Systematic Review: Non-medical Switching of Infliximab to CT-P13 in Inflammatory Bowel Disease

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Abstract

Background and Aims  Biosimilar approval, such as Inflectra™ (CT-P13) for treating ulcerative colitis (UC) and Crohn’s disease (CD), has reduced direct drug costs. Though clinicians are comfortable with biosimilar use in treatment-naïve patients, there are concerns in some jurisdictions that there are insufficient data from well-controlled trials to support non-medical switching. A systematic review, along with a critical assessment of the study design, was conducted to assess the potential impact of switching stable CD/UC patients from infliximab to CT-P13.

Methods  A literature search using PubMed and abstracts/posters from 3 major gastroenterology conferences from 2014 to 2018 was completed. Two individual reviewers extracted data from each relevant report and compiled it into evidence tables to facilitate descriptive analyses. Key randomized trial and observational study designs were critically assessed to contextualize data relevance.

Results  A total of 49 reports (3 randomized controlled trials, 40 observational trials, and 1 case series) were included. Most studies revealed no efficacy, safety, or immunogenicity concerns with non-medical switch. Limitations of supporting data include a small number of randomized controlled trials; predominance of observational studies with varying outcome assessments and lack of appropriate controls; and scarcity of research on biosimilar switch long-term effects.

Conclusions  The majority of studies suggested non-medical switch is safe. However, clinicians and regulatory bodies should be aware of differences and limitations in study designs when making inferences about the risks and benefits of switching stable IBD patients to biosimilars.

Keywords  Inflammatory bowel disease · Ulcerative colitis · Crohn’s disease · Infliximab · CT-P13 · Biosimilar

Introduction

The introduction of biologic therapies approximately 20 years ago resulted in unparalleled benefits for the treatment of chronic immune-mediated inflammatory diseases (IMIDs), such as rheumatoid arthritis (RA) and inflammatory bowel disease (IBD). Biologic drugs are complex macromolecules produced from living cells that are genetically modified for commercial-scale production and are estimated to account for approximately 32% of global pharmaceutical sales by 2023 [1]. Remicade® (infliximab) was the first biologic (an originator biologic) on the market for IBD. It is a murine chimeric anti-tumor necrosis factor α (TNFα) monoclonal antibody used for the treatment of RA, ankylosing spondylitis (AS), plaque psoriasis (PsO), psoriatic arthritis (PsA), ulcerative colitis (UC), and Crohn’s disease (CD) [2]. In 2016, global Remicade sales reached almost $8 billion (USD) [3]. In addition to market share growth, infliximab
has been increasingly used for IBD treatment to prevent UC and CD progression in some countries [4–7].

Over the past 5 years, a number of biosimilars have been approved for use by Health Canada. Biosimilars are biologic therapies that are highly similar, but not identical, to their respective originator biologic products. Although biosimilarity does not require all aspects of the biosimilar and originator products to be the same, biosimilars go through a rigorous comparative pre-approval testing process, where data demonstrating similarity between the originator and biosimilar must be provided [8]. As such, the type of data to support biosimilars is different than for a stand-alone biologic, and these attenuated clinical trial requirements mean that biosimilars can be offered at a lower price point versus originator molecules. Biologic therapies are complex, microheterogeneous molecules that are highly sensitive to changes in both raw materials and manufacturing conditions. Therefore, small differences exist between biosimilars and their originator products, which may lead to differences in efficacy, safety, or immunogenicity [8].

CT-P13 (marketed as Inflectra® in Canada and Remsima™ elsewhere) is the first infliximab biosimilar approved in Canada for similar indications as Remicade®, except for pediatric UC and CD, based on clinical trials in AS and RA. The availability of this lower-cost alternative may prompt payors to adopt strategies that prioritize biosimilars preferentially over innovator use to reduce healthcare expenditures [9]. In several countries, non-medical switch from an originator molecule to its biosimilar has emerged as a treatment strategy, typically driven by economic or access reasons. While the safety and efficacy of biosimilar use in infliximab-naïve patients is generally accepted, a “forced switch” to save costs concerns some gastroenterologists, who cite a lack of high-quality controlled trials to support such decisions in stable patients. The objective of this systematic review was to examine the current literature on biosimilar switch and its implications for guiding treatment decisions. To that end, we conducted a systematic review of studies in which patients with CD or UC were switched from infliximab to CT-P13 and reviewed the potential impact on efficacy, response, and safety. The study designs of key randomized clinical trials and observational studies were also critically assessed to contextualize the relevance of the available data.

Materials and Methods

The systematic review search strategy was developed to examine the following participants, interventions, comparisons, and outcomes (PICO) items: (P) randomized controlled trials and observational studies of patients diagnosed with CD or UC who were switched to (I) CT-P13 from (C) infliximab and the effect of non-medical switch on (O) efficacy/effectiveness, response, and safety as inferred from disease worsening, loss of response, and sustained remission rate data. This systematic review was not registered with PROSPERO. This systematic review is in accordance with the PRISMA checklist (Table 1).

Search Methods

A systematic review of the PubMed database was completed in April 2018, using the terms “((switch OR switching OR interchange OR interchanged [All Fields])) AND infliximab [All Fields]) AND (ulcerative colitis OR inflammatory bowel disease OR Crohn[MeSH Terms])” with no date restrictions, filtered by text availability (only full-text publications were included). Abstracts from the Congress of European Crohn’s and Colitis Organization (ECCO, 2013–2018) and Digestive Disease Week (DDW, 2013–2017) via the journal Gastroenterology were searched using the terms “infliximab AND biosimilar AND (inflammatory OR IBD)” and “infliximab AND biosimilar.” The absence of a searchable database for United European Gastroenterology Week (UEGW, 2013–2017) required oral and poster abstracts, in portable document format, to be scanned using the term “infliximab.”

Selection Criteria

Titles and abstracts were identified as potentially eligible by two independent individuals. Studies were eligible if they included data on patients with CD or UC treated with infliximab originator who were stable at the time of switch to CT-P13 and were written in the English language. Appropriate conference abstracts were analyzed to minimize publication bias. UEGW abstracts that included the term “infliximab” were further screened for eligibility criteria, as mentioned above. Review articles, meta-analyses, and other non-clinical papers were excluded. Two individuals then independently examined the full text of the remaining publications and conference abstracts deemed to match the eligibility criteria, with data extracted in duplicate. Any discrepancies were resolved by discussion.

Data Extraction

Data were extracted from each relevant report. Each publication was individually reviewed to identify data concerning: study information—primary author, year of publication, title of publication, type of study (randomized or observational); patient characteristics—total number of patients, number of included patients with CD or UC; and study data—interventions, duration of follow-up, efficacy outcomes (disease worsening, loss of response, and sustained remission rates), discontinuation rate, immunogenicity, and safety (adverse
## Table 1 PRISMA checklist

| Section/topic                        | # | Checklist item                                                                                                                                 |
|--------------------------------------|---|---------------------------------------------------------------------------------------------------------------------------------------------|
| **Title**                            | 1 | Identify the report as a systematic review, meta-analysis, or both                                                                       |
| **Abstract**                         | 2 | Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number |
| **Introduction**                     | 3 | Describe the rationale for the review in the context of what is already known                                                             |
| **Objectives**                       | 4 | Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS) |
| **Methods**                          | 5 | Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number |
| Protocol and registration            | 6 | Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale |
| Eligibility criteria                 | 7 | Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched |
| Information sources                  | 8 | Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated |
| Search                               | 9 | State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis) |
| Study selection                      | 10| Describe the method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators |
| Data items                           | 11| List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made |
| Risk of bias in individual studies   | 12| Describe the methods used for assessing the risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis |
| Summary measures                     | 13| State the principal summary measures (e.g., risk ratio, difference in means) |
| Synthesis of results                 | 14| Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., $I^2$) for each meta-analysis |
| Risk of bias across studies          | 15| Specify any assessment of the risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies) |
| Additional analyses                  | 16| Describe the methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified |
| **Results**                          | 17| Give the numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram |
| Study selection                      | 18| For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations |
| Risk of bias within studies          | 19| Present data on the risk of bias of each study and, if available, any outcome level assessment (see Item 12) |
| Results of individual studies        | 20| For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group, (b) effect estimates and confidence intervals, ideally with a forest plot |
| Synthesis of results                 | 21| Present the main results of the review. If meta-analyses are done, include for each, confidence intervals and measures of consistency |
| Updated in accordance with           |    |                                                                                                                                             |
| Risk of bias across studies          | 22| Present results of any assessment of the risk of bias across studies (see Item 15) |
| Additional analysis                  | 23| Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression (see Item 16)) |
Table 1 (continued)

| Section/topic         | #  | Checklist item                                                                                   |
|-----------------------|----|-------------------------------------------------------------------------------------------------|
| Discussion            |    | Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policymakers) |
| Summary of evidence   | 24 |                                                                                                  |
| Limitations           | 25 | Discuss limitations at study and outcome level (e.g., risk of bias) and at review level (e.g., incomplete retrieval of identified research, reporting bias) |
| Conclusions           | 26 | Provide a general interpretation of the results in the context of other evidence and implications for future research |
| Funding               |    | Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review |
| Funding               | 27 |                                                                                                  |

effects). Unstated data were designated as non-available (N/A). The extracted data were compiled into evidence tables to facilitate descriptive analysis.

Results

The literature search identified 2172 potential symposium abstracts and 144 potential publications from PubMed. A total of 27 symposium abstracts and 22 publications were deemed appropriate for final inclusion based on the eligibility criteria outlined in Fig. 1. This included 5 reports from 3 randomized controlled trials (1 publication; 4 abstracts) and 44 reports from 40 observational trials and 1 case series (21 publications; 23 abstracts), representing a total of 44 distinct studies.

Evidence from Randomized Controlled Trials

Data from a total of 3 distinct randomized controlled trials were reported in either published articles or abstract format and accounted for just 5 of the 49 publications included in this review. Of these 3 trials, data from the NOR-SWITCH trial were reported in 1 published article and 2 conference abstracts (Table 2). This randomized, double-blind, parallel group, non-inferiority trial evaluated the safety and efficacy of switching from infliximab to CT-P13 versus infliximab maintenance treatment across 6 inflammatory diseases [10].

Fig. 1 Flowchart of literature search outcomes
| Study | N  | Intervention                                                                 | Duration of follow-up | Efficacy outcomes                                                                 | Discontinuation rate                                                                 | Immunogenicity                                                                 | Safety                                                                 |
|-------|----|------------------------------------------------------------------------------|-----------------------|-----------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|--------------------------------------------------------------------------------|----------------------------------------------------------------------|
| **Published studies** |    |                                                                              |                       |                                                                                   |                                                                                        |                                                                                |                                                                      |
| Jørgensen KK et al., 2017, Norway (NOR-SWITCH study) | N = 482 | Maintenance on IFX, Switch to CT-P13                                           | 52 weeks              | Disease worsening: CD: 21.2% in IFX, 36.5% CT-P13                                        | 4% of the IFX group and 4% of the CT-P13 group at 52 weeks in the full analysis set   | New ADA in 7% of the IFX group and 8% of CT-P13 switch group in the full analysis set | No significant differences in TEAEs, or serious TEAEs                  |
| Abstracts |    |                                                                              |                       |                                                                                   |                                                                                        |                                                                                |                                                                      |
| Jørgensen KK et al., 2017, Norway (NOR-SWITCH subgroup analysis) | N = 481 | Maintenance on IFX, Switch to CT-P13                                           | 52 weeks              | CD PGA: -0.65 (-1.22, -0.07) CD PGA: -0.42 (-0.85, 0.001)                             | N/A                                                                                     | No significant difference in ADA, trough serum levels                  | No significant difference in AEs                                      |
| Jørgensen KK et al., 2017, Norway (NOR-SWITCH extension trial, subgroup analysis) | N = 198 | Maintenance on CT-P13 after DB phase, Switch to CT-P13 after DB phase          | 26 weeks              | Disease worsening: CD: 20.6% (13/63) maintenance, 13.1% (8/61) switch                | N/A                                                                                     | No significant difference in ADA, trough serum levels                  | No significant difference in AEs                                      |
| Kim YH et al., 2017, International study | N = 220 | Maintenance on IFX, Maintenance on CT-P13, Switch to IFX at week 30, Switch to CT-P13 at week 30 | 54 weeks              | No significant differences in CDAI-70 at week 54                                      | N/A                                                                                     | No significant difference in ADA                                      | No significant difference in AEs, infection rates                    |
| Volkers AG et al., 2017, Netherlands (SIMILAR trial) | N = 47  | Maintenance on IFX, Switch to IFX biosimilar                                   | 30 weeks              | 1 patient on IFX biosimilar relapsed                                                 | N/A                                                                                     | N/A                                                                           | No SAEs related to study drug                                        |

ADAs antidrug antibodies, AE adverse event, CD Crohn’s disease, CDAI Crohn’s Disease Activity Index, DB double-blind, IFX infliximab originator, N/A not available, PGA physician’s global assessment, PtGA patient’s global assessment, SAE severe adverse event, TEAE treatment-emergent adverse event, UC ulcerative colitis
Patients enrolled in the 52-week NOR-SWITCH trial were ≥ 18 years of age with RA, spondyloarthritis (SpA), PsA, UC, CD, or PsO, and clinically stable on infliximab for ≥ 6 months. The primary endpoint was a composite endpoint of disease worsening, presumably designed to increase event rate and reflect the pragmatic nature of the trial (UC: increase in partial Mayo [p-Mayo] score of ≥ 3 and a p-Mayo score of ≥ 5; CD: increase in Harvey-Bradshaw Index [HBI] of ≥ 4 and a HBI score of ≥ 7; RA/ PsA: increase in Disease Activity Score in 28 joints [DAS28] of ≥ 1.2 from randomization and a DAS28 of ≥ 3.2; AS/SpA: increase in Ankylosing Spondylitis Disease Activity Score [ASDAS] of ≥ 1.1 and ASDAS of ≥ 2.1; PsO: increase in Psoriasis Area and Severity Index [PASI] of ≥ 3 from randomization and a minimum PASI of ≥ 5). A 15% non-inferiority margin was used at 90% power [10].

The authors concluded that CT-P13 was non-inferior to infliximab based on the disease worsening composite endpoint (29.6% vs. 26.2%, respectively), with a risk difference of −4.4% (95% confidence interval [CI]; −12.7–3.9) adjusted for diagnosis and treatment duration of infliximab originator at baseline. Independently, results in 5 out of 6 disease categories failed to demonstrate non-inferiority. In CD, the risk difference for switching was −14.3% (95% CI −29.3–0.7) [10, 11]. In UC, disease worsening was observed in 9.1% and 11.9% of switch and infliximab maintenance patients, respectively, with a risk difference of −2.6% (95% CI −15.2–10.0) [10]. However, these analyses should be considered exploratory, as the trial design was not powered to examine subgroups.

Recently, Jørgensen et al. presented an IBD subgroup analysis from the 26-week, NOR-SWITCH open-label extension trial [12]. Stable switch patients on CT-P13 were enrolled into the maintenance arm after the initial 52-week trial, while those on infliximab maintenance were switched to CT-P13. The primary endpoint was disease worsening according to HBI for CD and the p-Mayo score for UC. Disease worsening occurred in 20.6% versus 13.1% CD patients (risk difference 7.9%, 95% CI −5.2–21) and 15.4% versus 2.9% UC patients (risk difference 12.4%, 95% CI −0.1–25) on CT-P13 maintenance versus those switching from infliximab, respectively. Though low enrollment numbers precluded conclusive implications to be drawn, the authors stated that efficacy, safety, and immunogenicity were similar between treatment arms [12].

The second randomized controlled trial included was presented by Kim et al. at ECCO 2017 (NCT02096861) [13]. This 54-week, phase III randomized, double-blind, parallel group, non-inferiority trial examined the efficacy and overall safety of CT-P13 versus infliximab in 220 patients with active CD. Patients were randomized to CT-P13 or infliximab and followed for 30 weeks. At week 30, patients in each respective arm were re-randomized to either continue their current therapy or switch, resulting in a total of 4 treatment arms. The primary objective of the study was to compare the efficacy between CT-P13 and infliximab in terms of Crohn’s Disease Activity Index-70 (CDAI-70) response rates, defined as a reduction in at least 70 points from baseline CDAI score at week 6. There was no difference in the primary endpoint between arms (CT-P13, 71.4%; infliximab, 75.2%; p = 0.5613). The study also examined secondary outcomes associated with CDAI-100 response (reduction in at least 100 points in baseline CDAI score) and clinical remission rates (absolute CDAI score of less than 150 points). Similar trends to CDAI-70 were observed for the more stringent and traditional outcome of CDAI-100 (CT-P13, 61.9%; infliximab, 64.4%; p = 0.7744) and clinical remission rates (CT-P13, 42.9%; infliximab, 44.6%; p = 0.8329) at week 6 [13]. A similar proportion of patients between treatment arms were found to have at least one treatment-emergent serious adverse event (CT-P13, 1.8% [2/111]; infliximab, 1.8% [2/109]) [13].

Preliminary results of the third and final included randomized controlled trial were presented by Volkers et al. at UEGW 2017 (NCT02452151) [14]. This randomized, controlled, double-blind, phase IV, non-inferiority trial examined CD and UC patients in clinical and biochemical remission while on infliximab therapy who were randomized to either CT-P13 or infliximab maintenance treatment. The primary endpoint was remission at week 30. A total of 47 patients (35 CD, 12 UC) had been recruited as of October 2017, of whom 21 completed the 30-week follow-up (15 received CT-P13 and 6 infliximab). One patient on CT-P13 experienced a loss of response, and no immunogenicity data were provided. Though preliminary, the authors concluded that switching from infliximab to the biosimilar is feasible and safe [14].

**Evidence from Published Observational Switch Studies**

Published observational studies accounted for 21 of the total 49 reports included in this review. Due to the significant inter-study heterogeneity of the patient populations, baseline characteristics, dosing schedules, and follow-up periods, a meaningful comparison between the published observational switch studies becomes difficult to conduct and interpret. For example, 6 out of 21 published observational studies (Table 3) reported a loss of effectiveness over the study period for both CD and UC patient populations, with values ranging from 6.3 to 30.2% and 0 to 22%, respectively [15–20]. Five studies reported a loss of response for IBD switch groups as a whole, varying from 0 to 24.6% [21–25]. The observational study by Ratnakumaran et al. was the only one that utilized a control group on infliximab maintenance therapy and also reported on the secondary loss of response.
| Study                          | N   | Intervention                                                                 | Duration of follow-up | Effectiveness of medication                                                                 | Response and/or remission criteria                                      | Discontinuation rate | Immunogenicity                                                                 | Safety                                      |
|-------------------------------|-----|------------------------------------------------------------------------------|-----------------------|-----------------------------------------------------------------------------------------------|--------------------------------------------------------------------------|----------------------|--------------------------------------------------------------------------------|--------------------------------------------|
| Avouac J et al., 2018, France | N = 64 | Switch to CT-P13                                                               | 3 biosimilar infusions | HBI (baseline; follow-up): 0.83; 0.56 PM (baseline; follow-up): 0.71; 0.33 No significant change in disease activity scores | N/A                                                                     | 22% for CD patients by 3rd infusion; 13% of all IBD at a mean follow-up of 34 weeks | No significant change in serum trough concentration after switch | No SAEs                                      |
| Binkhorst L et al., 2018, The Netherlands | N = 197 | Switch to CT-P13                                                               | 16 weeks              | Loss of response: 5.5% (n = 11) Increase in disease-related symptoms as established by the physician and/or changes in treatment | 10%                                                                     | No significant change in serum trough concentration after switch | No SAEs or unexpected AEs                         | 1 severe acute IRR                           |
| Fiorino G et al., 2018, Italy and Spain | IFX: n = 13 CT-P13: n = 9 Switch: n = 12 | Maintenance on IFX Initiated on CT-P13 Switch to CT-P13 | N/A                   | N/A                                                                                           | N/A                                                                     | N/A                  | 16.7% (2/12) developed ADA after switch to CT-P13; 100% cross-reactivity of ADAs between IFX, CT-P13, and SB2 | N/A                                        |
| Kang B et al., 2018, Korea     | N = 74 (pediatric) | Maintenance on IFX (n = 36) Switch to CT-P13 (n = 38) | 1 year                | Sustained remission rate: IFX: 77.8%, Switch: 78.9% (p = 1.00) Clinical remission lasting for at least 2 years plus mucosal healing identified on 2 consecutive annual follow-up colonoscopies | IFX: 13.9%, Switch: 7.9% (p = 0.649)                                      | Therapeutic trough levels: IFX: 90.3%, Switch: 88.6%; New ADA: IFX: n = 2, Switch: n = 1 | 27 AEs in IFX vs. 30 in switch group          | N/A                                        |
| Schmitz EMH et al., 2018, The Netherlands | N = 133 | Switch to CT-P13                                                               | 1 year                | Disease worsening: 9% (n = 12; with 2 deemed not switch-related)                               | N/A                                                                     | 26%                  | 3 developed ADAs after switch                                                   | 9.8% discontinued treatment due to AEs     |
| Strik A et al., 2018, Belgium and The Netherlands (SECURE study) | N = 120 | Switch to CT-P13                                                               | 16 weeks              | Loss of response at 16 weeks: CD: 12% UC: 24%                                                  | Increase in HBI > 4 in CD patients and in SCCAI > 2 in UC patients      | Geometric mean ratio of drug concentration (week 16 vs. baseline): CD: 107.6% (97.4–118.8); UC: 110.1% (90% CI 96.0–126.3); 3 developed new ADAs after switch | 6 SAEs, with only 1 (perianal abscess) deemed related to study treatment | N/A                                        |
| Arguelles Arias F et al., 2017, Spain | N = 120 | Switch to CT-P13 (n = 98) Naive patients initiated on CT-P13 (n = 22) | 6 months              | Sustained remission rate in switch patients: CD: 87.5% at 3 months, 83.9% at 6 months UC: 92% at 3 months, 91.3% at 6 months | No need for steroids, surgery or increased dose in both CD and UC switch patients | N/A                  | SAEs in 7.5% of total study population                                          | N/A                                        |
| Study                          | N          | Intervention          | Duration of follow-up | Effectiveness of medication | Response and/or remission criteria | Discontinuation rate | Immunogenicity | Safety          |
|-------------------------------|------------|-----------------------|-----------------------|-----------------------------|-----------------------------------|----------------------|---------------|----------------|
| Argüelles-Arias F et al., 2017, Spain | N=98  
n=67 CD  
n=31 UC | Switch to CT-P13 | 1 year | Disease worsening at 1 year: CD: 30.2% UC: 19%; Sustained remission rates: CD: 83.6% at baseline, 62.7% at 1 year UC: 90.6% at baseline, 64.3% at 1 year | Maintenance of HBI ≤ 4 in CD patients or PM ≤ 2 in UC patients, without the need for steroids, surgery or increased dose after switching | N/A                      | CD: 9%  
UC: 19.4% | N/A | SAEs in 11.2% of patients |
| Buer LCT et al., 2017, Norway  | N=143  
n=99 CD  
n=44 UC | Switch to CT-P13 | 6 months | Sustained remission rates: CD: 87% at baseline, 81% at 6 months UC: 88% at baseline, 95% at 6 months | Maintenance of HBI ≤ 4 in CD patients and PM < 2 in UC patients | 3%                    | 3 developed ADAs after switch | Rate of IRRs 0.7%  
(5/750) | 17 AEs reported, 2 resulting in treatment discontinuation |
| Eberl A, et al., 2017, Finland | N=62  
n=32 CD  
n=30 UC | Switch to CT-P13 | 3 biosimilar infusions | No change in disease activity (HBI, PM score) | N/A                              | N/A                      | 2 developed ADAs after switch; Trough levels: CD: 5.75 mg/L at baseline, 5.5 mg/L at 3rd infusion (p=0.05) UC: 5.2 mg/L at baseline, 4.25 mg/L at 3rd infusion (p=0.019) | Rate of IRRs 2.5%  
(4/156 infusions) |
| Fiorino G et al., 2017, Italy (PROSIT-BIO cohort) | N=97  
n=53 CD  
n=44 UC | Switch to CT-P13 | 6.1 ± 2.2 months (mean ± SD) | Kaplan–Meier estimate for the probability of response: 94.5% at 8 weeks, 90.8% at 16 weeks, 78.9% at 24 weeks | Kaplan–Meier estimate for the probability of treatment persistence: 100% at 8 weeks, 98.7% at 16 weeks, 92.1% at 24 weeks | N/A                      | N/A | SAEs in 12.4% of patients; discontinuation due to SAEs in 3.1% IRRs in 7.2% of patients; discontinuation due to IRRs in 2.1% |
| Guerrero Puente L et al., 2017, Spain | N=36  
n=23 CD  
n=13 UC | Switch to CT-P13 | 8.4 ± 3.5 months (mean ± SD) | Loss of response: 13.9% (mean time to relapse= 2.4 months); Relapse-free survival at 6 months: 85.7% | Increased disease activity as assessed by the clinic, radiology, laboratory and endoscopy findings | N/A                      | No significant difference in pre- and post-switch drug levels at 8 and 16 weeks of follow-up | 3 (8.3%) of patients experienced AEs; 1 resulted in treatment discontinuation No IRR, malignancy or death |
| Study                        | N   | Intervention | Duration of follow-up | Effectiveness of medication | Response and/or remission criteria | Discontinuation rate | Immunogenicity | Safety                     |
|-----------------------------|-----|--------------|-----------------------|-----------------------------|-----------------------------------|---------------------|----------------|---------------------------|
| Kolar M et al., 2017, Czech Republic | N=74 | Switch to CT-P13 | 56 weeks             | Sustained remission rates: 72.2% at baseline, 77.8% at week 56 (p = 0.55) | Maintenance of HBI ≤ 4 in CD patients and PM < 2 in UC patients | N/A                | ADA positive patients: 9.5% at baseline, 6.0% at week 56 | 2 patients with CD discontinued due to AEs |
| Ratnakumaran R et al., 2017, UK | N=210 | Switch to CT-P13 (n = 191) | 1 year | Loss of response: Switch: 24.6% IFX: 42.1% (p = 0.10); Sustained remission rate: Switch: 58.1% IFX: 47.4% (p = 0.37) | Increased disease activity according to physician’s global assessment. Asymptomatic patients in clinical assessment were considered in remission | N/A                | N/A                              | Incidence of AEs were 0% for the IFX group and 4.7% for the switch group (p = 1.0) |
| Razanskaite V et al., 2017, UK | N=143 | Switch to CT-P13 | Up to 1 year | IBD-Control-8 Score: 11 at switch, 14 at 3rd infusion (p = 0.041) | N/A                              | 27% at 1 year | Mean ADA levels: Pre: 73.2 ± 83.9 AU/mL Post: 96.1 ± 102.0 AU/mL (p = 0.24); no significant difference in pre- and post-switch trough level | The most common AEs at 3rd infusion were headaches (17%), joint pain (13.8%) and infections (13.8%) |
| Smits LJT et al., 2017, The Netherlands | N=83 | Switch to CT-P13 | 52 weeks             | Sustained remission rates: 64% at baseline, 73% at week 52 | Maintenance of HBI ≤ 4 in CD patients and SCCAI ≤ 3 in UC patients | 18%                | 2 developed ADAs after switch; trough levels unaffected by switch | 7% (6) discontinued due to AEs |
| Sieczkowska J et al., 2016, Poland | N=39 (pediatric) | Switch to CT-P13 | CD: 8 ± 2.6 months (mean ± SD) UC: 5 ± 3.6 months (mean ± SD) | Loss of response: CD: 6.3% UC: 14.2%; Sustained remission rate CD: 69% at baseline, 88% at last infusion Sustained remission rate UC: 29% at baseline, 57% at last infusion Mean PCDAI (range): At switch: 8.8 (0–35) 32 weeks: 6.6 (0–43) Mean PUCAI (range): At switch: 16.4 (0–30) 4th infusion: 2 (0–5) | N/A                              | CD: 37% UC: 43% | N/A                          | 2 IRRs after switch, 1 resulting in treatment discontinuation 1 UC patient discontinued due to varicella zoster infection after 1st CT-P13 infusion CT-P13 safety profile consistent with IFX |
| Study | N | Intervention | Duration of follow-up | Effectiveness of medication | Response and/or remission criteria | Discontinuation rate | Immunogenicity | Safety |
|-------|---|-------------|----------------------|-----------------------------|-----------------------------------|----------------------|---------------|--------|
| Smits LJT et al., 2016, The Netherlands | N=83  
  n=57 CD  
  n=24 UC  
  n=2 IBD-u | Switch to CT-P13 | 16 weeks | Sustained remission rate: CD: 60% at baseline, 67% at week 16  
 UC: 73% at baseline, 62% at week 16 | Maintenance of HBI ≤ 4 in CD patients and SCCAI ≤ 3 in UC patients. | 6% | Median trough level: 3.5 μg/ml (range 0–18) at baseline, 4.2 μg/ml (range 0–21) at week 16 (p=0.010); 2 developed ADAs post-switch | No SAEs |
| Jung YS et al., 2015, Republic of Korea | N=36  
  n=27 CD  
  n=9 UC | Switch to CT-P13 | 54 weeks | Loss of response: CD: 7.4%  
 UC: 11.1% | Response in CD patients: ≥ 70-point CDAI decrease from baseline and ≥ 25% reduction in total CDAI score;  
 Response in UC patients: 3-point decrease in total Mayo score or 2-point decrease in PM, along with 1-point decrease in rectal bleeding subscore or absolute rectal bleeding subscore ≤ 1 | 13.9% | N/A | 1 discontinued due to AEs |
| Kang YS et al., 2015, Republic of Korea | N=9  
  n=5 CD  
  n=4 UC | Switch to CT-P13 | 8 weeks | Loss of response: CD: 20%  
 UC: 0% | Response in CD patients: > 70-point decrease in CDAI  
 Response in UC patients: > 30% decrease in Mayo score, along with decreased rectal bleeding and endoscopy subscore | N/A | N/A | 1 discontinued due to AEs |
Evidence from Observational Studies in Conference Abstracts

Data from observational studies presented as posters or abstracts at the selected international gastroenterology conferences accounted for 23 of the 49 total reports included in this review (Table 4) [36–58]. Similarly to the published empirical data, there was significant heterogeneity in patient populations, concomitant medications, and follow-up periods. Results demonstrated high variability in effectiveness, immunogenicity, and safety, with no studies utilizing an infliximab maintenance control arm. Loss of effectiveness for the combined IBD switch population was specifically highlighted in just 6 out of 23 studies, ranging from 4.2 to 25.0% [40, 43, 44, 46, 52, 54, 57]. Suk et al. was the only study to clearly specify loss of response in the CD and UC switch populations, at 14% and 33%, respectively [58]. Variability in the reported discontinuation rates and immunogenicity outcomes was also observed (Table 4).

Ilias et al. recently presented an abstract addressing the paucity of evidence on reverse switch to infliximab (i.e., switching back to infliximab after initial switch to CT-P13) [49]. Since 2014, all infliximab-naïve patients, and those who had not been exposed for at least 12 months, were mandated to start CT-P13 in Hungary. In August 2017, policy changes resulted in a mandatory reverse switch for all patients on CT-P13, back to originator infliximab. Data were collected from 117 stable CD and UC patients on CT-P13 maintenance from four IBD centers at the time of the reverse switch and 24 weeks post-reverse switch. Concomitant immunomodulators and steroids were used in 53.8/45.8% and 7.5/37.5% of CD/UC patients at the reverse switch, respectively. All but one patient, who experienced an infusion-related reaction, tolerated the first infusion after the reverse switch.

Evidence from Published Observational Studies with Updated Data in Conference Abstracts

Currently, the observational study with the most robust enrollment is for the PROSIT-BIO cohort, which used a post-switch. The authors concluded that observed inter-arm differences were statistically insignificant [22]. Discontinuation rates were reported for 12 out of 21 studies and ranged from 3 to 27% for the IBD switch groups and 9 to 37% and 19.4 to 43% for CD and UC switch groups, respectively [16, 19–21, 24–31]. Immunogenicity and safety data across the studies were diverse, but often lacked a control arm for comparison [15–35]. This variability makes it difficult for broad generalizations regarding the safety and effectiveness of non-medical switch to be made.

Table 3 (continued)

| Study | Intervention | Duration of follow-up | Effectiveness of medication | Response and/or remission criteria | Discontinuation rate | Immunogenicity | Safety |
|-------|--------------|-----------------------|----------------------------|-----------------------------------|---------------------|---------------|--------|
| Park SH et al., 2015, Republic of Korea | Switch to CT-P13 | 30 weeks | Disease worsening:CD: 12.9% FCD: 25% UC: 0%; Sustained remission rate:CD: 80.6% FCD: 50% UC: 45.5% | AD1, 14.7% of patients receiving 5 mg/kg dose had a TEAE; 6.7% receiving > 5 mg/kg had a TEAE | N/A | N/A | 11.7% of patients receiving 5 mg/kg dose had a TEAE; 6.7% receiving > 5 mg/kg had a TEAE |

AD1: antidrug antibodies, AE: adverse event, CD: Crohn’s disease, CDMA: Crohn’s disease activity index, IFX: infliximab originator, PCDAI: Pediatric Crohn’s disease activity index, PM: partial Mayo score, PUCAI: pediatric ulcerative colitis activity index, SAEC: severe adverse event, SCCAI: simple clinical colitis activity index, TEAE: treatment-emergent adverse event, UC: ulcerative colitis.
| Study | N     | Intervention | Duration of follow-up | Effectiveness of medication | Discontinuation rate | Immunogenicity | Safety |
|-------|-------|--------------|-----------------------|-----------------------------|----------------------|----------------|--------|
| Daperno M et al., 2018, Italy | N = 69  
 n = 33 CD  
 n = 36 UC | Switch to CT-P13  
 (n = 54)  
 Biologic-naïve patients started on CT-P13  
 (n = 15)  | <2 years | N/A | N/A | No significant difference in ADA and trough levels between the switch and naïve groups | The incidence of AEs is reduced in the switch vs. naïve group (p < 0.05) |
| Guerra Veloz MF et al., 2018, Spain | N = 167  
 n = 116 CD  
 n = 51 UD | Switch to CT-P13 | 1 year | Loss of response: 15.7%; Sustained remission rate (baseline vs. 1 year): CD: 88.8% vs. 69.7%  
 (p > 0.05)  
 UC: 84.3% vs. 76.7%  
 (p > 0.05) | 9% | N/A | SAEs reported in 7.2% of patients |
| Hoivic ML et al., 2018, Norway | N = 143  
 n = 99 CD  
 n = 44 UC | Switch to CT-P13 | 18 months | No significant change in disease activity scores | 8% | N/A | N/A |
| Ilias A et al., 2018, Hungary | N = 117  
 n = 98 CD  
 n = 19 UC | Reverse switch from CT-P13 to IFX | 24 weeks | N/A | N/A | N/A | N/A |
| Petitdidier N et al., 2018, France | N = 113  
 n = 83 CD  
 n = 28 UC | Switch to CT-P13 | 1 year | Loss of response: 12.4% | N/A | N/A | N/A |
| Pierik MJ et al., 2018, The Netherlands | N = 119  
 n = 77 CD  
 n = 42 UC | Switch to CT-P13 | 6 infusions | Mean (± SEM) change in disease activity compared with baseline: SCCAI: 0.1 ± 0.3  
 HBI: 0.5 ± 0.7 | N/A | N/A | N/A |
| Smits L et al., 2018, The Netherlands | N = 83  
 n = 57 CD  
 n = 24 UC  
 n = 2 IBD-u | Switch to CT-P13 | 2 years | Loss of response: 12%  
 (n = 10);  
 No significant change in SCCAI or HBI | 32% | N/A | 53 AEs, including 3 non-related SAEs (melanoma, subarachnoid hemorrhage, hospitalization with fever) 8 discontinued CT-P13 due to AEs |
| Choe YH et al., 2017, Republic of Korea | N = 32 (pediatric)  
 n = 25 CD  
 n = 7 UC | Switch to CT-P13 | 30 weeks | Sustained remission rate: CD: 86.4%  
 UC: 100%  | N/A | N/A | 2 had TEAEs  
 1 TESAE  
 1 had an IRR |
| Choe YH et al., 2017, Republic of Korea | N = 83  
 n = 8 FCD:  
 n = 75 CD: | Switch to CT-P13 | N/A | Sustained remission rate: FCD: 87.5%  
 CD: 80% | N/A | N/A | N/A |
### Table 4 (continued)

| Study                      | N          | Intervention                        | Duration of follow-up | Effectiveness of medication | Discontinuation rate | Immunogenicity | Safety                          |
|----------------------------|------------|-------------------------------------|-----------------------|----------------------------|----------------------|----------------|---------------------------------|
| Guidi L et al., 2017, Italy (PROSIT study) | N = 680 | Switch to CT-P13 (n = 109)           | 2+ infusions in switch group; safety analysis up to 1 year | No efficacy analysis specific for the switch group; Loss of response (whole cohort): 10.3%; Deep remission rate (whole cohort): CD: 57%; UC: 50% | N/A                  | N/A                          | 92 SAEs (13.5%) were reported in whole cohort, leading to discontinuation in 73 patients (10.7%) |
| Huoponen S et al., 2017, Finland | N = 56 | Switch to CT-P13 (n = 24 CD)         | 16 weeks              | No significant change in HBI or PM scores post-switch | N/A                  | N/A                          | No SAEs observed                  |
| Nugent S et al., 2017, Ireland | N = 33 | Switch to CT-P13 (n = 62 CD)         | 1 year                | N/A                        | 15%                  | N/A                          | N/A                              |
| Rodriguez Glez GE et al., 2017, Spain | N = 72 | Switch to IFX biosimilar (n = 60 UC) | 1 year                | Loss of response: 5.6%; Sustained remission rate (baseline vs. 1 year): 86% vs. 80.5% | 9.7%                 | N/A                          | 10 patients reported AEs; none resulted in treatment discontinuation |
| Sladek M et al., 2017, Italy & Poland | N = 45 (pediatric) | Switch to CT-P13 (n = 38 CD)       | 24–36 weeks           | No significant change in disease activity after switch | One patient developed ADA post-switch | N/A                          | 1 discontinued due to AE with high ADA, which presented before switch to CT-P13 |
| Bettey M et al., 2016, UK | N = 134 | Switch to CT-P13 (n = 62 CD)         | 16 weeks              | N/A                        | 1.5%                 | N/A                          | Similar incidence of AEs before and after switch |
| Bettey M et al., 2016, UK | N = 134 | Switch to CT-P13 (n = 62 CD)         | 16 weeks              | N/A                        | 1.5%                 | N/A                          | Similar incidence of AEs before and after switch |
| Díaz Hernández L et al., 2016, Spain | N = 72 | Switch to IFX biosimilar (n = 60 UC) | 6 months              | Loss of response: 4.2% (1 primary non-response, 2 secondary loss of response); Sustained remission rate: Baseline: 86.1% 6 months: 86.1% | 4.2%                 | N/A                          | 5 AEs reported (2 infection, 1 cutaneous injury, 1 cephalgia, 1 fever allergic reaction) None discontinued due to AEs |
| Guerra Veloz MF et al., 2016, Spain | N = 31 UC | Switch to CT-P13 (n = 29 switch/1 naïve) | 6 months              | Loss of response: 3 months (total 29 switch/1 naïve): 26.3% 6 months (total 15 switch/1 naïve): 30% | N/A                  | N/A                          | Only mild AEs reported |
| Guerra Veloz MF et al., 2016, Spain | N = 71 CD | Switch to CT-P13 (n = 60 switch/2 naïve) | 6 months              | Loss of response: 3 months (total 60 switch/2 naïve): 27.7% 6 months (total 29 switch/2 naïve): 30% | N/A                  | N/A                          | Only mild AEs reported |
| Hamanaka S et al., 2016, Japan | N = 3 | Switch to NK (n = 2 CD)              | 24 weeks              | Sustained remission rate: 100% | N/A                  | N/A                          | N/A                              |
Table 4 (continued)

| Study                           | N     | Intervention                               | Duration of follow-up | Effectiveness of medication                          | Discontinuation rate | Immunogenicity                     | Safety                      |
|---------------------------------|-------|--------------------------------------------|-----------------------|------------------------------------------------------|----------------------|------------------------------------|-----------------------------|
| Hlavaty T et al., Slovak Republic | N = 12| Switch to CT-P13 (n = 12)                  | Up to 48 weeks       | Loss of response: 24 weeks: 0% (0/12)                | N/A                  | N/A                                | Developed psoriasiform dermatitis and discontinued treatment |
| Sieczkowska J et al., Poland    | N = 16| Switch to IFX biosimilar                   | 5 infusions           | N/A                                                  | N/A                  | N/A                                | 100% (15/15) of patients maintained therapeutic drug levels; 26.7% (4/15) had ADA > 2 ng/mL |
| Kierkus J et al., Poland        | N = 32| Switch to CT-P13                           | 8 weeks               | PCDAI (mean [range]): At switch: 8.5 (0–35)         | N/A                  | N/A                                | No change in the incidence of mild AEs |
| Suk JY et al., Republic of Korea| N = 42| Switch to CT-P13                           | ≥1 infusion           | Loss of response: CD: 14% UC: 33%                    | N/A                  | N/A                                | N/A                         |

ADA antidual antibody, AE adverse event, CD Crohn's disease, FCD fistulizing Crohn's disease, HBI Harvey-Bradshaw Index, IBD-u unclassified inflammatory bowel disease, IFX infliximab originator, IRR infusion-related reaction; N/A not available, NK Nippon Kayaku/Celltrion infliximab biosimilar (Japan), PCDAI Pediatric Crohn's Disease Activity Index, PM partial Mayo score, PUCAI Pediatric Ulcerative Colitis Activity Index, SAE severe adverse event, SCCAI Simple Clinical Colitis Activity Index, SEM standard error of mean, TEAE treatment-emergent adverse event, TESAE treatment-emergent serious adverse event, TNFi tumor necrosis factor inhibitor, UC ulcerative colitis

Data on long-term outcomes are usually scarce and typically limited to one year following non-medical switch. Studies involving long-term follow-up of patients who have undergone a non-medical switch (NM-S) have been published more recently. Some studies have presented data at the 2018 Congress of ECCO on the effectiveness and safety of switch from infliximab to CT-P13 in patients with IBD [26]. More recently, Smits et al. examined the long-term efficacy, safety, and immunogenicity profile of a non-medical switch from infliximab to CT-P13 in patients with IBD [26]. Data from a single-center, prospective, open-label study consisted of 57 patients with CD, 24 UC, and 2 "IBD unclassified" patients, who were initially followed for 2 weeks post-switch. No control arm was used in this study. The primary endpoint was a change in disease activity scores at week 52 compared to baseline, as measured by HBI for CD or Simple Clinical Colitis Activity Index (SCCAI) for UC and unclassified IBD. Remission was defined as an HBI < 15 for CD, < 4 for UC, and < 5 for IBD-u. At week 52, 85% of patients with CD (56/66), 78% of patients with UC (67/87), and 81% of patients with IBD-u (52/65) were in remission, and 80% (53/66, 67/87, 52/65) were in remission as of week 52 compared to baseline. In this study, the authors also examined the potential impact of non-medical switch on the efficacy and safety of CT-P13 in patients with IBD [26]. Patients enrolled in this study were anti-TNFα naïve patients (n = 311), 139 had previous exposure to one or more biologics (median duration of drug holiday was 9 months for infliximab and 10 months for other anti-TNFs), and 97 were switched from infliximab to CT-P13 therapy (hereafter referred to as switch patients). No control arm was utilized for this study. Among switch patients, 53 of 79 (68%) patients remained on CT-P13 at week 52, while 24 UC patients from 31 referral centers treated with CT-P13 were enrolled. After a median of 6 months of follow-up, there were 94.5%, 90.8%, and 78.9%, respectively, for the switch patients (log-rank test; p = 0.005). The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients. The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients. The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients. The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients. The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients. The majority of these patients were anti-TNFα naïve; 12% were CD, 24 UC, and 2 "IBD unclassified" patients.
Median trough levels at baseline and weeks 52 and 104 were 3.6 μg/ml (interquartile range [IQR] 1.7–5.5), 3.7 μg/ml (IQR 2.1–5.8), and 3.9 μg/ml, respectively (IQR 2.2–5.7; \( p = 0.664 \)). Antidrug antibodies (ADAs) were present in 5 of 83 (6%) patients at baseline (prior to switching) and in 2 (2%) patients before week 52. No subsequent ADAs were detected up to week 104. The majority of IBD patients (68%) continued CT-P13 beyond 2 years after switching from innovator infliximab [57].

Discussion

We conducted a systematic review of randomized controlled trials, observational studies, and case series, in which stable patients with UC or CD were switched from originator infliximab to the CT-P13 biosimilar. In total, 49 reports from 44 distinct studies available up to April 2018 were included, expanding the literature examined on infliximab biosimilar switch presented over the last few years [59–63]. Given the paucity of published data from randomized controlled trials, most systematic reviews on infliximab biosimilar switch have referenced a limited number of observational studies [62, 63]. Some publications have suggested that the current data support the safety of switching from originator infliximab to biosimilar in IBD patients [59, 60], whereas others have emphasized the need for studies on bidirectional biosimilar switch, as well as those that use appropriate control arms [61]. We present a comprehensive review evaluating the impact of switching to CT-P13 on efficacy/effectiveness and safety in CD and UC patients. In addition, this systematic review critically appraises methodologies of the studies generating the data that may influence the decision of switching stable patients from infliximab to CT-P13, and highlights their limitations.

Only 3 of the studies included in this systematic review were found to be randomized controlled trials, with NOR-SWITCH being the only trial fully published to date. Designed as a pragmatic real-life study, NOR-SWITCH examined treatment switch across 6 indications in approximately 500 patients (CD, \( n = 155 \); UC, \( n = 93 \); SpA, \( n = 91 \); RA, \( n = 78 \); PsO, \( n = 35 \); and PsA, \( n = 30 \)). Limitations of the study design have been previously discussed [64]. For example, a composite measure of disease worsening was used, which included 6 highly heterogeneous inflammatory disease states each employing different disease activity measures, varying infliximab dosing regimens, and concomitant therapies. This, in addition to patient inclusion being largely based on a “clinical diagnosis” of stable disease with ill-defined disease-specific criteria, introduces the potential for bias and limits the inferences that can be extrapolated to IBD. The primary endpoint of disease worsening was also estimated to occur in 29.6% of patients treated with CT-P13 versus 26.2% on infliximab therapy. As the total enrollment number allowed for a 15% non-inferiority margin at 90% power, theoretically, 50% of CT-P13-treated patients could experience disease worsening without the primary endpoint being deemed inferior. Finally, while subgroup efficacy analyses were underpowered to detect differences, the trend toward increased disease worsening in CD patients has caused some concern in the gastrointestinal community and a desire for more robust data in IBD patients.

In order to address the potential concerns and limitations of NOR-SWITCH, Kim et al. conducted a double-blind study that enrolled a homogenous patient population with active CD, utilized appropriate control arms, and allowed for dose escalation [13]. It was the first randomized controlled trial that examined the efficacy and safety of a one-way switch from CT-P13 to infliximab in this patient population and investigated a number of secondary endpoints (e.g., pharmacokinetics, safety, ADA development, and quality of life). Randomization of 220 patients across 4 switch arms results in approximately 55 patients per arm (not accounting for attrition). Assuming a power level of 80% and \( p < 0.05 \) for statistical significance, the margin of difference would be approximately 20% at 6 weeks. A more stringent non-inferiority margin of 5–10% would require a larger sample size than the 220 patients included in this trial. This study also utilized CDAI-70 response at week 6 as the primary endpoint [13]. Standard CDAI assesses remission, not response, typically being measured at weeks 8 or 12, and typically a more stringent definition of CDAI-100 is used for response. It is unclear whether 6 weeks is a sufficient duration for a response signal to manifest and if the shorter duration to primary endpoint creates a bias toward demonstrating non-inferiority.

Volkers et al. are currently conducting the first phase IV, double-blind, randomized, non-inferiority clinical trial comparing IBD patients who were switched from infliximab to CT-P13 [14]. Inclusion criteria required patients to be in clinical remission (HBI < 5 and Mayo < 2) and have a fecal calprotectin < 250 mg/g. The primary endpoint is the number of patients in remission at week 30. Though utilization of the HBI and Mayo questionnaires to determine remission may facilitate the feasibility of a phase IV trial, it does not allow for endoscopic verification of remission, particularly when switching stable patients.

Real-world data can offer valuable insight into the clinical effectiveness and safety of a therapy, prescribing patterns, and quality of life outcomes. The major disadvantage of these studies is the lack of randomization or appropriate control arms. Efficacy and safety assessments are also unlikely to be as robust as those in randomized controlled trials. Though many patients in published observational switch studies appeared to experience disease control, uncontrolled variables may influence perceived
drug–outcome correlations, and models may not entirely account for confounding factors, rendering therapies to be seemingly equivalent. This underscores the caution that must be taken when switching stable IBD patients. A recent meta-analysis attempted to address these concerns by examining published observational switch studies and analyzing data by disease type and duration of follow-up. Due to the heterogeneity of the patient populations, each analyzed group was comprised of only 1–2 studies, making inferences of the analysis difficult to assess [62]. The lack of published data has also been a limiting factor for meta-analyses. For instance, among the observational studies included in this systematic review, only half had been published by the time of submission. Unfortunately, this leads to a narrow understanding of important study details, the knowledge of which could affect readers’ perception of study results.

The present systematic review has several limitations, including the small number of randomized controlled trials; predominance of observational studies with varying outcome assessments, lack of appropriate controls, and absence of data on loss of response and treatment discontinuation; and scarcity of research on long-term effects of biosimilar switch in CD and UC patients. The inclusion of conference abstracts can also be considered a limitation. While their inclusion may help to minimize publication bias, conference abstracts do not provide a complete description of methods and results to allow an evaluation of the rigor in which the research was conducted. In addition, the risk of bias for individual studies used in this review could not be assessed, as this would be subjective due to the biased nature of observational studies, which make up the majority of the studies included. Since this review mainly provides a critique of the methodologies used to generate the majority of data on biosimilar switch from infliximab to CT-P13, the lack of risk assessment does not affect its conclusions.

To date, the totality of evidence would suggest that switching infliximab patients to CT-P13 is generally safe and effective for most individuals. It remains unclear what the long-term impact is on those who are negatively affected by non-medical switching, the role that immunogenicity may play, and what physicians are willing to consider an acceptable risk.

**Impact of Non-medical Switch on Immunogenicity**

ADAs are a common occurrence with anti-TNFα therapies [65]. Indeed, the pivotal trials from a number of anti-TNFα products have demonstrated that the rate of ADAs ranges from approximately 1 to 30% [2, 66–69]. Despite the propensity of ADA development, there are a number of challenges in terms of their quantification and impact assessment.

A recent European Medicines Agency guideline stated that ADA assays may often bind to the therapeutic product itself and thus require significant modifications to accommodate for inaccurate signals. However, such modifications can decrease the sensitivity and accuracy of the assay [70]. It has also been suggested that comparative assessment of immunogenicity be comprised of more than incidence rates between originator and biosimilar and include titers and titer distribution [71]. These factors, combined with the inherent differences within patient populations, hamper the comparability of ADAs across clinical trials.

Above all, the precise factors that cause an immunogenic response in patients are likely due to various treatment-, product-, and patient-related considerations [72, 73]. Gonczi et al. demonstrated the potential impact of drug holidays on those using infliximab. CD patients exposed to infliximab over 12 months prior to initiating CT-P13 therapy demonstrated significantly reduced clinical response and remission up to week 54 (p < 0.005) and higher ADA levels in early treatment (week 2, p < 0.001). Previously treated UC patients demonstrated significantly reduced clinical remission (p ≤ 0.03) and ADA development up to week 6 (p < 0.02) versus infliximab-naïve UC patients [74]. No differences were observed at weeks 14 or 30. Final one-year results of this study confirmed the clinical impact of previous infliximab exposure and highlighted the increased likelihood to develop infusion reactions [75, 76].

To date, real-world infliximab switch studies in IBD do not have robust or long-term ADA assessments and are not powered to detect significant differences in immunogenicity. The lack of clarity regarding immunogenicity, combined with the inherent variability in ADA assays and loss of response rates observed thus far, reinforces a need for high-quality studies.

**Potential Multiple Switch Scenarios**

A number of infliximab biosimilars are likely to gain market authorization over the next few years. Several regulatory agencies have started to establish more stringent guidelines for demonstrating interchangeability between a reference drug and its biosimilar [77]. However, with very limited data on single reverse switches, and no trials examining multiple switches with infliximab, healthcare practitioners are currently ill-informed of the ADA risks and complications that may arise from frequent non-medical switch due to preferential listing or product tendering.

It is prudent that regulatory bodies take appropriate steps to ensure patient safety in this changing environment, as the existence of multiple infliximab biosimilars would make clinical trials extremely complex, and examination of all product-switch permutations unfeasible. Moreover, the increase in untested exposure to potentially different
epitopes of infliximab and its biosimilars (particularly if the structures diverge over time), combined with the variability in patient-related factors, is likely to increase the unpredictability of immune responses [64].

Conclusion

Despite the potential limitations of the NOR-SWITCH trial, it is currently the only published randomized controlled trial available to glean insights regarding the non-medical switch. Considering this paucity of well-controlled data, the gastroenterology community has largely based clinical decisions on evidence from heterogeneous patient populations in observational trials, which lack controls. Though the majority of these studies suggest that a non-medical switch is safe and effective for most individuals, healthcare professionals and regulatory bodies should be aware of the limitations of study designs when making inferences about the risks and benefits of switching stable IBD patients. Full publication of additional randomized controlled trials in IBD populations will be critical to further our understanding of the impact that such clinical or economic decisions will have on patients.

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Compliance with Ethical Standards

Conflict of interest

Edmond-Jean Bernard has served as a speaker and consultant for Janssen, Abbvie, Takeda, Shire, Ferring, Allergan, Pfizer, and Merck. Richard N. Fedorak has served as a speaker and consultant for AbbVie Canada, Inc. and Janssen Canada, Inc. Vipul Jairath has received consulting fees from AbbVie, Eli Lilly, GlaxoSmithKline, Arena pharmaceuticals, Genetech, Pendopharm, Sandoz, Merck, Takeda, Janssen, Robarts Clinical Trials, Topivert, and Celltrion, and speaker’s fees from Takeda, Janssen, Shire, Ferring, Abbvie, and Pfizer.

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