Comparison of the efficacy and risk of discontinuation between non-TNF-targeted treatment and a second TNF inhibitor in patients with rheumatoid arthritis after first TNF inhibitor failure

Dong-Jin Park, Sung-Eun Choi, Ji-Hyoun Kang, Kichul Shin, Yoon-Kyoung Sung and Shin-Seeok Lee

Abstract
Objectives: Despite improved care for rheumatoid arthritis (RA) patients, many still experience treatment failure with biologic disease-modifying antirheumatic drugs (bDMARDs) or targeted synthetic DMARDs (tsDMARDs; typically Janus kinase inhibitors [JAKi]), and eventually switch to other agents. We compared the efficacy of a second tumor necrosis factor inhibitor (TNFi) and non-TNF-targeted treatment as the second-line treatment in patients showing an insufficient response to the first TNFi.

Methods: Patients were included if they had received at least one prescription for a TNFi, and at least one follow-up prescription for a second TNFi or non-TNF-targeted treatment after discontinuation of the first drug. In total, 209 patients were analyzed, including 69 with a second TNFi and 140 with a non-TNF-targeted treatment (106 non-TNF biologics and 34 JAKi). Cox regression was used to estimate the hazard ratio (HR) for discontinuation.

Results: The mean follow-up period after switching was 28.0 (range: 0–80) months and 24.4% of the 209 patients switched or discontinued the second drug. In multivariate Cox proportional hazard analysis, the non-TNF-targeted treatment group had a lower likelihood of discontinuing their treatment than the second TNFi group [HR = 0.326, 95% confidence interval (CI): 0.170–0.626, \( p = 0.001 \)]. When analyzed separately, the risk of discontinuation was significantly lower in both the non-TNF biologic (HR = 0.318, 95% CI: 0.160–0.633, \( p = 0.001 \)) and JAKi (HR = 0.356, 95% CI: 0.129–0.980, \( p = 0.046 \)) groups than in the second TNFi group.

Conclusion: Our study supported switching to a non-TNF-targeted treatment instead of TNF cycling in patients with RA showing an inadequate response to initial TNFi.

Keywords: JAK inhibitor, rheumatoid arthritis, switching, treatment continuation, TNF inhibitor

Introduction
Rheumatoid arthritis (RA) is one of the most common autoimmune inflammatory diseases, and is characterized by pathological synovial hypertrophy, joint inflammation, and structural damage.\(^1,2\) RA has an incidence of up to 2% worldwide and often causes substantial functional impairment and decreased health-related quality of life relative to the general population.\(^3,4\) Although conventional synthetic disease-modifying antirheumatic drugs (csDMARDs), including methotrexate (MTX), remain the first-line therapy for RA, considerable advances have been made in the treatment of RA over the last few decades. The development of targeted treatments, such as biologic disease-modifying antirheumatic
drugs (bDMARDs) and targeted synthetic DMARDs (tsDMARDs), represented by Janus kinase inhibitors (JAKis), showed a therapeutic revolution in the treatment of RA.5,6

Tumor necrosis factor inhibitors (TNFis) remain the first-line targeted treatment following csDMARD therapy failure,7–9 as these agents are effective in reducing the signs and symptoms of RA and inhibiting the progression of structural joint damage.10,11 Despite the marked treatment effect of TNFi, 40–50% of RA patients discontinue their first TNFi within 3 years of initiation.12,13 For patients discontinuing their first TNFi due to a lack of efficacy or intolerance, both cycling to another TNFi (cycling strategy) or switching to other targeted agent with a different mechanism of action (MOA) (swap strategy) may be considered as alternative strategies.7,8 An observational cohort study reported similar outcomes between RA patients who cycled to another TNFi and those who initiated non-TNFi biologic agent with a different MOA (abatacept).14 A retrospective study reported that switching from first TNFi (infliximab) to either another MOA bDMARD (tocilizumab) or second TNFi (etanercept) in patients with RA showed no significant difference in efficacy, as measured by disease activity.15 However, randomized controlled trials suggested that switching to non-TNF biologic agents, such as rituximab, tocilizumab, and abatacept, could be an alternative option for RA patients with an inadequate response to one or more TNFIs.16–18 Moreover, an observational study reported that, after failure of a first TNFi for RA, switching to a new MOA bDMARD (rituximab) was more effective than cycling to another TNFi.19 Furthermore, another randomized trial showed that a swap strategy using non-TNF biologic agents may be more effective than use of a second TNFi in patients with an insufficient response to the first TNFi.20 With this background, the 2021 American College of Rheumatology (ACR) guideline conditionally recommended a swap strategy over cycling to another TNFi in patients who showed an insufficient response to a previous TNFi.21 However, as this recommendation is based on very low-certainty evidence, the therapeutic choice after failure of the first TNFi is largely dependent on the experience of the treating physician or the patient’s preference.21

Therefore, further studies are needed to better understand the outcomes of either cycling to another TNFi or switching to non-TNF-targeted treatment, such as a different MOA bDMARD or JAKi. Using data from the KORean nationwide BIOlogics and targeted therapy (KOBIO) registry, a nationwide real-world prospective cohort to assess outcomes of RA patients treated with any targeted treatment,8 this study was performed to compare the effectiveness of the cycling and swap strategies in terms of the drug discontinuation rate, and to clarify the predictors of discontinuation of second-line treatment in RA patients who discontinued their prior TNFi.

**Patients and methods**

*Study design and population*

This study was performed using data from the KOBIO registry, a nationwide multicenter, hospital-based observational registry maintained by the Korean College of Rheumatology (KCR). The aim of the registry was to prospectively assess the clinical manifestations and outcomes, including adverse events, of RA patients who had received any targeted treatment, such as bDMARDs or tsDMARDs.8 All patients eligible for the study were classified as having RA by their treating rheumatologist, and fulfilled the 2010 ACR criteria for RA.2 The RA patients were enrolled from 47 tertiary academic and community rheumatologic centers across the country and had follow-up assessments at approximately 12-month intervals.8 For this study, subjects were identified from the baseline and follow-up data of the KOBO registry.

Although the decision to switch targeted treatment with a certain drug or certain MOA was made solely by the treating rheumatologist, these decisions were based on the Korean National Health Insurance (KNHI) reimbursement criteria. The criteria for initiation and maintenance of targeted agents, including bDMARDs and JAKis, are summarized in Supplementary Table 1. If the maintenance criteria are met after 6 months of a targeted agent, that drug can be reimbursed for an additional 6 months. If the maintenance criteria are not satisfied or serious adverse events occur, treatments should be switched to other agents.

A total of 2356 RA patients receiving a bDMARD or tsDMARD were enrolled in the KOBIO registry from December 2013 to November 2020 (Figure 1). Of these, RA patients who did not
switch their first targeted treatment \( (n=1499) \) or multiple switcher \( (n=319) \), or those who were lost to follow-up \( (n=178) \) were excluded. Among patients switching their first targeted therapy, those receiving non-TNFα treatment as the first targeted therapy \( (n=130) \) and those with missing follow-up data \( (n=21) \) were also excluded. Finally, a total of 209 patients were analyzed in this study, consisting of 69 patients in the second TNFα group and 140 in the non-TNF-targeted treatment group (106 in the non-TNFi biologic group and 34 in the JAKi group). Patients were followed up at least once from the time of initiation of the subsequent TNFα or non-TNF-targeted treatment until discontinuation. This study adhered to all relevant principles of the Declaration of Helsinki. The study protocol and data collection forms were approved by the institutional review board or local ethics committee of all participating institutions, including that of Chonnam National University Hospital (Approval No. CNUH-2012-239). All participants provided written informed consent for enrollment in the KOBIO registry. The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement guidelines for reporting observational studies were followed.22

**Data collection**

All patient data were transferred, by individual investigators, into the KOBIO web server (http://www.rheum.or.kr/kobio/). RA patients were interviewed using a structured questionnaire that captured sociodemographic data and concomitant medications. The following data were collected: age, sex, disease duration, education level, smoking status, blood pressure, body mass index (BMI), presence of hypertension and diabetes mellitus, and laboratory findings, such as the erythrocyte sedimentation rate (ESR), C-reactive protein (CRP), rheumatoid factor (RF), and anticyclic citrullinated peptide (CCP) antibody. Radiographs of the hands and feet were also obtained to evaluate erosion and joint space narrowing of these joints at the time of enrollment.
Disease activity was evaluated using validated composite measures, and physical examinations were based on the clinical finding of one or more tender and swollen joints (44 joints). The joint assessments were performed by trained investigators at each institution. The results of 10-cm visual analog scales (VAS) for patient global assessment (PGA) and physician global assessment (PhGA) were recorded. Quantitative assessments of RA disease activity, such as Disease Activity Score of 28 joints (DAS28), were evaluated at the time of the initiation of the second TNFi or non-TNF-targeted treatment. Following 1 year of treatment, achievement of remission or low disease activity (LDA) was also evaluated.

Concomitant csDMARD use was also recorded after initiation of the second TNFi or non-TNF-targeted treatment. Use of csDMARDs was determined based on any use of oral or subcutaneous MTX, sulfasalazine, hydroxychloroquine (HCQ), leflunomide, tacrolimus, or cyclosporine. The first TNFi was also recorded and reasons for switching were classified into the following mutually exclusive categories: inefficacy, adverse events, or other (such as patient preference, financial issues, and concerns regarding safety or comorbidity).

The primary outcome was the discontinuation of second TNFi or non-TNF-targeted bDMARD/tsDMARD in patients with RA who discontinued their first TNFi. In addition, the change in DAS28-ESR score between baseline and 1 year, and the proportion of patients who achieved DAS28-ESR remission or LDA at 1 year, were obtained to assess the effectiveness of targeted therapies. Predictors of discontinuation of the second-line targeted treatment in RA patients were also evaluated.

Statistical analysis
Descriptive statistical analyses were performed. Values are shown as means ± standard deviation (SD) or percentages. Data were analyzed using the χ² test for categorical variables and Mann–Whitney U test or one-way analysis of variance (ANOVA) for continuous variables. The cut-off value for LDA was defined as DAS28 ≤ 3.2 and that for clinical remission was defined as DAS28 < 2.6. Kaplan–Meier curves were used to examine the duration of treatment, and the log-rank test was used to compare drug continuation between the TNFi cycling and new MOA non-TNF-targeted swapping groups. Multivariable Cox proportional hazards models were used to evaluate potential predictors of drug discontinuation. Variables significant at p < 0.5 in univariable analysis, along with age, sex, disease duration, and concomitant use of csDMARDs, were included in multivariable analysis to evaluate predictors of discontinuation of second-line targeted treatment. Furthermore, subgroup analyses were performed to determine values for predicting discontinuation of the second TNFi drug and non-TNF-targeted treatment. In these analyses, multivariable Cox regression analysis was performed including variables significant at p < 0.5 in univariable analysis, that is concomitant DMARDs, age, sex, and disease duration. Multivariable analysis used a backward inclusion methodology, and hazard ratio (HRs), 95% confidence interval (CIs), and p-values were used to interpret the results. In all analyses, p < 0.05 was taken to indicate statistical significance; the Bonferroni correction was applied when performing multiple comparisons of the 1-year treatment responses among the three groups. Statistical analyses were performed using SPSS for Windows software (ver. 21.0; SPSS Inc., Chicago, IL, USA).

Multiple imputation
The multiple imputations method was applied to address missing baseline data, using the Estimation Maximization (EM) algorithm; five imputed datasets were thus created. The complete variables included age at commencement, gender, disease duration, BMI, smoking, diabetes mellitus, hypertension, educational status, duration of TNFi treatment, reason for discontinuing first TNFi treatment, tender and swollen joint counts, PGA and PhGA levels, and elevated ESR/CRP and DAS28-ESR/CRP scores. Missing variables (number of missing data items) included erosion or joint space narrowing evident in X-ray (n = 65), RF positivity (n = 3), and anti-CCP positivity (n = 27).

Results
The baseline characteristics of the patients are shown in Table 1. At the time of enrollment, the mean age was 46.3 ± 12.9 years, and most of the patients were women (85.2%; n = 178). The mean time since initial diagnosis of RA (disease duration) was 71.1 ± 77.9 months. Of the total of 209 patients with RA, 69 were second TNFi cyclers (second TNFi group) and 140 comprised the non-TNF-targeted treatment group. The
Table 1. Demographic and clinical features of RA patients receiving second-line targeted treatments.

|                                | All patients n = 209 | Second TNFi group n = 69 | Non-TNFi biologic group n = 106 | JAKi group n = 34 | p-value |
|--------------------------------|----------------------|--------------------------|---------------------------------|------------------|---------|
| Age at commencement, years    | 46.3 ± 12.9          | 45.7 ± 12.1              | 46.9 ± 12.2                     | 45.3 ± 16.4      | 0.688   |
| Men                            | 31 [14.8]            | 8 [11.6]                 | 16 [15.1]                       | 7 [20.6]         | 0.479   |
| Disease duration, months       | 71.1 ± 77.9          | 70.7 ± 63.4              | 71.3 ± 85.4                     | 71.4 ± 82.2      | 0.447   |
| BMI, kg/m²                      | 22.3 ± 3.10          | 22.8 ± 3.25              | 22.3 ± 3.80                     | 22.3 ± 3.09      | 0.715   |
| Current smoker                 | 14 [6.7]             | 5 [7.2]                  | 8 [7.5]                         | 1 [2.9]          | 0.630   |
| Diabetes mellitus              | 27 [12.9]            | 9 [13.0]                 | 16 [15.1]                       | 2 [5.9]          | 0.378   |
| Hypertension                   | 53 [25.4]            | 18 [26.1]                | 26 [24.5]                       | 9 [26.5]         | 0.961   |
| Education, years               | 12.0 ± 3.79 [n = 202]| 11.7 ± 3.88              | 12.0 ± 3.72 [n = 103]           | 13.1 ± 3.79 [n = 30]| 0.281   |
| Duration of first TNFi treatment, months | 12.3 ± 10.7 | 14.2 ± 12.6 | 10.4 ± 9.13 | 14.2 ± 10.55 | 0.072   |
| First TNFi                     |                      |                         |                                 |                  | 0.004   |
| Etanercept                     | 58 [27.8]            | 25 [36.2]                | 27 [25.5]                       | 6 [17.6]         |         |
| Infliximab                     | 70 [33.5]            | 28 [40.6]                | 35 [33.0]                       | 7 [20.6]         |         |
| Adalimumab                     | 64 [30.6]            | 13 [18.8]                | 37 [34.9]                       | 14 [41.2]        |         |
| Golimumab                      | 17 [8.1]             | 3 [4.3]                  | 7 [6.6]                         | 7 [20.6]         |         |
| Reason for discontinuing first TNFi |                  |                         |                                 |                  | 0.205   |
| Inefficacy                     | 147 [70.3]           | 44 [63.8]                | 78 [73.6]                       | 25 [73.5]        |         |
| Adverse events                 | 55 [26.3]            | 20 [29.0]                | 27 [25.5]                       | 8 [23.5]         |         |
| Other                          | 7 [3.3]              | 5 [7.2]                  | 1 [0.9]                         | 1 [2.9]          |         |
| Use of concomitant csDMARDs    | 188 [90.0]           | 66 [95.7]                | 90 [84.9]                       | 32 [94.1]        | 0.047   |
| Use of concomitant corticosteroids | 182 [87.1]       | 60 [87.0]                | 94 [88.7]                       | 28 [82.4]        | 0.632   |
| Erosion or joint space narrowing on X-ray | 85/144 [59.0]    | 28/43 [65.1]             | 44/75 [58.7]                    | 13/26 [50.0]     | 0.463   |
| RF positivity                  | 177/206 [85.9]       | 54/67 [80.6]             | 93/106 [87.7]                   | 30/33 [90.9]     | 0.281   |
| Anti-CCP positivity            | 162/182 [89.0]       | 46/59 [78.0]             | 91/96 [94.8]                    | 25/27 [92.6]     | 0.004   |
| Swollen joint count (44 joints)| 5.23 ± 6.84          | 5.72 ± 8.77              | 5.08 ± 5.80                     | 4.71 ± 5.33      | 0.573   |
| Tender joint count (44 joints) | 6.45 ± 7.67          | 6.38 ± 9.16              | 6.37 ± 6.82                     | 6.85 ± 7.10      | 0.266   |
| PGA                            | 5.79 ± 2.58          | 5.00 ± 2.58              | 6.00 ± 2.63                     | 6.74 ± 2.59      | 0.004   |
| PhGA                           | 5.53 ± 2.55          | 4.88 ± 2.39              | 5.65 ± 2.71                     | 6.44 ± 2.02      | 0.008   |
| Elevated ESR                  | 157 [75.1]           | 51 [73.9]                | 78 [73.6]                       | 28 [82.4]        | 0.566   |
| Elevated CRP                   | 135 [64.6]           | 40 [58.0]                | 71 [67.0]                       | 24 [70.6]        | 0.346   |
| DAS28-ESR⁺                     | 4.76 ± 1.72          | 4.45 ± 1.86              | 4.84 ± 1.66                     | 5.12 ± 1.51      | 0.181   |
| DAS28-CRP⁺                     | 4.13 ± 1.57          | 3.78 ± 1.68              | 4.27 ± 1.51                     | 4.39 ± 1.45      | 0.062   |

Anti-CCP, anti-cyclic citrullinated peptide; BMI, body mass index; CRP, C-reactive protein; csDMARDs, conventional synthetic disease-modifying antirheumatic drugs; DAS28, Disease Activity Score-28; ESR, erythrocyte sedimentation rate; JAKi, Janus kinase inhibitor; PGA, patient global assessment; PhGA, physician global assessment; RF, rheumatoid factor; TNFi, tumor necrosis factor inhibitor. Except where otherwise indicated, data are shown as n (%) or mean ± standard deviation. *Missing data were excluded from the analyses.
non-TNF-targeted treatment group was further subdivided into two subgroups: non-TNFi biologic switcher (non-TNFi biologic group; 106 patients) and JAKi switcher (JAKi group; 34 patients) groups. There were no significant differences in age or sex distribution among the three groups. In all patients, infliximab (33.5%) was the most commonly prescribed first TNFi, followed by adalimumab (30.6%), etanercept (27.8%), and golimumab (8.1%). In the second TNFi group, infliximab (40.6%) was the most common first TNFi drug, but adalimumab was the most commonly prescribed drug in both the non-TNFi biologic and JAKi groups (34.9% and 41.2%, respectively); these differences were statistically significant ($p < 0.004$). The most commonly reported reason for stopping prior TNFi was inefficacy (70.3%), followed by adverse events (26.3%), and other miscellaneous causes (mostly patient choice due to infection-related problems, family planning, or financial issues). There were no significant differences in the reason for discontinuation of the first TNFi among the groups. The rate of concomitant csDMARD use after switching was significantly lower in the non-TNFi biologic group (84.9%) than the TNFi and JAKi groups (95.7% and 94.1%, respectively) ($p = 0.047$).

With regard to clinical features, the TNFi group had significantly lower anti-CCP positivity than the non-TNFi biologic and JAKi groups (78.0%, 94.8%, and 92.6%, respectively) ($p = 0.004$). With regard to disease activity at the time of switching, both PGA and PhGA were higher in the JAKi group (6.74 ± 2.59 and 6.44 ± 2.02, respectively) than the TNFi group (5.00 ± 2.58 and 4.88 ± 2.39, respectively) and non-TNFi biologic group (6.00 ± 2.63 and 5.65 ± 2.71, respectively) ($p = 0.004$ and $p = 0.008$, respectively). Although both DAS28-ESR and DAS28-CRP tended to be higher in the JAKi group than in both the second TNFi group and non-TNFi biologic group, the differences were not statistically significant (all $p$s $> 0.05$). Furthermore, there were no significant group differences in the 44 tender-swollen joint count, 44 swollen joint count, levels of ESR and CRP, or presence of erosion or joint space narrowing on radiographic findings.

Table 2 shows a comparison of the response to treatment at 1 year among the groups. At the 1-year follow-up, the rate of continuation of their targeted treatment was significantly higher in the non-TNFi biologic group (80.2%) than the second TNFi group (62.3%) and JAKi group (76.5%) ($p = 0.030$). Furthermore, the proportion of patients who achieved remission or LDA was higher in the non-TNFi biologic group (72.9%) than the second TNFi group (46.5%) and JAKi group (44.0%) ($p = 0.003$). In addition, DAS28-ESR at 1 year was lower in the non-TNFi biologic group (2.79 ± 1.14) than the second TNFi group (3.37 ± 1.18) and JAKi group (3.24 ± 0.88) ($p = 0.015$). The change in DAS28-ESR score between baseline and 1 year was significantly greater in the non-TNFi biologic group (2.01 ± 2.00) and JAKi group (2.08 ± 1.47) than the second TNFi group (0.95 ± 1.63) ($p = 0.006$). However, the delta DAS28-ESR was not different between the non-TNFi biologic and JAKi groups ($p > 0.05$).

The mean duration of follow-up after drug switching was 28.0 (range: 0–80) months. Overall, 156 (74.6%) patients continued their second targeted treatment, while 53 (24.4%) switched to another drug or discontinued treatment. In detail, 46 (66.7%) patients in the TNFi group, 83 (78.3%) in the non-TNFi biologic group, and 27 (77.4%) in the JAKi group continued their drug. Kaplan–Meier analysis showed that the duration of treatment after drug switching was significantly longer after non-TNF-targeted treatment swapping compared to second TNFi cycling ($p = 0.015$) [Figure 2(a)]. In fact, the second TNFi group showed the lowest rate of drug continuation among the three groups ($p = 0.049$), while there were no statistically significant differences in the duration of treatment between the non-TNFi biologic and JAKi groups ($p > 0.05$) [Figure 2(b)]. Although the data are not shown, there were no significant differences in rates of continuation between the different MOA biologic agents in the non-TNFi biologic group ($p > 0.05$).

The results of univariable and multivariable Cox proportional hazard analyses for discontinuation of targeted treatments used after TNFi failure are presented in Table 3. In multivariable Cox proportional hazard analysis adjusted for baseline demographic and clinical characteristics, patients with non-TNF-targeted treatments, including the non-TNFi biologic and JAKi groups, had a lower risk of discontinuation of treatment than the second TNFi group (HR = 0.326, 95% CI: 0.170–0.626, $p = 0.001$). Even when analyzed separately, the risk of discontinuation was significantly lower in both the non-TNFi biologic group (HR = 0.318,
95% CI: 0.160–0.633, \( p = 0.001 \) and JAKi group (HR = 0.356, 95% CI: 0.129–0.980, \( p = 0.046 \)) than in the second TNFi group. Furthermore, after the inclusion of first TNFi and PGA, both of which had a \( p \)-value < 0.05 (Table 1), the results of the multivariable Cox regression analysis were unchanged.

BMI, Further subgroup analyses were performed to determine the predictor of discontinuation of the second TNFi drug and non-TNF-targeted treatment (Table 4). We found that anti-CCP positivity (HR = 0.273, 95% CI: 0.092–0.809, \( p = 0.019 \)) was an independent predictor of discontinuation of the second TNFi drug, while use of concomitant csDMARDs (HR = 0.131, 95% CI: 0.027–0.634, \( p = 0.011 \)) was a significant predictor of discontinuation of non-TNF-targeted treatment.

TNFi, Discussion

In this real-world analysis of patients with RA, we found that switching to non-TNF-targeted treatment (both a non-TNFi biologic agent and JAKi) was associated with a lower risk of discontinuation of treatment and a better 1-year treatment response than cycling to a second TNFi. In addition, anti-CCP positivity was significantly associated with a lower risk of discontinuation of the second TNFi, while use of concomitant csDMARDs was associated with a lower risk of discontinuation of non-TNF-targeted treatment.

Our study was generally consistent with previous reports showing that switching to non-TNF-targeted drugs was more beneficial than cycling to a second TNFi. To date, observational and registry studies have suggested that RA patients swapping to non-TNFi treatments, such as rituximab, abatacept, tocilizumab, or tofacitinib, following an inadequate response to a first TNFi, showed a favorable outcome compared to those cycling to subsequent TNFis.\(^{14,25,26}\) In an Italian study, Favalli \textit{et al.}\(^{26}\) showed that RA patients swapping to different MOA bDMARDs (abatacept, rituximab, or tocilizumab) had a higher retention rate (HR = 2.258, 95% CI: 1.507–3.385) than those cycling to a second TNFi. Moreover, real-world data from the United States showed that during a median follow-up of 29.9 months, TNFi cyclers were more likely to be non-persistent (HR = 1.511, 95% CI: 1.196–1.908, \( p = 0.001 \)) than new non-TNFi switchers, and switching to a non-TNF-targeted treatment (abatacept, anakinra, rituximab, tocilizumab or tofacitinib) showed a trend toward greater disease activity reduction.\(^{27}\) A recent US study of
large-scale claim-based data also found that, although the treatment cost tended to be lower for TNFi cycling, patients who swapped to a non-TNF-targeted treatment (abatacept, anakinra, rituximab, tocilizumab, or tofacitinib) had longer latencies to discontinuation than those who cycled to a second TNFi. Moreover, in another US claim-based analysis, Bonafede et al. showed that switching to non-TNF-targeted treatments (abatacept, tocilizumab, or tofacitinib) was associated with better outcomes, that is continuation at 12 months, than cycling to TNFi. In addition, a recent pragmatic randomized trial showed that non-TNF biologic agents (abatacept, tocilizumab, and rituximab) achieved a better treatment response than a
### Table 3. Univariable and multivariable Cox proportional hazard models of baseline variables predictive of discontinuation of second-line targeted treatment.

|                                | Univariable analysis | p     | Multivariable analysis | p     |
|--------------------------------|----------------------|-------|------------------------|-------|
| **Age at commencement, years** | 0.998 [0.977–1.319]  | 0.831 |                        |       |
| **Men**                        | 1.439 [0.723–2.865]  | 0.300 |                        |       |
| **Disease duration, months**   | 1.002 [0.999–1.005]  | 0.202 |                        |       |
| **BMI, kg/m²**                 | 0.981 [0.904–1.064]  | 0.638 |                        |       |
| **Current smoker**             | 1.095 [0.395–3.035]  | 0.861 |                        |       |
| **Diabetes mellitus**          | 0.613 [0.244–1.541]  | 0.299 |                        |       |
| **Hypertension**               | 0.957 [0.511–1.789]  | 0.890 |                        |       |
| **Education**                  | 1.021 [0.949–1.098]  | 0.585 |                        |       |
| **Duration of TNFi treatment, months** | 1.018 [0.993–1.043] | 0.154 |                       |       |
| **First TNFi**                 |                      |       |                        |       |
| Etanercept                     | Reference group      |       |                        |       |
| Infliximab                     | 0.821 [0.433–1.554]  | 0.544 |                        |       |
| Adalimumab                     | 0.517 [0.249–1.077]  | 0.078 |                        |       |
| Golimumab                      | 0.528 [0.155–1.794]  | 0.306 |                        |       |
| **Reason for discontinuing first TNFi** |                    |       |                        |       |
| Inefficacy                     | Reference group      |       |                        |       |
| Adverse events                 | 1.240 [0.674–2.279]  | 0.489 |                        |       |
| Other                          | 3.559 [1.251–10.122] | 0.017 |                        |       |
| Use of concomitant csDMARDs    | 0.563 [0.275–1.154]  | 0.117 | 0.382 [0.170–0.858]   | 0.020 |
| Use of concomitant corticosteroids | 0.899 [0.405–1.995] | 0.793 |                        |       |
| Erosion or joint space narrowing on X-ray^a | 1.560 [0.764–3.187] | 0.222 |                        |       |
| RF positivity                  | 0.563 [0.289–1.096]  | 0.091 |                        |       |
| Anti-CCP positivity            | 0.403 [0.187–0.871]  | 0.021 |                        |       |
| Swollen joint counts (44 joints) | 1.034 [1.002–1.068] | 0.036 |                        |       |
| Tender joint counts (44 joints) | 1.024 [0.994–1.055] | 0.124 |                        |       |
| PGA                            | 1.101 [0.988–1.228]  | 0.082 |                        |       |
| PhGA                           | 1.123 [1.005–1.255]  | 0.040 | 1.199 [1.055–1.362]   | 0.005 |
| Elevated ESR^a                 | 0.869 [0.472–1.602]  | 0.653 |                        |       |
| Elevated CRP^a                 | 1.255 [0.681–2.204]  | 0.498 |                        |       |
| DAS28-ESR^a                    | 1.153 [0.975–1.364]  | 0.097 |                        |       |

(continued)
In summary, our study supported the use of non-TNF-targeted treatment in RA patients with initial TNFi failure. To further investigate the efficacy of swap treatment, we stratified swap patients according to the MOA non-TNF-targeted biologic agent used. However, we found no statistically significant differences in drug continuation among abatacept, rituximab, and tocilizumab in the non-TNF biologic group.

Although available data support the use of non-TNF-targeted treatment after first TNFi failure, there is no clear evidence of the superiority of any particular non-TNF bDMARD. In RA patients with an inadequate response to TNFi, a prospective cohort study showed that short-term drug continuation was better in patients treated with rituximab or tocilizumab than in those treated with abatacept. However, similar to our results, Favalli et al. reported no significant differences in retention rates among these three non-TNF biologic agents in the swap group. Direct head-to-head randomized clinical trials comparing abatacept, tocilizumab, and tofacitinib are needed.

Second TNFi in patients with an insufficient response to the first TNFi. In summary, our study supported the use of non-TNF-targeted treatment in RA patients with initial TNFi failure.
with an inadequate response to a prior TNFi. As there have been no randomized trials directly comparing these drugs, a network meta-analysis was performed to compare bDMARDs, including TNFis (golimumab, abatacept, rituximab, and tocilizumab), with tofacitinib in RA patients showing an inadequate response to TNFi or treatment failure. This analysis suggested that tofacitinib was comparable to non-TNFi bDMARDs (abatacept, rituximab, and tocilizumab) in terms of efficacy, based on the ACR response rates at Weeks 12 and 24, and improvement in the Health Assessment Questionnaire Disability Index (HAQ-DI) at Week 12; in this previous study, efficacy was not different between tofacitinib and golimumab as a second-TNFi. In addition, although a claim-based analysis supported switching to non-TNFi-targeted treatments, including tofacitinib, rather than cycling to TNFi, that study did not present a comparison between JAKi and non-TNFi biologic agents. In our study, although the proportion of patients achieving remission or LDA at the 1-year follow-up was not higher in the JAKi than second TNFi group, the JAKi group showed marked improvement in the DAS28-ESR score compared to the second TNFi group. This may have been due to the higher disease activity at the time of enrollment in the JAKi than TNFi group. Taken together, our results provide additional evidence supporting the use of JAKi as an effective therapy for patients with first TNFi treatment failure. However, as our study did not include upadacitinib, which showed greater improvement of disease activity than TNFi [28], further studies are needed to confirm our findings.

In this study, anti-CCP positivity and concomitant use of csDMARDs were associated with continuation of the second TNFi and non-TNFi-targeted treatment after initial TNFi failure, respectively. Previous studies aiming to identify predictors of the response to targeted treatments, such as bDMARDs and tsDMARDs, have not been fully validated and yielded inconsistent results. A large observational study indicated that patients with anti-CCP positivity showed greater clinical improvement after the initiation of TNFi than those negative for anti-CCP antibodies. However, a meta-analysis showed that anti-CCP status was not associated with the response to TNFi. In addition, other studies demonstrated that anti-CCP-negative patients showed better continuation of, or a greater response to, TNFi compared to patients who were anti-CCP-positive. Rather than predicting a response to TNFi, the presence of anti-CCP antibodies was a predictive factor of better responses to non-TNFi biologics, including abatacept and rituximab. With regard to concomitant csDMARDs, studies have shown that the use of csDMARDs, such as MTX, was associated with clinical benefits, including improved continuation of TNFi treatment and disease activity compared to TNFi alone. In patients with high-disease activity, concomitant MTX use was associated with a higher remission rate (adjusted odds ratio 2.54) with tocilizumab treatment. On the one hand, no significant differences were found in the clinical outcomes of abatacept between patients with and without concomitant MTX treatment. However, these studies mostly focused on RA patients treated with first bDMARDs. Due to differences in characteristics between patients treated with first- and second-line targeted treatment, such as the presence of anti-drug antibodies, disease duration, and alterations of immunogenicity, predictors of the response to these drugs may be different. Although further studies are needed to confirm our results, this study provided useful information for clinical decision-making as it pertains to the choice of second-line targeted therapy for patients with RA.

This study had several strengths and limitations. We analyzed KOBIO data collected up to the end of 2020, and the KOBIO data included newer classes of biologic agents and JAKis. Furthermore, the KOBIO registry enrolled RA patients during routine clinical practice, and data were collected prospectively from both academic and community centers. Therefore, our data reflect the actual patterns of targeted treatment in recent years and represent various prescribing patterns and reasons for switching targeted treatments in real-world settings. However, our study also had several limitations. First, any open, non-randomized study has an inherent limitation in terms of the assignment of targeted treatments; selection bias is possible. Although all decisions to initiate or switch targeted treatments were based on the Korean National Health Insurance (KNHI) reimbursement criteria, the choice of targeted treatment could vary depending on the preference of the treating rheumatologist. Furthermore, Kearsley-Fleet et al. suggested that increased class availability and higher expectations for bDMARDs in the more recent cohort could have led to selection bias. Therefore, because of differences in the availability of targeted treatments over time,
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selection bias might also be present in our study. At the time of cohort establishment, all targeted treatments (with the exception of JAKi) were approved as first-line targeted treatments after csDMARDs failure. In May 2017, JAKis were approved as first-line targeted agents that could be initiated after csDMARDs failure, but rituximab remained a second-line targeted agent. Thus, differences in availability among targeted treatments over time may have created selection bias. In an effort to avoid this, the non-TNF-targeted treatment group was divided into two subgroups (non-TNFi biologic and JAKi subgroups). Therefore, we believe that with exception of JAKi, the class availability and expectations for targeted treatments were generally consistent during the study period. Second, RA patients switching to non-TNFi-targeted treatments exhibited a higher anti-CCP-positive rate than the other patients, and those switching to non-TNFi bDMARDs received a first TNFi treatment for an average of 4 months less than the other patients. As this was an observational study, we could not strictly control the RA treatment; also, we may have failed to control for some confounding factors that can be better balanced in randomized controlled trials. In addition, the number of enrolled patients was relatively small, causing issues with certain analyses. Third, there are large disparities in the use of targeted drugs across countries. This discrepancy may be due to differences in healthcare systems and the accessibility of RA treatment among countries. Therefore, more recent data from around the world are needed to improve the generalizability of our results.

In conclusion, the results of this study showed that, after initial TNFi failure, switching to different MOA non-TNF-targeted treatments was associated with significantly better treatment outcomes and drug continuation than cycling to another TNFi. Although various targeted treatments, including TNFIs, have revolutionized the treatment of RA, switching targeted drugs may be unavoidable. Therefore, understanding whether swapping to non-TNF-targeted treatment and cycling to another TNFi have different effects is important to establish evidence-based guidelines for switching to targeted treatments after first TNFi failure. This study provided useful data to develop better protocols for switching to targeted treatment of RA.

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Author contribution(s)
Dong-Jin Park: Conceptualization; Data curation; Formal analysis; Investigation; Methodology; Project administration; Writing – original draft; Writing – review & editing.
Sung-Eun Choi: Data curation; Formal analysis; Investigation; Methodology; Resources; Visualization.
Ji-Hyoun Kang: Data curation; Formal analysis; Investigation; Methodology; Resources; Visualization
Kichul Shin: Conceptualization; Data curation; Formal analysis; Investigation; Methodology; Resources; Visualization
Yoon-Kyoung Sung: Conceptualization; Data curation; Investigation; Project administration; Resources; Software; Supervision; Visualization
Shin-Seok Lee: Conceptualization; Data curation; Formal analysis; Funding acquisition; Investigation; Methodology; Project administration; Supervision; Visualization; Writing – review & editing.

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**ORCID iDs**

Yoon-Kyoung Sung  https://orcid.org/0000-0001-6691-8939

Shin-Seok Lee  https://orcid.org/0000-0001-6810-7355

**Supplemental material**

Supplemental material for this article is available online.

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