananthakrishnan[20] found that patients with high ADAbs titers exhibited similar rates of clinical efficacy to ADA therapy compared to those with low titers (at 3 mo and 12 mo *P* = 0.81 and 0.62 respectively). This may mean IFX and ADA have similar efficacy in previous anti-TNF exposed CD patients. Our findings indicated that either in naïve or non-naïve patients ADA and IFX had similar clinical response and remission. More studies conducted on previous anti-TNF exposure CD patients will be necessary.

Co-immunosuppression affected the results of the analysis. The finding that combination therapy with an immunomodulator is superior with IFX but not with ADA was reported in Kestens *et al*[4], Benmassaoud *et al*[7], and Doecke *et al*[16]. The possible reason is that IFX combined with immunomodulator treatment reduces its immunogenicity. However, clinical efficacy of ADA combination therapy did not differ from that of ADA monotherapy (71.8% *vs* 68.1% at week 26, *P* = 0.63)[21]. Therefore, more patients in the IFX group were combined with immunomodulator treatment than in the ADA group in the Narula *et al*[9] study. No change was found in results after sensitivity analysis was conducted. Patients were on concomitant immunomodulation at anti-TNF induction to improve the efficacy of the induction of the remission and discontinued co-therapy due to adverse effects or intolerability (from the beginning). When loss of response occurred, concomitant therapy was resumed (later add on). Only the Cosnes *et al*[12] study used immunomodulators later. No different results were found after sensitivity analysis was performed. Furthermore, CD patients who lost response were allowed to shorten intervals and double dosage. These optimization strategies also impacted the results. We conducted subgroup analyses comparing the outcomes between using dose optimization and not and found the clinical effect of ADA was similar to IFX.

Similar to the findings of many studies[4,10,17], the significantly higher rate of overall adverse events can be seen in patients using IFX, which could be attributed to infusion or allergic reactions. Benmassaoud *et al*[7] reported that IFX group patients were more likely to have infusion or injection reactions than ADA. A higher rate of allergic reactions in the IFX was observed in a study by Narula *et al*[9]. However, we noted that the difference did not exist in anti-TNF-α non-naïve patients and with long follow-up time. We were unable to evaluate long-term safety due to the different follow-up times of each study. Larger and long-term comparison studies will be necessary. In addition, the instability of the results also require further studies to establish these findings.

Additionally, we failed to evaluate long-term results due to the different follow-up times of each study. Inokuchi *et al*[22] performed a retrospective study to evaluate long-term prognosis. They observed that the rates of cumulative steroid-free remission rates and surgery-free did not differ significantly between the two groups after a median observation period of 64.2 mo (*P* = 0.42 and *P* = 0.74, respectively). The goal of CD treatment requires more than clinical healing. Mucosal healing and tissue healing are expected to stop disease progression and reduce recurrence. Tursi *et al*[15] found that mucosal healing and histological healing were comparable between the two groups (*P* = 0.946 and *P* = 0.895, respectively).

Although biologic agents targeting TNF-α have achieved remarkable progress in treating CD, some patients do not respond to the induction therapy or lose response over time (secondary loss of response). The anti-drug antibodies or low serum drug concentrations play a critical part in the loss of response[23]. If ADA is superior to IFX for remission, ADA should have a lower rate of secondary loss of response than IFX. However, we failed to find a difference in the secondary loss of response between the two groups, which contradicted our hypothesis. It was further demonstrated that both have similar effects.

This work is the first direct comparison meta-analysis to evaluate the comparative effectiveness and safety of ADA and IFX in CD. Previous network meta-analyses addressed similar outcomes in the Bayesian setting indirect comparison. In our study, we enrolled comparative trial data resulting in more credible results. Furthermore, head-to-head clinical trials comparing ADA and IFX would not be feasible in the future; therefore, our studies will help guide optimal therapies.

Our current study has some limitations. First, we only included observational studies and failed to control adequately confounders, such as disease severity, disease phenotype, steroid use, *etc*. In addition to clinical benefits, we should consider other factors, such as patients’ preferences and costs. Future studies are needed to address these questions.

**CONCLUSION**

IFX and ADA have similar response characteristics either in anti-TNF naïve and non-naïve CD patients, and ADA therapy has fewer overall adverse events. Our study indicates that IFX or ADA can be freely chosen as treatment based on physician and patient agreement. Eventually, the decision of which treatment to start may depend on factors such as patient preference and cost.

**ARTICLE HIGHLIGHTS**

***Research background***

Infliximab (IFX) is often selected as the first-line anti-tumor necrosis factor-α (TNF-α) agent for Crohn’s disease (CD), despite the lack of data showing its superiority over adalimumab (ADA).

***Research motivation***

By comparing the effectiveness and safety between ADA and IFX, we wanted to determine if IFX or ADA is superior to the other for treatment of CD.

***Research objectives***

The present meta-analysis was performed to evaluate the comparative effectiveness and safety of ADA and IFX for CD to assist clinicians in making treatment choices.

***Research methods***

The clinical studies that compared the effectiveness or safety of ADA and IFX in the treatment of CD were searched in PubMed, Embase, Cochrane Library, and Web of Science databases.

***Research results***

Our meta-analysis of CD patients who were naïve or non-naïve to anti-TNF-α agents found no significant differences between IFX and ADA on many measures of effectiveness, including clinical response, clinical remission, and secondary loss of response. Interestingly, we observed a higher rate of overall adverse events in patients using IFX compared to ADA.

***Research conclusions***

IFX and ADA are comparable in clinical outcomes for patients with CD who are naïve or non-naïve to anti-TNF-α antagonists. However, fewer overall adverse events are noted in ADA patients.

***Research perspectives***

Our study provide reassurance to clinicians by synthesizing current literature suggesting that the ADA and IFX have similar effectiveness in “real-world” use. Larger, long-term, and prospective head-to-head comparison studies will be necessary to confirm these results. More research also will be necessary to explore the cost of anti-TNF-α agents.

**ACKNOWLEDGEMENTS**

We would like to thank all authors of the included primary studies.

**REFERENCES**

1 **Cushing K**, Higgins PDR. Management of Crohn Disease: A Review. *JAMA* 2021; **325**: 69-80 [PMID: 33399844 DOI: 10.1001/jama.2020.18936]

2 **Danese S**, Vuitton L, Peyrin-Biroulet L. Biologic agents for IBD: practical insights. *Nat Rev Gastroenterol Hepatol* 2015; **12**: 537-545 [PMID: 26284562 DOI: 10.1038/nrgastro.2015.135]

3 **Singh S**, Garg SK, Pardi DS, Wang Z, Murad MH, Loftus EV Jr. Comparative efficacy of biologic therapy in biologic-naïve patients with Crohn disease: a systematic review and network meta-analysis. *Mayo Clin Proc* 2014; **89**: 1621-1635 [PMID: 25441399 DOI: 10.1016/j.mayocp.2014.08.019]

4 **Kestens C**, van Oijen MG, Mulder CL, van Bodegraven AA, Dijkstra G, de Jong D, Ponsioen C, van Tuyl BA, Siersema PD, Fidder HH, Oldenburg B; Dutch Initiative on Crohn and Colitis (ICC). Adalimumab and infliximab are equally effective for Crohn's disease in patients not previously treated with anti-tumor necrosis factor-α agents. *Clin Gastroenterol Hepatol* 2013; **11**: 826-831 [PMID: 23376000 DOI: 10.1016/j.cgh.2013.01.012]

5 **Otake H**, Matsumoto S, Mashima H. Does long-term efficacy differ between infliximab and adalimumab after 1 year of continuous administration?: A STROBE-compliant retrospective cohort study. *Medicine (Baltimore)* 2017; **96**: e6635 [PMID: 28422861 DOI: 10.1097/MD.0000000000006635]

6 **Zorzi F**, Zuzzi S, Onali S, Calabrese E, Condino G, Petruzziello C, Ascolani M, Pallone F, Biancone L. Efficacy and safety of infliximab and adalimumab in Crohn's disease: a single centre study. *Aliment Pharmacol Ther* 2012; **35**: 1397-1407 [PMID: 22519466 DOI: 10.1111/j.1365-2036.2012.05100.x]

7 **Benmassaoud A**, Al-Taweel T, Sasson MS, Moza D, Strohl M, Kopylov U, Paradis-Surprenant L, Almaimani M, Bitton A, Afif W, Lakatos PL, Bessissow T. Comparative Effectiveness of Infliximab Versus Adalimumab in Patients with Biologic-Naïve Crohn's Disease. *Dig Dis Sci* 2018; **63**: 1302-1310 [PMID: 29243105 DOI: 10.1007/s10620-017-4874-6]

8 **Varma P**, Paul E, Huang C, Headon B, Sparrow MP. A retrospective comparison of infliximab *vs* adalimumab as induction and maintenance therapy for Crohn disease. *Intern Med J* 2016; **46**: 798-804 [PMID: 26865349 DOI: 10.1111/imj.13040]

9 **Narula N**, Kainz S, Petritsch W, Haas T, Feichtenschlager T, Novacek G, Eser A, Vogelsang H, Reinisch W, Papay P. The efficacy and safety of either infliximab or adalimumab in 362 patients with anti-TNF-α naïve Crohn's disease. *Aliment Pharmacol Ther* 2016; **44**: 170-180 [PMID: 27226407 DOI: 10.1111/apt.13671]

10 **Macaluso FS**, Fries W, Privitera AC, Cappello M, Siringo S, Inserra G, Magnano A, Di Mitri R, Mocciaro F, Belluardo N, Scarpulla G, Magrì G, Trovatello A, Carroccio A, Genova S, Bertolami C, Vassallo R, Romano C, Citrano M, Accomando S, Ventimiglia M, Renna S, Orlando R, Rizzuto G, Porcari S, Ferracane C, Cottone M, Orlando A; Sicilian Network for Inflammatory Bowel Diseases [SN-IBD]. A Propensity Score-matched Comparison of Infliximab and Adalimumab in Tumour Necrosis Factor-α Inhibitor-naïve and Non-naïve Patients With Crohn's Disease: Real-Life Data From the Sicilian Network for Inflammatory Bowel Disease. *J Crohns Colitis* 2019; **13**: 209-217 [PMID: 30295785 DOI: 10.1093/ecco-jcc/jjy156]

11 **Kaniewska M**, Rosołowski M, Rydzewska G. Efficacy, tolerability, and safety of infliximab biosimilar in comparison to originator biologic and adalimumab in patients with Crohn disease. *Pol Arch Intern Med* 2019; **129**: 484-489 [PMID: 31316042 DOI: 10.20452/pamw.14901]

12 **Cosnes J**, Sokol H, Bourrier A, Nion-Larmurier I, Wisniewski A, Landman C, Marteau P, Beaugerie L, Perez K, Seksik P. Adalimumab or infliximab as monotherapy, or in combination with an immunomodulator, in the treatment of Crohn's disease. *Aliment Pharmacol Ther* 2016; **44**: 1102-1113 [PMID: 27666569 DOI: 10.1111/apt.13808]

13 **Di Domenicantonio R**, Trotta F, Cascini S, Agabiti N, Kohn A, Gasbarrini A, Davoli M, Addis A. Population-based cohort study on comparative effectiveness and safety of biologics in inflammatory bowel disease. *Clin Epidemiol* 2018; **10**: 203-213 [PMID: 29440933 DOI: 10.2147/CLEP.S150030]

14 **Bau M**, Zacharias P, Ribeiro DA, Boaron L, Steckert Filho A, Kotze PG. Safety profile of anti-TNF therapy in Crohn's disease management: A Brazilian single-center direct retrospective comparison between infliximab and adalimumab. *Arq Gastroenterol* 2017; **54**: 328-332 [PMID: 28954043 DOI: 10.1590/S0004-2803.201700000-43]

15 **Tursi A**, Elisei W, Picchio M, Penna A, Lecca PG, Forti G, Giorgetti G, Faggiani R, Zampaletta C, Pelecca G, Brandimarte G. Effectiveness and safety of infliximab and adalimumab for ambulatory Crohn's disease patients in primary gastroenterology centres. *Eur J Intern Med* 2014; **25**: 485-490 [PMID: 24631020 DOI: 10.1016/j.ejim.2014.02.010]

16 **Doecke JD**, Hartnell F, Bampton P, Bell S, Mahy G, Grover Z, Lewindon P, Jones LV, Sewell K, Krishnaprasad K, Prosser R, Marr D, Fischer J, R Thomas G, Tehan JV, Ding NS, Cooke SE, Moss K, Sechi A, De Cruz P, Grafton R, Connor SJ, Lawrance IC, Gearry RB, Andrews JM, Radford-Smith GL; Australian and New Zealand Inflammatory Bowel Disease Consortium. Infliximab vs. adalimumab in Crohn's disease: results from 327 patients in an Australian and New Zealand observational cohort study. *Aliment Pharmacol Ther* 2017; **45**: 542-552 [PMID: 27995633 DOI: 10.1111/apt.13880]

17 **Ma C**, Huang V, Fedorak DK, Kroeker KI, Dieleman LA, Halloran BP, Fedorak RN. Crohn's disease outpatients treated with adalimumab have an earlier secondary loss of response and requirement for dose escalation compared to infliximab: a real life cohort study. *J Crohns Colitis* 2014; **8**: 1454-1463 [PMID: 24947334 DOI: 10.1016/j.crohns.2014.05.007]

18 **Ji CC**, Takano S. Clinical efficacy of adalimumab *vs* infliximab and the factors associated with recurrence or aggravation during treatment of anal fistulas in Crohn's disease. *Intest Res* 2017; **15**: 182-186 [PMID: 28522947 DOI: 10.5217/ir.2017.15.2.182]

19 **Vande Casteele N**, Abreu MT, Flier S, Papamichael K, Rieder F, Silverberg MS, Khanna R, Okada L, Yang L, Jain A, Cheifetz AS. Patients With Low Drug Levels or Antibodies to a Prior Anti-Tumor Necrosis Factor Are More Likely to Develop Antibodies to a Subsequent Anti-Tumor Necrosis Factor. *Clin Gastroenterol Hepatol* 2022; **20**: 465-467.e2 [PMID: 33421628 DOI: 10.1016/j.cgh.2021.01.006]

20 **Sasson AN**, Ananthakrishnan AN. High Anti-Infliximab Antibody Titers Do Not Impact Response to Subsequent Adalimumab Treatment in Inflammatory Bowel Diseases. *Dig Dis Sci* 2021 [PMID: 34117949 DOI: 10.1007/s10620-021-07088-x]

21 **Matsumoto T**, Motoya S, Watanabe K, Hisamatsu T, Nakase H, Yoshimura N, Ishida T, Kato S, Nakagawa T, Esaki M, Nagahori M, Matsui T, Naito Y, Kanai T, Suzuki Y, Nojima M, Watanabe M, Hibi T; DIAMOND study group. Adalimumab Monotherapy and a Combination with Azathioprine for Crohn's Disease: A Prospective, Randomized Trial. *J Crohns Colitis* 2016; **10**: 1259-1266 [PMID: 27566367 DOI: 10.1093/ecco-jcc/jjw152]

22 **Inokuchi T**, Takahashi S, Hiraoka S, Toyokawa T, Takagi S, Takemoto K, Miyaike J, Fujimoto T, Higashi R, Morito Y, Nawa T, Suzuki S, Nishimura M, Inoue M, Kato J, Okada H. Long-term outcomes of patients with Crohn's disease who received infliximab or adalimumab as the first-line biologics. *J Gastroenterol Hepatol* 2019; **34**: 1329-1336 [PMID: 30724387 DOI: 10.1111/jgh.14624]

23 **Roda G**, Jharap B, Neeraj N, Colombel JF. Loss of Response to Anti-TNFs: Definition, Epidemiology, and Management. *Clin Transl Gastroenterol* 2016; **7**: e135 [PMID: 26741065 DOI: 10.1038/ctg.2015.63]

**Footnotes**

**Conflict-of-interest statement:** Nothing to disclosed.

**PRISMA 2009 Checklist statement:** The authors have read the PRISMA 2009 Checklist, and the manuscript was prepared and revised according to the PRISMA 2009 Checklist.

**Open-Access:** This article is an open-access article that was selected by an in-house editor and fully peer-reviewed by external reviewers. It is distributed in accordance with the Creative Commons Attribution NonCommercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: https://creativecommons.org/Licenses/by-nc/4.0/

**Provenance and peer review:** Unsolicited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** December 29, 2021

**First decision:** January 25, 2022

**Article in press:** April 30, 2022

**Specialty type:** Medicine, research and experimental

**Country/Territory of origin:** China

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): 0

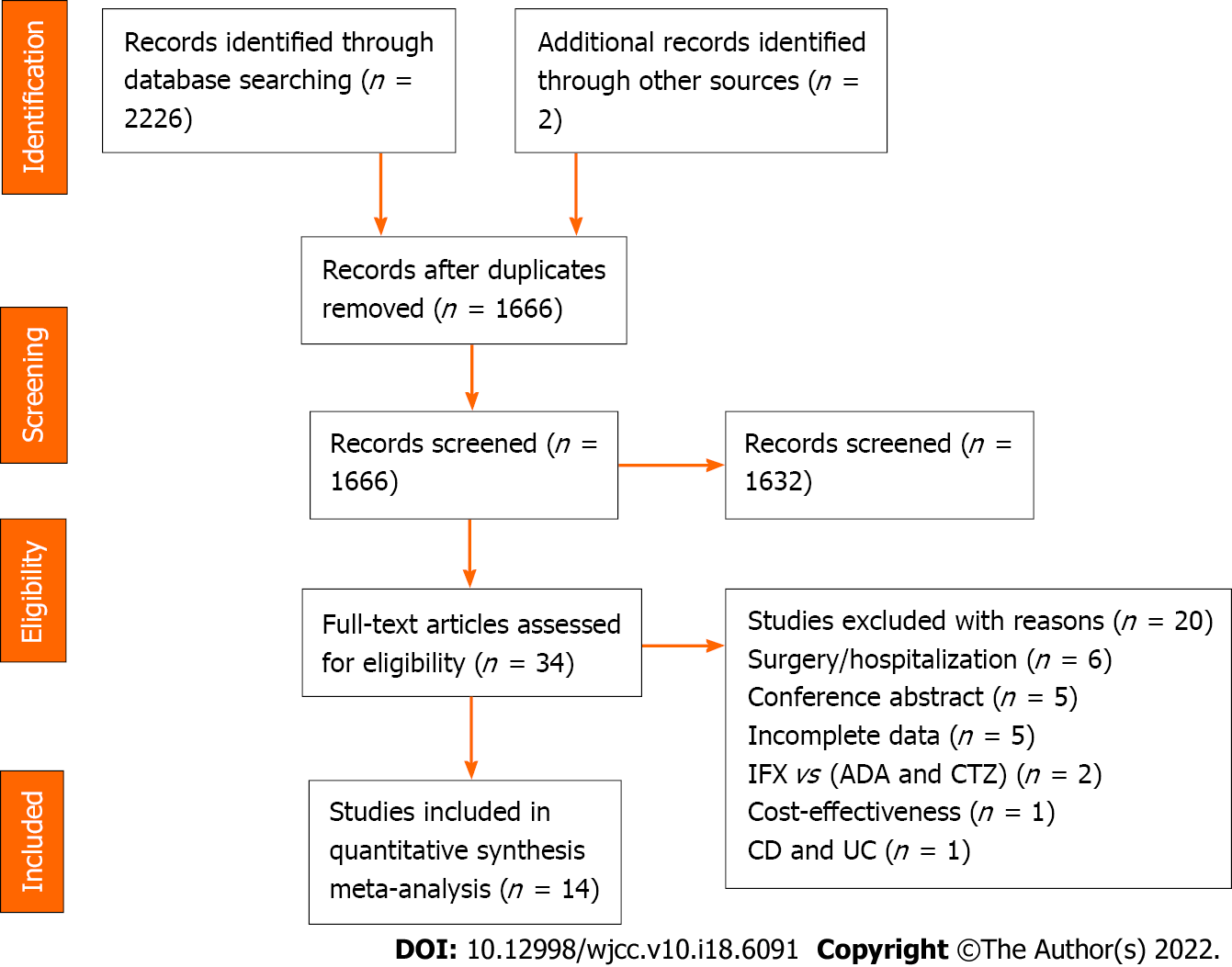
Grade C (Good): C, C

Grade D (Fair): 0

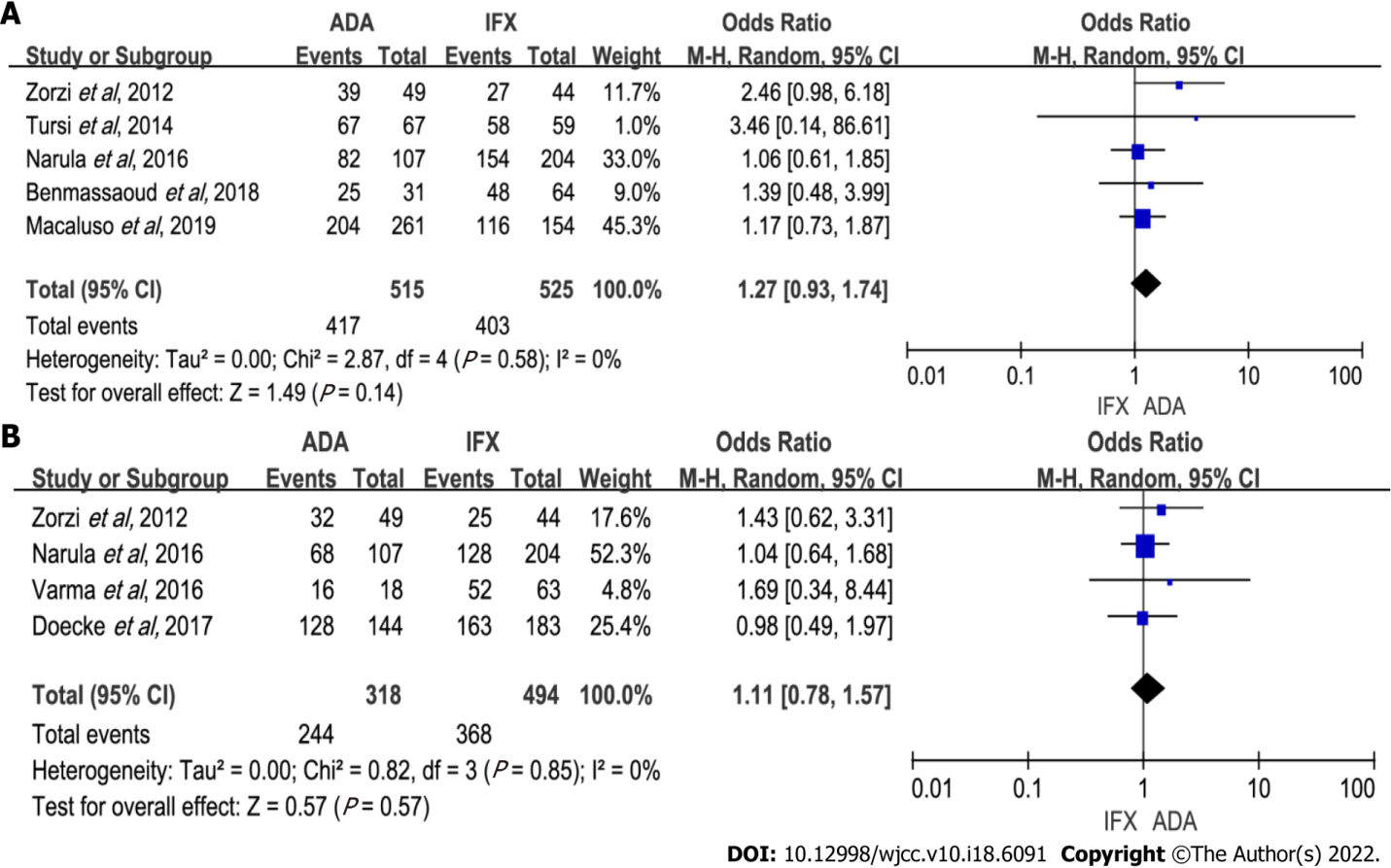
Grade E (Poor): 0

**P-Reviewer:** Fukata M, Japan; Ribeiro IB, Brazil **S-Editor:** Fan JR **L-Editor:** Filipodia **P-Editor:** Fan JR

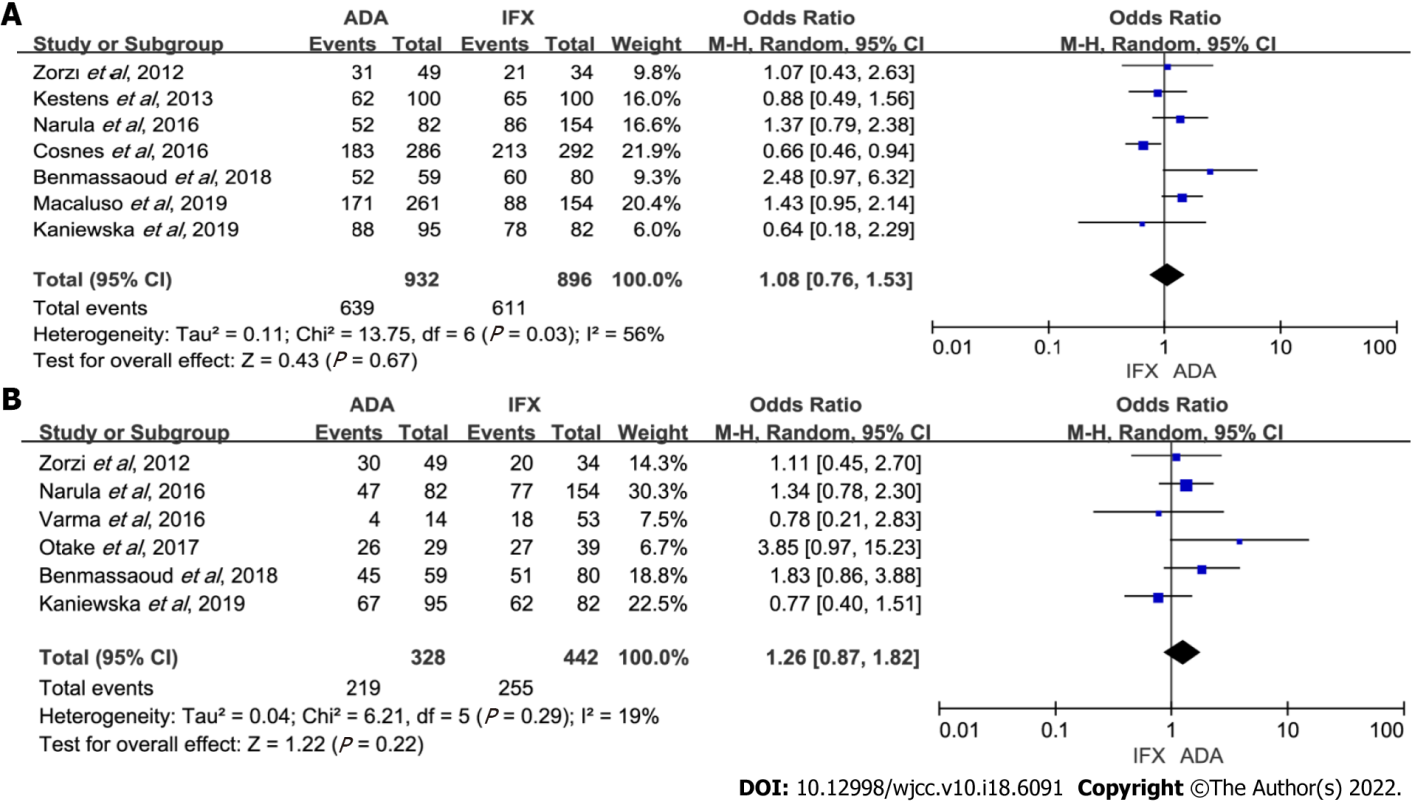
**Figure Legends**



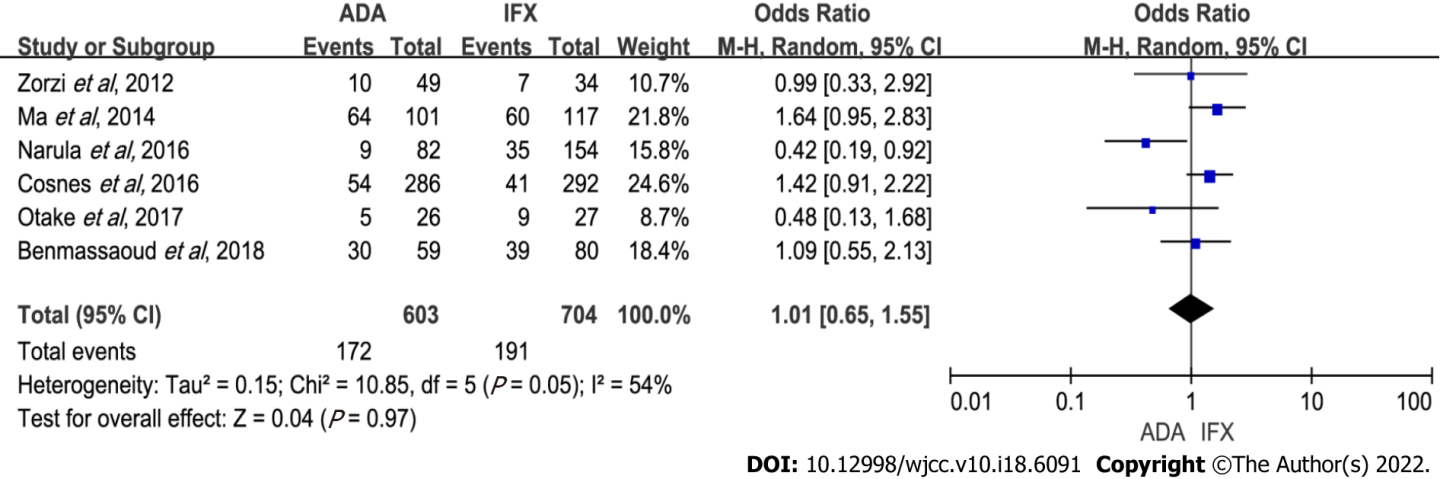
**Figure 1 Flow chart for literature search.** IFX: Infliximab; ADA: Adalimumab; CTZ: Certolizumab; CD: Crohn’s disease; UC: Ulcerative colitis*.*

**

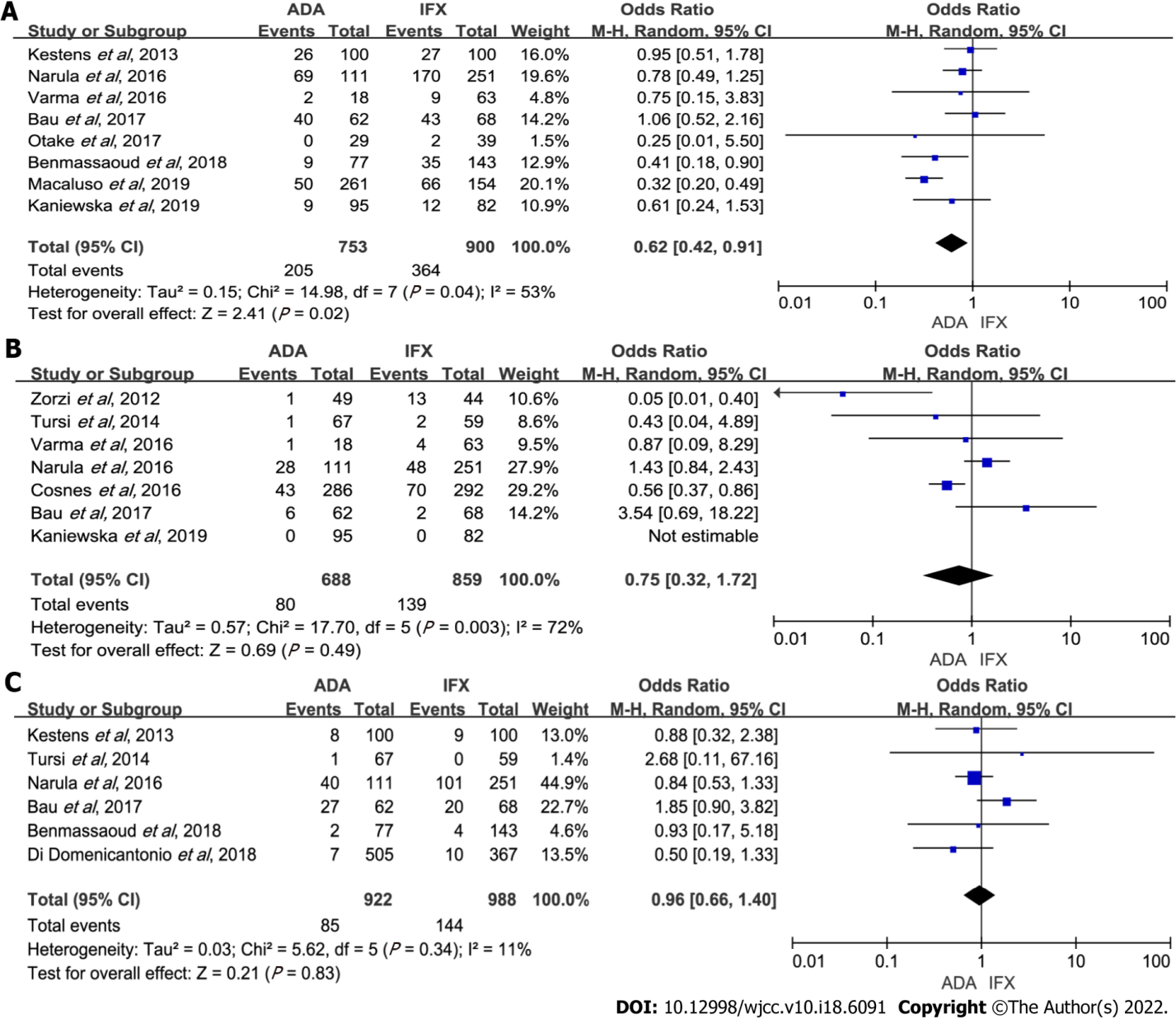
**Figure 2 Forest plot for induction efficacy comparing adalimumab and infliximab.** A: Induction of response; B: Induction of remission. ADA: Adalimumab; IFX: Infliximab; CI: Confidence interval.



**Figure 3 Forest plot for maintenance efficacy comparing adalimumab and infliximab.** A: Maintenance of response; B: Maintenance of remission. ADA: Adalimumab; IFX: Infliximab; CI: Confidence interval.



**Figure 4 Forest plot comparing infliximab and adalimumab for the incidence of secondary loss of response.** ADA: Adalimumab; IFX: Infliximab; CI: Confidence interval.



**Figure 5 Forest plot for comparisons of the rate of adverse events for adalimumab and infliximab.** A: Overall adverse events; B: Severe adverse events; C: Opportunistic infection. ADA: Adalimumab; IFX: Infliximab; CI: Confidence interval.



**Figure 6 Funnel plot.** A: Maintenance of response; B: Maintenance of remission; C: Secondary loss of response; D: Overall adverse events. OR: Odds ratio; SE: Standard error.

**Table 1 Characteristics of selected studies**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Ref.** | **Study design** | **Patient**  **inclusion criteria** | **ADA/IFX, *n*** | **Definition of remission** | **Definition of secondary loss of response** | **Induction of response/remission in wk** | **Maintenance of response/remission in wk** | **Adverse events** | **NOS** |
| Zorzi *et al*[6], 2012 | Retrospective | Active CD | 49/44 | CDAI < 150 | No improvement or worsening | 4/6 | 54 | Multiple | 6 |
| Kestens *et al*[4], 2013 | Retrospective | Naïve CD | 100/100 | NS | NS | NS | 54 | Multiple | 9 |
| Ma *et al*[17], 2014 | Retrospective | Naïve CD | 101/117 | NS | Requiring dose escalation | NS | NS | NS | 8 |
| Tursi *et al*[15], 2014 | Retrospective | CD | 67/59 | HBI ≤ 5 | NS | 6-14 | NS | Multiple | 8 |
| Cosnes *et al*[12], 2016 | Prospective | Naïve CD | 264/127 | CDAI < 150 | Disease activity | NS | 26 | Multiple | 8 |
| Varma *et al*[8], 2016 | Retrospective | Naïve CD | 18/63 | CDAI < 150 | NS | 12 | 48 | Multiple | 7 |
| Narula *et al*[9], 2016 | Prospective | Naïve CD | 111/251 | HBI < 5 | Dose escalation | 12 | 48 | Multiple | 8 |
| Bau *et al*[14], 2017 | Retrospective | Refractory CD | 62/68 | NS | NS | NS | 168 | Multiple | 5 | |
| Otake *et al*[5], 2017 | Retrospective | CD | 29/39 | CDAI < 150 | Multiple | NS | 54 | NS | 8 |
| Doecke *et al*[16], 2017 | Retrospective | CD | 144/183 | CDAI ≤ 150 | NS | 14 | NS | NS | 7 |
| Benmassaoud *et al*[7], 2018 | Retrospective | Naïve CD | 77/143 | HBI ≤ 4 | Need for dose escalation | 12 | 48 | Multiple | 8 |
| Di Domenicantonio *et al*[13], 2018 | Retrospective | Naïve CD | 505/367 | NS | NS | NS | NS | Multiple | 9 |
| Macaluso *et al*[10], 2019 | Retrospective | Naïve and non-naïve CD | Naïve: 214/107; non-naïve: 47/47 | NS | NS | 12 | 48 | Multiple | 9 |
| Kaniewska *et al*[11], 2019 | Retrospective | CD | 95/82 | CDAI < 150 | NS | NS | 48 | Multiple | 7 |

CD: Crohn's disease; ADA: Adalimumab; IFX: Infliximab; CDAI: Crohn's disease activity index; HBI: Harvey Bradshaw Index; NOS: Newcastle-Ottawa Quality Assessment Scale; NS: Not stated.

**Table** **2 Subgroup analysis**

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Outcomes of interest** | **Subgroup analysis** | | | | | | | |
| **Grouping criteria** | **Categories** | **Studies, *n*** | **Patients, *n*** | **OR** | **95%CI** | ***I*2, %** | ***P* value** |
| Induction of response | Anti-TNF naivety | Naïve | 3 | 727 | 1.17 | (0.80-1.70) | 0 | 0.41 |
| Non-naïve | 3 | 313 | 1.44 | (0.55-3.78) | 47 | 0.46 |
| Use optimization | Yes | 5 | 1040 | 1.27 | (0.93-1.74) | 0 | 0.14 |
| No | 0 | 0 | - | - | - | - |
| Induction of remission | Anti-TNF naivety | Naïve | 2 | 392 | 1.08 | (0.68-0.72) | 0 | 0.75 |
| Non-naïve | 2 | 420 | 1.15 | (0.67-1.96) | 0 | 0.62 |
| Use optimization | Yes | 3 | 731 | 1.08 | (0.76-1.55) | 0 | 0.66 |
| No | 1 | 81 | 1.69 | (0.34-8.44) | - | 0.52 |
| Maintenance of response | Anti-TNF naivety | Naïve | 5 | 1468 | 1.08 | (0.72-1.62) | 63 | 0.71 |
| Non-naïve | 3 | 354 | 1.10 | (0.64-1.90) | 0 | 0.73 |
| Use optimization | Yes | 6 | 1645 | 1.12 | (0.77-1.63) | 62 | 0.57 |
| No | 1 | 177 | 0.64 | (0.18-2.29) | - | 0.50 |
| Maintenance of remission | Anti-TNF naivety | Naïve | 3 | 442 | 1.39 | (0.92-2.11) | 0 | 0.12 |
| Non-naïve | 3 | 328 | 1.24 | (0.56-2.72) | 53 | 0.59 |
| Use optimization | Yes | 3 | 458 | 1.41 | (0.95-2.09) | 0 | 0.09 |
| No | 3 | 312 | 1.18 | (0.46-2.99) | 55 | 0.73 |
| Secondary loss of response | Anti-TNF naivety | Naïve | 3 | 353 | 1.09 | (0.54-2.18) | 42 | 0.81 |
| Non-naïve | 3 | 947 | 0.91 | (0.46-1.80) | 72 | 0.78 |
| Use optimization | Yes | 5 | 1247 | 1.07 | (0.69-1.67) | 56 | 0.75 |
| No | 1 | 53 | 0.48 | (0.13-1.68) | 54 | 0.99 |
| Overall adverse events | Anti-TNF naivety | Naïve | 5 | 1184 | 0.67 | (0.50-0.89) | 1 | 0.005 |
| Non-naïve | 4 | 469 | 0.41 | (0.31-1.31) | 79 | 0.13 |
| Assessment time | ≤ 48 wk | 6 | 1323 | 0.50 | (0.33-0.76) | 41 | 0.001 |
| > 48 wk | 2 | 330 | 1.00 | (0.62-1.60) | 0 | 0.98 |
| Severe adverse events | Anti-TNF naivety | Naïve | 3 | 1021 | 0.88 | (0.40-1.92) | 73 | 0.74 |
| Non-naïve | 4 | 526 | 0.45 | (0.03-6.51) | 81 | 0.56 |
| Assessment time | ≤ 48 wk | 4 | 746 | 1.32 | (0.80-2.19) | 0 | 0.28 |
| > 48 wk | 3 | 801 | 0.52 | (0.09-3.05) | 80 | 0.47 |
| Opportunistic infections | Anti-TNF naivety | Naïve | 4 | 1654 | 0.78 | (0.54-1.14) | 0 | 0.21 |
| Non-naïve | 2 | 256 | 1.88 | (0.93-3.82) | 0 | 0.08 |
| Assessment time | ≤ 48 wk | 3 | 782 | 0.85 | (0.56-1.28) | 0 | 0.43 |
| > 48 wk | 3 | 1128 | 1.12 | (0.38-3.24) | 57 | 0.84 |

anti-TNF: Anti-tumor necrosis factor; OR: Odds ratio; CI: Confidence interval.

**Table 3 GRADE evidence profile**

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Quality assessment-No. of studies** | **Quality assessment-study design** | **Quality assessment-risk of bias** | **Quality assessment-inconsistency** | **Quality assessment-indirectness** | **Quality assessment-imprecision** | **Quality assessment-Publication bias** | **Summary of findings-number of patient, with IFX** | **Summary of findings-number of patient, with ADA** | **Summary of findings-effect, relative (95%CI)** | **Summary of findings-effect, absolute (95%CI)** | **Summary of findings-effect, Quality** |
| Induction of response | | | | | | | | | | | |
| 5 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 403/525 | 417/515 | OR: 1.27 (0.93-1.74) | 768 *per* 1000 | ⨁⨁◯◯ |
| Induction of remission | | | | | | | | | | | |
| 4 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 368/494 | 244/318 | OR: 1.11 (0.78-1.57) | 745 *per* 1000 | ⨁⨁◯◯ |
| Maintenance of response | | | | | | | | | | | |
| 7 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 611/896 | 639/932 | OR: 1.02 (0.83-1.25) | 682 *per* 1000 | ⨁⨁◯◯ |
| Maintenance of remission | | | | | | | | | | | |
| 6 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 255/442 | 219/328 | OR: 1.26 (0.87-1.82) | 577 *per* 1000 | ⨁⨁◯◯ |
| Secondary loss of response | | | | | | | | | | | |
| 6 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 191/704 | 172/603 | OR: 1.01 (0.65-1.55) | 271 *per* 1000 | ⨁⨁◯◯ |
| Overall adverse events | | | | | | | | | | | |
| 8 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 364/900 | 205/753 | OR: 0.62 (0.42-0.91) | 404 *per* 1000 | ⨁⨁◯◯ |
| Severe adverse events | | | | | | | | | | | |
| 7 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 139/859 | 80/688 | OR: 0.75 (0.32-1.72) | 162 *per* 1000 | ⨁⨁◯◯ |
| Opportunistic infection | | | | | | | | | | | |
| 6 | Observational study | Not serious | Not serious | Not serious | Not serious | Not found | 144/988 | 85/922 | OR: 0.96 (0.66-1.40) | 146 *per* 1000 | ⨁⨁◯◯ |

GRADE Working Group grades of evidence: High quality (⨁⨁⨁⨁): Further research is unlikely to change our confidence in the estimate of effect; Moderate quality (⨁⨁⨁◯): Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate; Low quality (⨁⨁◯◯): Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate; Very low quality (⨁◯◯◯): We are very uncertain about the estimate. ADA: Adalimumab; IFX: Infliximab; OR: Odds ratio; CI: Confidence interval.



Published by **Baishideng Publishing Group Inc**

7041 Koll Center Parkway, Suite 160, Pleasanton, CA 94566, USA

**Telephone:** +1-925-3991568

**E-mail:** bpgoffice@wjgnet.com

**Help Desk:** https://www.f6publishing.com/helpdesk

https://www.wjgnet.com



**© 2022 Baishideng Publishing Group Inc. All rights reserved.**