Discontinuation of Azathioprine could be considered in pediatric patients with Crohn’s disease who have sustained clinical and deep remission

Tae Jong Jeong1,6, Eun Sil Kim1,6, Yiyoung Kwon1, Seonwoo Kim2, Sang Won Seo3,4,5, Yon Ho Choe1* & Mi Jin Kim1*

Few studies have demonstrated treatment strategies about the duration and cessation of medications in patients with Crohn’s disease (CD). We investigated factors affecting clinical relapse after infliximab (IFX) or azathioprine (AZA) withdrawal in pediatric patients with CD on combination therapy. Pediatric patients with moderate-to-severe CD receiving combination therapy were analyzed retrospectively and factors associated with clinical relapse were investigated. Discontinuation of IFX or AZA was performed in patients who sustained clinical remission (CR) for at least two years and achieved deep remission. A total of 75 patients were included. Forty-four patients (58.7%) continued with combination therapy and 31 patients (41.3%) discontinued AZA or IFX (AZA withdrawal 10, IFX withdrawal 15, both withdrawal 6). Cox proportional-hazards regression and statistical internal validation identified three factors associated with clinical relapse: IFX cessation (hazard ratio; HR 2.982, \( P = 0.0081 \)), IFX TLs during maintenance therapy (HR 0.581, \( P = 0.003 \)), 6-thioguanine nucleotide (6-TGN) level (HR 0.978, \( P < 0.001 \)). However, AZA cessation was not associated with clinical relapse (\( P = 0.9021 \)). Even when applied in pediatric patients who met stringent criteria, IFX cessation increased the relapse risk. However, withdrawal of AZA could be contemplated in pediatric patients with CD who have sustained CR for at least 2 years and achieved deep remission.

Crohn’s disease (CD) is a chronic, relapsing disease of the gastrointestinal tract that can cause serious morbidity and disability1. Current treatment trends recommend the early introduction of biologics in pediatric patients with CD exhibiting severe disease2. Infliximab (IFX), one type of anti-tumor necrosis factor (TNF) agent, can be appropriate initial therapy in moderate-to-severe pediatric CD patients3. It is well known that corticosteroid-free clinical remission (CR) is more likely to be achieved in patients with moderate-to-severe CD when treated with a combination therapy that includes IFX plus azathioprine (AZA) or with IFX monotherapy, than in those receiving AZA monotherapy4. Early introduction of IFX, known as top-down therapy, is an important option for pediatric patients with moderate-to-severe CD and has become the mainstay of treatment for CD5,6. However, since long-term use of AZA increases the risk of complications such as leukopenia, opportunistic infections, and lymphoma in children, concomitant use of immunomodulators and biologics has raised safety concerns7,8. In addition, combination therapy is not clearly associated with high efficacy after achieving long-term remission9. Despite the increasing use of combination therapy, few studies have established the optimum strategy in terms of duration and cessation of medications after achieving remission, which is still a matter of discussion10. Furthermore, although it is well known that concomitant treatment with immunomodulator reduces the formation of antibody-to-infliximab (ATI) and improves the pharmacokinetics of IFX, other factors affecting ATI formation are not clearly identified11.

1Department of Pediatrics, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea. 2Statistics and Data Center, Samsung Medical Center, Seoul, Korea. 3Department of Neurology, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea. 4Neuroscience Center, Samsung Medical Center, Seoul, Korea. 5Department of Clinical Research Design and Evaluation, SAIIHST, Sungkyunkwan University, Seoul, Korea. 6These authors contributed equally: Tae Jong Jeong and Eun Sil Kim. *email: i101016@skku.edu; mijin1217.kim@samsung.com
The aim of this study was to determine factors affecting clinical relapse after achieving remission in pediatric CD and to estimate the proper duration and time of cessation of early combination therapy in a selected population of pediatric patients with CD in a real-life cohort.

Materials and methods

Study design and patients. This was a retrospective observational study conducted at the Department of Pediatrics of Samsung Medical Center in the Republic of Korea, between January 2012 and March 2018. Eligible patients had moderate-to-severe luminal CD and were aged less than 19 years at both time of diagnosis and the time of their first treatment with IFX. CD was diagnosed in accordance with the revised Porto criteria of the European Society for Pediatric Gastroenterology, Hepatology and Nutrition\textsuperscript{12} and disease phenotype classification was based on the Paris classification\textsuperscript{13}. Patients on early combination therapy were included in the analysis, and patients underwent endoscopy on a regular basis every 1–2 years.

Patients were excluded if they were primary non-responders to IFX, or developed ATI during induction therapy of IFX. In addition, patients who were poor metabolizers of AZA according to thiopurine methyltransferase (TPMT) and nucleotide triphosphate diphosphatase 15 (NUDT15) genotype, were excluded for accurate analysis of effects of combination therapy\textsuperscript{14}.

IFX was administered according to the scheduled induction regimen of 5 mg/kg at weeks 0, 2, and 6, and scheduled maintenance IFX was repeated every 8 weeks. Twenty-six patients were treated with originator Remicade\textsuperscript{®}, 35 patients treated with Remsima\textsuperscript{®}, a biosimilar version of IFX, and 14 patients switched from Remicade to Remsima during the follow up period. Since there was no difference in the clinical efficacy of Remicade and Remsima according to the previous studies\textsuperscript{16–18}, it was considered that there was no difference according to the type of administered IFX. IFX doses were adjusted by IFX TL during maintenance therapy; if IFX TL was less than 3 μg/ml, dose intensification was performed. Azathioprine was given at doses of 0.5–1 mg/kg/day and was later modified when required according to TPMT and NUDT15 genotype and thiopurine metabolite levels of 6-thioguanine nucleotide (6-TGN) levels (therapeutic range: 235–450)\textsuperscript{19}.

Drug cessation was performed among patients who maintained CR more than two years and at the same time achieved deep remission. The decision to discontinue either IFX or AZA was determined by various requirements, such as concerns about side effects from long-term use of the medication, cost effectiveness, and patients’ and parental demands. We did not selectively discontinue specific medications for specific patients.

Data collection and definitions. Demographic and clinical data at diagnosis, at combination therapy initiation, at drug cessation, and during follow-up were collected from electronic medical records: age, sex, disease phenotype, body mass index (BMI), pediatric Crohn’s disease activity index (PCDAI), laboratory results, simple endoscopic score for CD (SES-CD), histologic results and concomitant medication. Date related to AZA or IFX cessation, date of clinical relapse, 6-TGN, IFX TLs, and formation of ATI were collected retrospectively from electronic charts or electronic test results. The data described above was used to determine factors affecting clinical relapse and to estimate the appropriate duration and time of cessation of early combination therapy.

Serum samples were obtained prior to each infusion for measurement of IFX TLs and ATI. IFX TLs and ATI had been quantified using an enzyme-linked immunosorbent assay (ELISA; Matriks Biotek Laboratories, Ankara, Turkey). We used sandwich ELISA to measuring IFX TLs and free ATI. We measured free ATI which is known to be associated with low serum IFX TLs and probability of active disease using drug-sensitive ATI assays\textsuperscript{17}. AZA concentration was estimated indirectly every 3–6 months by measuring 6-TGN levels which are considered to be the primary active metabolite of AZA.

CR was defined as a PCDAI < 10 points. Deep remission was defined as the absence of mucosal ulceration (SES-CD < 2), i.e., mucosal healing, at the same time as the achievement of histologic remission which means the absence of microscopically active inflammation in all gastrointestinal tissue obtained by endoscopy. Clinical relapse was defined as a PCDAI score of ≥ 10 with a change of at least 10 points from the previous visit with the need for treatment intensification, defined as either i) the addition of a new medication, ii) dose escalation of maintenance treatment and (iii) intestinal surgery because of strictureing or penetrating CD.

Statistical analysis. Baseline characteristics of subjects were explored with descriptive statistics through frequencies (proportion) for categorical variables, or medians (interquartile range; IQR) for continuous variables. Data related to clinical characteristics, the duration and cessation timing of medication, 6-TGN, and IFX TLs were analyzed. Data with normal distribution and/or equal variances (Levene’s and F-test for equality of variances) were analyzed with 1-way ANOVA or t-test for independent samples, whereas data without normal distribution were analyzed with Kruskal–Wallis H-test or Mann–Whitney U-test. The risk factors associated with clinical relapse were identified by Cox proportional-hazards regression models and internally validated by bootstrap resampling\textsuperscript{19}. Time-varying covariance occurs when covariates change over time during the follow-up period. Drug-related factors were more appropriate to be analyzed as time-varying covariates rather than time-independent covariates because variables such as drug administration (or discontinuation) and drug concentration vary over time. Therefore, cessation and duration of medication, 6-TGN, and IFX TLs were analyzed using time-varying covariates with the Cox regression model to estimate these effects on survival time. For this, we organized the data in a counting process style. Hazard ratio (HR) for each variable was derived within 95% confidence intervals (CIs). Relapse-free curve was estimated using an extended Kaplan–Meier method that can be used with time-varying covariates\textsuperscript{21}. Data were considered significantly different if two-sided P was < 0.05. All statistical analyses were performed using SAS software (version 9.4; SAS Institute Inc., Cary, NC, USA) and R software (version 3.6.1; R Foundation, Vienna, Austria).
Ethics declarations. This study was approved by the Institutional Review Board of the Samsung Medical Center (IRB file No. 2020–06–128–001), and was conducted in accordance with the Declaration of Helsinki. All patients and parents and/or legal guardian of subjects who are under 18 provided written informed consent. We confirmed that all methods were performed in accordance with the approved guidelines and regulations. We reported and presented data according to the STROBE statement.

Results
Baseline characteristics. From January 2012 to March 2018, a total of 216 pediatric patients were diagnosed with CD and of these 75 patients were finally considered eligible for analysis as shown in the flow diagram for patient selection (Fig. 1). Among the study participants, 48 patients (64.0%) were male and the median age of subjects at diagnosis was 14.2 years (IQR 12.0–17.0). The median initial PCDAI at diagnosis was 39.7 (IQR 37.5–45.0) and median initial SES-CD was 16.9 (IQR 11.0–24.0). The median observational duration was 41.5 months (IQR 23.0–58.7 months). Other baseline characteristics are described in detail in Table 1.

Relapse rate of patients according to withdrawal of medications. Of 75 patients, 31 (41.3%) patients met the criteria of sustained CR more than two years and the definition of deep remission, and discontinued AZA or IFX according to various requirements. Sixteen patients withdrew AZA, 21 patients IFX, and among them, six patients discontinued both. The remaining 44 patients (58.7%) who achieved CR but did not reach deep remission, continued combination therapy with IFX and AZA. The mean durations of AZA and IFX therapy were 38.0 ± 19.3 months and 32.0 ± 18.9 months respectively. In the drug discontinuation group, the mean follow-up duration after AZA and IFX withdrawal was 14.0 ± 9.5 months and 28.0 ± 22.9 months respectively. When comparing the group that withdrew AZA with the group who discontinued IFX, there was no significant difference in disease activity and laboratory results at the time of diagnosis and at the time of drug discontinuation (Table 2).

Among 75 patients, 31 (41.3%) were clinically relapsed and 44 patients (58.7%) maintained CR during the study period. Four of 16 patients who withdrew AZA experienced clinical relapse (4/16, 25.0%), 15 of 21 patients who withdrew IFX experienced relapse (15/21, 71.4%), and four of six patients who withdrew all drugs experienced relapse (4/6, 66.6%). Sixteen of 44 patients (16/44, 36.4%) who continued IFX and AZA during study period experienced relapse.

Factors associated with clinical relapse. A Cox proportional-hazards regression identified four factors associated with clinical relapse. IFX cessation (HR 2.982, 95% CI 1.322–6.485, P = 0.0081), formation of ATI (HR 3.120, 95% CI 1.069–9.103, P = 0.0373), low IFX TLs during maintenance therapy (HR 0.581, 95%
CI 0.432–0.781, \( P = 0.0003 \) and low 6-TGN levels (HR 0.978, 95% CI 0.968–0.987, \( P < 0.0001 \)) were found to increase the risk of clinical relapse. AZA cessation (HR 1.078, 95% CI 0.327–3.550, \( P = 0.9021 \)) was not shown to increase the risk of clinical relapse (Table 3). In the analysis using bootstrapped data for internal validation, IFX cessation (HR 3.178, 95% CI 1.294–6.312), low 6-TGN level (HR 0.941, 95% CI 0.267–0.988), low IFX TLs (HR 0.544, 95% CI 0.031–0.765) were confirmed to be associated with clinical relapse, whereas ATI formation was not shown to increase risk of clinical relapse in bootstrapped data (HR 5.869, 95% CI 0.775–22.741).

To evaluate the relapse-free curve in relation to discontinuation of medication and ATI formation, Kaplan–Meier survival curves were calculated (Fig. 2). Patients who discontinued IFX (Fig. 2B, \( P = 0.0081 \)) or developed ATI (Fig. 2C, \( P = 0.0373 \)) had significantly poorer outcomes compared to those who did not. However, there was no difference in survival rate between patients who continued or discontinued AZA (Fig. 2A, \( P = 0.9021 \)).

**Factors associated with formation of antibody-to-infliximab.** The formation of ATI was detected in nine patients (12.0%) receiving combination therapy. Factors associated with formation of ATI during maintenance therapy were evaluated. According to Cox proportional-hazards regression, only 6-TGN level was related to the formation of ATI (HR 0.988, 95% CI 0.980–0.996, \( P = 0.0048 \)). AZA cessation (HR 17.99, 95% CI 0.194–1663.717, \( P = 0.2109 \)) was not related to formation of ATI. Within the therapeutic range (235–450 pmol/\( 8 \times 10^8 \) RBC)\(^{18} \), high 6-TGN level is a factor that lowers the risk of developing ATI (HR 0.988, 95% CI 0.980–0.996, \( P = 0.0048 \), Table 4). In addition, analysis using bootstrap for internal validation also found that low 6-TGN is associated with risk of ATI formation (HR 0.978, 95% CI 0.929–0.998). When comparing patients according to

| CI 0.432–0.781, \( P = 0.0003 \) and low 6-TGN levels (HR 0.978, 95% CI 0.968–0.987, \( P < 0.0001 \)) were found to increase the risk of clinical relapse. AZA cessation (HR 1.078, 95% CI 0.327–3.550, \( P = 0.9021 \)) was not shown to increase the risk of clinical relapse (Table 3). In the analysis using bootstrapped data for internal validation, IFX cessation (HR 3.178, 95% CI 1.294–6.312), low 6-TGN level (HR 0.941, 95% CI 0.267–0.988), low IFX TLs (HR 0.544, 95% CI 0.031–0.765) were confirmed to be associated with clinical relapse, whereas ATI formation was not shown to increase risk of clinical relapse in bootstrapped data (HR 5.869, 95% CI 0.775–22.741).

To evaluate the relapse-free curve in relation to discontinuation of medication and ATI formation, Kaplan–Meier survival curves were calculated (Fig. 2). Patients who discontinued IFX (Fig. 2B, \( P = 0.0081 \)) or developed ATI (Fig. 2C, \( P = 0.0373 \)) had significantly poorer outcomes compared to those who did not. However, there was no difference in survival rate between patients who continued or discontinued AZA (Fig. 2A, \( P = 0.9021 \)).

**Factors associated with formation of antibody-to-infliximab.** The formation of ATI was detected in nine patients (12.0%) receiving combination therapy. Factors associated with formation of ATI during maintenance therapy were evaluated. According to Cox proportional-hazards regression, only 6-TGN level was related to the formation of ATI (HR 0.988, 95% CI 0.980–0.996, \( P = 0.0048 \)). AZA cessation (HR 17.99, 95% CI 0.194–1663.717, \( P = 0.2109 \)) was not related to formation of ATI. Within the therapeutic range (235–450 pmol/\( 8 \times 10^8 \) RBC)\(^{18} \), high 6-TGN level is a factor that lowers the risk of developing ATI (HR 0.988, 95% CI 0.980–0.996, \( P = 0.0048 \), Table 4). In addition, analysis using bootstrap for internal validation also found that low 6-TGN is associated with risk of ATI formation (HR 0.978, 95% CI 0.929–0.998). When comparing patients according to
Our data that IFX cessation in patients with CD was associated with a high risk of clinical relapse is consistent with the results of other previously published studies. A recent retrospective study conducted in Korea on adults evaluated the long-term outcomes following cessation of anti-TNF-α treatment in IBD patients with CR. After cessation of anti-TNF-α treatment for CD patients, the cumulative relapse rates at 1, 3, and 5 years were 11.3%, 46.7%, and 62.5%. In this cohort, mucosal healing rate before discontinuation of anti-TNF-α treatment was 22.5%. Another recent retrospective study performed by Luca and colleagues investigated the clinical course after anti-TNF-α treatment discontinuation in selected pediatric patients with IBD who achieved deep remission similar to our study. Among these patients, relapse-free survival rates at 12, 24, and 36 months were 83.3%, 71.1%, and 23.7%, respectively. In contrast, none of the patients with CD who maintained anti-TNF-α treatment after achieving mucosal healing experienced clinical relapse.

### Table 2. Comparison between patients discontinuing infliximab or azathioprine. Baseline characteristics of subjects who withdrew infliximab or azathioprine were explored with descriptive statistics through frequencies (proportion) for categorical variables or medians (interquartile range[IQR]) for continuous variables. Chi-square test; Kruskal–Wallis test. ESR, Erythrocyte sedimentation rate; PCDAI, Pediatric Crohn’s disease activity index; SES-CD, Simple endoscopic score for Crohn’s disease; AZA, azathioprine; IFX, infliximab.

|                     | IFX withdrawal group (n=15) | AZA withdrawal group (n=10) | IFX and AZA withdrawal group (n=6) | P-value |
|---------------------|----------------------------|-----------------------------|-----------------------------------|---------|
| Sex, male (%)       | 7 (31.80)                  | 9 (40.90)                   | 6 (27.30)                         | 0.018^2 |
| At diagnosis        |                            |                             |                                   |         |
| White blood cell    | 7.80 (7.43, 9.71)           | 9.29 (8.23, 10.72)          | 8.61 (7.80, 9.19)                 | 0.673^6 |
| Hematocrit, %       | 35.60 (33.10, 39.10)        | 35.20 (32.30, 39.10)        | 40.00 (37.90, 41.70)              | 0.069 K |
| ESR, mm/h           | 53.00 (41.50, 78.00)        | 63.00 (39.30, 83.50)        | 25.00 (23.00, 29.30)              | 0.071 K |
| C-reactive protein, | 1.21 (0.64, 7.60)           | 2.37 (1.84, 3.52)           | 2.33 (1.10, 3.46)                 | 0.712 K |
| Albumin, g/dl       | 3.80 (3.60, 4.30)           | 3.80 (3.50, 4.30)           | 4.20 (4.00, 4.40)                 | 0.479 K |
| PCDAI at diagnosis  | 40.00 (33.30, 46.30)        | 35.00 (30.60, 41.90)        | 35.00 (30.00, 41.00)              | 0.515 K |
| SES-CD at diagnosis | 16.00 (11.50, 19.50)        | 18.00 (14.30, 24.00)        | 13.50 (7.00, 17.80)               | 0.298 K |
| At IFX or AZA withdrawal |                     |                             |                                   |         |
| White blood cell    | 5.52 (4.80, 7.07)           | 6.11 (5.45, 6.88)           | 5.67 (5.11, 5.82)                 | 0.830 K |
| Hematocrit, %       | 39.40 (37.10, 43.80)        | 45.40 (44.50, 46.60)        | 43.20 (42.50, 45.50)              | 0.174 K |
| ESR, mm/h           | 12.00 (4.50, 20.50)         | 4.50 (2.30, 9.00)           | 5.00 (2.80, 11.80)                | 0.243 K |
| C-reactive protein, | 0.03 (0.03, 0.08)           | 0.04 (0.03, 0.04)           | 0.13 (0.03, 0.36)                 | 0.437 K |
| Albumin, g/dl       | 4.50 (4.40, 4.70)           | 4.70 (4.60, 4.90)           | 4.50 (4.50, 4.60)                 | 0.196 K |
| PCDAI at drug       | 0 (0, 0)                    | 0 (0, 0)                    | 0 (0, 0)                          | 0.669 K |
| SES-CD at drug withdrawal | 0 (0, 0.45)                  | 0 (0, 0)                    | 0 (0, 0.25)                      | 0.496 K |

### Discussion

We investigated the factors affecting clinical relapse and estimated the optimum duration and time of cessation of medication in moderate to severe pediatric CD treated with combination therapy. We identified three factors related to elevated risk of clinical relapse: IFX cessation, low IFX TLs, low 6-TGN level. Interestingly, withdrawal of AZA was not directly associated with clinical relapse in pediatric CD patients with sustained CR of at least two years and who had achieved deep remission.

In clinical practice, physicians may encounter questions about the feasibility of therapeutic de-escalation in CD patients who have achieved deep remission, on account of safety, adverse events and cost. There is a paucity of data related to the optimum duration and the appropriate time to cease medication after achieving remission in patients with CD, particularly in pediatric patients.

It is now known that early introduction of biologics has tremendous advantages in terms of maintaining tight control of inflammation and catching the therapeutic window. A randomized controlled trial (RCT) revealed that co-treatment with IFX and AZA significantly increases the rate of CR and mucosal healing compared to AZA monotherapy in CD patients. In addition, recent studies have reported that endoscopic and histological mucosal healing should be treat-to-target in CD patients instead of symptomatic control to achieve better outcomes and minimize future complications. With the development of treatment strategies, clinicians are being confronted with a crossroads decision about when to stop or reduce the therapy dose once patients achieve remission.

A systematic review of cessation studies for patients with inflammatory bowel disease (IBD) concluded that about 50 percent of patients who withdrew anti-TNF-α agents after combination therapy with immunomodulators maintained remission for 24 months. Reenaers et al. reported that among patients with CD who withdrew IFX in stable CR state, twenty-one percent did not restart biologics including IFX, and sustained CR for seven years after IFX cessation. Therefore, it seems reasonable to conclude that there may be a subgroup of patients who are good candidates for treatment withdrawal.

Our data that IFX cessation in patients with CD was associated with a high risk of clinical relapse is consistent with the results of other previously published studies. A recent retrospective study conducted in Korea on adults evaluated the long-term outcomes following cessation of anti-TNF-α treatment in IBD patients with CR. After cessation of anti-TNF-α treatment for CD patients, the cumulative relapse rates at 1, 3, and 5 years were 11.3%, 46.7%, and 62.5%. In this cohort, mucosal healing rate before discontinuation of anti-TNF-α treatment in CD patents was 22.5%. Another recent retrospective study performed by Luca and colleagues investigated the clinical course after anti-TNF-α treatment discontinuation in selected pediatric patients with IBD who achieved deep remission similar to our study. Among these patients, relapse-free survival rates at 12, 24, and 36 months for CD were 83.3%, 71.1%, and 23.7%, respectively. In contrast, none of the patients with CD who maintained anti-TNF-α treatment after achieving mucosal healing experienced clinical relapse.
Table 3. Factors affecting clinical relapse in pediatric patients with Crohn’s disease. The risk factors associated with clinical relapse were identified by Cox proportional-hazards regression models. In this model, cessation and duration of medication, 6-TGN, and IFX TL were analyzed as time-varying covariates. Hazard ratio (HR) for each variable was derived within the 95% confidence intervals (CIs). *Variables were analyzed as time varying covariates. PCDAI, pediatric Crohn’s disease activity index; SES-CD, simple endoscopic score for Crohn’s disease; AZA, azathioprine; IFX, infliximab; 6-TGN, 6-thioguanine nucleotide.

| Variables                                      | Univariate Cox analysis | Bootstrapped data (n=500) |
|------------------------------------------------|-------------------------|----------------------------|
| Age                                            | 0.965                   | 0.967                      |
| Sex                                            |                         |                            |
| Male                                           | Reference               | Reference                   |
| Female                                         | 1.203                   | 1.287                      |
| Initial laboratory findings                    |                         |                            |
| White blood cell counts                        | 0.676                   | 0.897                      |
| Hematocrit                                     | 0.960                   | 0.962                      |
| Albumin                                        | 0.751                   | 0.779                      |
| Erythrocyte sedimentation rate                 | 1.009                   | 1.009                      |
| C-reactive protein                             | 0.964                   | 0.969                      |
| Initial PCDAI                                   | 1.004                   | 1.005                      |
| Initial SES-CD                                 | 0.995                   | 0.997                      |
| Initial disease phenotype                      |                         |                            |
| L1 (Ileal) + L3 (Fleacoliconic)                 | Reference               | Reference                   |
| L2 (Colonic)                                   | 1.996                   | 5.190                      |
| Initial disease behavior                       |                         |                            |
| B2 (Strictureing)                              | Reference               | Reference                   |
| B1 (Inflammatory)                              | 0.891                   | 0.954                      |
| Initial growth impairment                      | 0.669                   | 0.722                      |
| *AZA cessation                                 | 1.078                   | 1.264                      |
| *IFX cessation                                 | 2.928                   | 3.178                      |
| *Mesalazine cessation                          | 1.165                   | 1.334                      |
| *antibody-to-IFX formation                     | 3.120                   | 5.869                      |
| *IFX Trough levels                             | 0.581                   | 0.544                      |
| *6-TGN levels                                  | 0.978                   | 0.941                      |

Figure 2. Relapse-free curve according to drug cessation and development of antibody-to-infliximab. AZA, azathioprine; IFX, infliximab; ATI, antibody-to-infliximab.
Table 4. Factors affecting development of antibody-to-infliximab. The risk factors associated with formation of ATI were identified by Cox proportional-hazards regression models. In this model, cessation of medication and 6-TGN were analyzed as time-varying covariates. Hazard ratio (HR) for each variable was derived within the 95% confidence intervals (CIs). *Variables were analyzed as time varying covariates. PCDAI, pediatric Crohn’s disease activity index; SES-CD, simple endoscopic score for Crohn’s disease; AZA, azathioprine; 6-TGN, 6-thioguanine nucleotide.

| Variables                        | Univariate Cox analysis | Bootstrapped data (n = 500) |
|----------------------------------|-------------------------|----------------------------|
|                                  | HR 95% CI | P-value | HR 95% CI |
| Age                              | 1.132 | 0.872–1.468 | 0.351 | 1.169 | 0.852–1.696 |
| **Sex**                          |           |       |           |           |       |
| Male                             | Reference |       |       | Reference |       |
| Female                           | 1.561 | 0.419–5.813 | 0.507 | 2.116 | 0.381–6.628 |
| **Initial laboratory findings**  |           |       |           |           |       |
| White blood cell counts          | 0.456 | 0.041–5.108 | 0.523 | 1.001 | 0.044–4.311 |
| Hematocrit                       | 1.033 | 0.900–1.185 | 0.646 | 1.048 | 0.911–1.237 |
| Albumin                          | 0.859 | 0.303–2.436 | 0.775 | 0.924 | 0.401–1.960 |
| Erythrocyte sedimentation rate   | 1.004 | 0.983–1.025 | 0.719 | 1.004 | 0.986–1.024 |
| C-reactive protein               | 0.992 | 0.812–1.211 | 0.934 | 0.981 | 0.720–1.178 |
| Initial PCDAI                    | 1.001 | 0.943–1.063 | 0.965 | 1.001 | 0.959–1.040 |
| Initial SES-CD                   | 0.950 | 0.875–1.030 | 0.215 | 0.940 | 0.816–1.041 |
| **Initial disease phenotype**    |           |       |           |           |       |
| L1 (ileal) + L3 (ileocolonic)    | Reference |       |       | Reference |       |
| L2 (Colonic)                     | 3.297 | 0.410–26.513 | 0.262 | 6.503 | 0.874–24.412 |
| **Initial disease behavior**     |           |       |           |           |       |
| B2 (Strictureing) + B3 (Penetrating) | Reference |       |       | Reference |       |
| B1 (Inflammatory)                | 0.661 | 0.137–3.183 | 0.605 | 0.924 | 0.126–2.800 |
| Initial growth impairment        | 1.115 | 0.279–4.462 | 0.877 | 1.508 | 0.159–4.441 |
| *AZA cessation                    | 17.987 | 0.194–1663.717 | 0.924 | 2.890–19.997 |
| *6-TGN level                     | 0.988 | 0.980–0.996 | 0.978 | 0.929–0.998 |

Figure 3. 6-Thioguanine nucleotide levels according to formation of antibody-to-infliximab. 6-TGN, 6-thioguanine nucleotide; ATI, antibody-to-infliximab.
However, the association between discontinuation of AZA and clinical relapse is controversial. In one reported study, which was focused on CD in CR under IFX-AZA combination therapy, AZA cessation was associated with a high risk of relapse in those patients treated with combination therapy for less than 27 months\(^2\). In contrast, two other RCTs and one meta-analysis suggested that discontinuing AZA from a combination therapy regimen may not differ in clinical relapse rates compared to continuing with combination therapy\(^9,26,33\). However, both RCTs included subjects comprised primarily of those who had previously failed immunomodulators mono-therapy. It could be argued that in these cohorts, the subsequent cessation of this agent would not be expected to have a significant effect compared to a patient cohort in which patients were initially treated with combination therapy\(^10\). Nevertheless, most studies of patients with CD who discontinued AZA after combination therapy revealed that there were no differences between AZA withdrawal from combination therapy and continuation of combination therapy in terms of clinical relapse. In addition, the European Crohn’s and colitis organization guideline provide the opinion that AZA withdrawal in patients treated with combination therapy is inappropriate in patients with high risk/refractory disease or in patients at risk of biologic failure\(^4\). In other words, it can be said that in patients with sustained deep remission, controlled disease activity and low risk of biologic failure, azathioprine could be withdrawn.

When subjects were treated with combination therapy and maintained CR for at least two years, mucosal healing was observed in 82.7% and deep remission was achieved in 41.3% in our study. Based on these findings, we attempted drug cessation assuming that most of the patients who satisfied both conditions had received sufficient treatment and had a low risk of relapse. In our study, AZA cessation was not shown to increase the risk of clinical relapse (HR 1.078, 95% CI 0.327–3.55, \(P=0.9021\)). This result could be partially explained by another study which revealed that withdrawal of AZA after at least six months of combination therapy does not reduce IFX TLs in patients with CD\(^35\). The mean durations of AZA therapy were 38 ± 19.3 months. We also wanted to know the change in the risk of relapse according to the treatment continuation of AZA, but the number of patients who discontinued AZA was small, so further statistical analysis was impossible.

According to our results and those of other reports, IFX TLs and 6-TGN levels affect clinical relapse\(^36,37\). High drug concentrations within the therapeutic range could be a factor in lowering the risk of relapse. Especially, it has been reported that IFX TLs are positively associated with mucosal healing and could be a factor in lowering the risk of relapse during maintenance treatment with IFX\(^38\). Pursuing a high therapeutic concentration may be of concern due to adverse drug reactions. However, using therapeutic drug monitoring (TDM) for personalizing therapy for CD patients, drug concentration could be maintained within the therapeutic range. The average IFX TLs and 6-TGN levels during the follow-up period were 6.1 ± 5.4 μg/mL and 208.5 ± 114.2 pmol/8 × 10^8 RBC respectively. Since patients who were poor metabolizers of AZA and primary non-responders to IFX were excluded from this study, the relationship between higher drug concentrations above therapeutic range of 6-TGN and occurrence of adverse events was not addressed in this study. Interestingly, in our study, IFX TLs at the time of IFX cessation were significantly higher in patients who experienced relapse than patients who maintained CR (5.4 ± 2.8 vs. 2.8 ± 0.8, \(P<0.01\)). The present findings correlate with the findings of previous studies, which indicate higher IFX TLs at IFX cessation were associated with relapse\(^39,40\).

In adults, the development of ATI occurs in up to 65.3% in patients with IBD\(^41\), while ATI have been reported in 8% to 43% of pediatric patients with IBD\(^42\). The results of our study are consistent with these data, as the formation of ATI was detected in 12.0% of patients receiving combination therapy. The development of ATI can neutralize IFX by direct binding of neutralizing antibodies, or accelerating the clearance of the IFX by the binding of non-neutralizing antibodies\(^43,44\). These mechanisms lower IFX TLs and can lead to loss of response during IFX administration\(^45\). Similarly, patients with ATI formation had a lower relapse free survival rate than those with negative ATI in our study (Fig. 3C). ATI formation was shown to increase risk of clinical relapse in univariate Cox analysis in our data, whereas it was not shown to increase risk of clinical relapse in bootstrapped data. Since the number of ATI positive patient is small (\(n=9\)), the estimation and results are considered unreliable. Therefore, it is thought that confirmation through a study using a larger cohort is necessary in the future.

It is well known that concomitant use of immunomodulators reduces ATI formation, and improves the pharmacokinetics of IFX\(^46,47\). This finding is interesting because it appears to be related with the role of AZA in preventing immunogenicity. Recent cross-sectional study of 72 patients receiving maintenance therapy with IFX and a thiopurine for IBD concluded that patients with 6-TGN levels less than 125 pmol/8 × 10^8 RBC were significantly more likely to have ATI (odds ratio 1.3, 95% CI 2.3–72.5, \(P<0.01\))\(^9\). 6-TGN levels of 125 pmol/8 × 10^8 RBC or higher were best predictive of the absence of ATIs; higher 6-TGN concentrations did not provide additional benefit at least in terms of ATI formation. However, other factors affecting ATI formation were not clearly identified. We specifically analyzed whether there is a relationship between the factors affecting clinical relapse and ATI formation. According to our analysis, only 6-TGN level was consistent with an increased risk of formation of ATI and this result is consistent with another study\(^48\).

In our study, AZA cessation (HR 17.99, \(P=0.2109\)) was not associated with formation of ATI. However, with a hazard ratio of 17.99, this affirmation does not seem reliable, in spite of \(P>0.05\) because of small population. In addition, from the results estimated from the bootstrapped data, AZA cessation was found to increase the risk of ATI development. Considering the role of AZA in preventing immunogenicity, AZA cessation might be a risk for ATI formation theoretically. It is thought that this is because the number of ATI-positive patients in outcome and the number of patients who discontinued AZA were small, and the estimation and results could be unreliable.

Although the present study is retrospective and relatively small-sized, it sought a more analytical approach to drug cessation and drug concentration. Drug-related factors were analyzed as time-varying covariates rather than time-independent covariates. Conventional survival analysis is generally applied to the time-independent data, where the exposure variables of interest are often treated as time-fixed\(^8\). However, values of these exposure variables can vary over time and time-fixed analysis may cause bias over time potentially altering the conclusions of the study. To the best of our knowledge, none of the studies to determine when to withdraw drugs in
patients with CD have used time-varying covariates. The factors which are known to affect relapse such as 6-TGN levels, IFX TLs, and timing of drug cessation are all variables that change with time after patient observation starts, therefore, the time-varying covariate approach is preferable for survival analysis in our study. This method increased the reliability of the study because it avoids biases associated with different timing of drug cessation in different patients.

The current study has a few limitations. First, this was a single-center, retrospective study and, consequently, had relatively unstructured follow-up of patients with certain limitations compared to studies using prospective design. However, all subjects received regular examinations such as endoscopy and biopsy on the same principles and IFX TLs or 6-TGN levels were examined at regular intervals. Therefore, the extraction of clinical disease activity using PCDAI, laboratory results, endoscopic and histologic results from medical records was possible at all outpatient visits. Although the decision to withdraw either IFX or AZA was not randomly assigned, there were no differences in disease activity and baseline as described in Table 2. Moreover, our findings are meaningful because these are real-world data from a single center cohort. Second, selection bias might have been introduced because the number of participating patients was relatively small and the observation period was not as long as might be desired. Also, because of adverse events such as lymphoma that may occur in male patients with long-term use of AZA, the group who withdrew AZA had a statistically larger number of male patients than the group who discontinued IFX, resulting in a selection bias of gender. However, in order to compensate for the shortcomings of the small number of subjects, it was attempted to increase the verification power through bootstrapping. Third, since multiple time-varying covariates made multivariate analysis too complicated to perform, only univariate results were presented.

Despite these limitations, our study reports the outcome of drug cessation in strictly selected cohort who sustained sufficient CR and achieved deep remission in real clinical settings. Maintaining the 6-TGN levels and IFX TLs high within the therapeutic range lowers the risk of clinical relapse. In conclusion, even when applied in pediatric patients who meet strict criteria after a sufficient CR period and deep remission, IFX cessation in pediatric CD should be considered more carefully. However, withdrawal of AZA could be contemplated in selected pediatric patients with CD sustaining CR for at least two years and achieved deep remission.

Received: 31 May 2021; Accepted: 14 December 2021
Published online: 11 January 2022

References
1. Cosnes, J. et al. Long-term evolution of disease behavior of Crohn’s disease. Inflamm. Bowel Dis. 8, 244–250 (2002).
2. Walters, T. D. et al. Increased effectiveness of early therapy with anti-tumor necrosis factor-α vs an immunomodulator in children with Crohn’s disease. Gastroenterology 146, 383–391 (2014).
3. Bouguen, G. et al. Efficacy and safety of tumor necrosis factor antagonists in treatment of internal fistulizing Crohn’s disease. Clin. Gastroenterol. Hepatol. 16, 628–636 (2020).
4. Colombel, J. F. et al. Infliximab, azathioprine, or combination therapy for Crohn’s disease. N. Engl. J. Med. 362, 1383–1395 (2010).
5. Khanna, R. et al. Early combined immunosuppression for the management of Crohn’s disease (REACT): A cluster randomised controlled trial. Lancet 366, 1825–1834 (2015).
6. Kang, B. & Cho, Y. Early biologic treatment in pediatric Crohn’s disease: Catching the therapeutic window of opportunity in early disease by treat-to-target. Pediatr Gastroenterol. Hepatol. Nutr. 21, 1–11 (2018).
7. Ashworth, L. A. et al. Lymphoma risk in children and young adults with inflammatory bowel disease: Analysis of a large single-center cohort. Inflamm. Bowel Dis. 18, 838–843 (2012).
8. Smith, M. A. et al. malignancy on thiopurine treatment with special reference to inflammatory bowel disease. Aliment Pharmacol. Ther. 32, 119–130 (2010).
9. Van Assche, G. et al. Withdrawal of immunosuppression in Crohn’s disease treated with scheduled infliximab maintenance: A randomized trial. Gastroenterology 134, 1861–1868 (2008).
10. Boyapati RK, Torres J, Palmela C et al. Withdrawal of immunosuppressant or biologic therapy for patients with quiescent Crohn’s disease. Cochrane Database Syst. Rev. 2018.
11. Vermeire, S. et al. Effectiveness of concomitant immunosuppressive therapy in suppressing the formation of antibodies to infliximab in Crohn’s disease. Gut 56, 1226–1231 (2007).
12. Levine, A. et al. ESPGHAN revised porto criteria for the diagnosis of inflammatory bowel disease in children and adolescents. J Pediatr Gastroenterol Nutr 58, 795–806 (2014).
13. Levine, A. et al. Pediatric modification of the Montreal classification for inflammatory bowel disease: The Paris classification. Inflamm. Bowel Dis. 17, 1314–1321 (2011).
14. Black, A. J. et al. Thiopurine methyltransferase genotype predicts therapy-limiting severe toxicity from azathioprine. Ann. Intern. Med. 129, 716–718 (1998).
15. Ye, B. D. et al. Efficacy and safety of biosimilar CT-P13 compared with originator infliximab in patients with active Crohn’s disease: An international, randomised, double-blind, phase 3 non-inferiority study. Lancet 393, 1699–1707 (2019).
16. Buer, L. C. et al. Switching from Remicade® to Remsima® is well tolerated and feasible: A prospective, open-label study. J. Crohns Colitis 11, 297–304 (2017).
17. Kang, B. et al. Long-term outcomes after switching to CT-P13 in pediatric-onset inflammatory bowel disease: A single-center prospective observational study. Inflamm. Bowel Dis. 24, 607–616 (2018).
18. Dervieux, T. et al. Liquid chromatography–tandem mass spectrometry analysis of erythrocye thiopurine nucleotides and effect of thiopurine methyltransferase gene variants on these metabolites in patients receiving azathioprine/6-mercaptopurine therapy. Clin. Chem. 51, 2074–2084 (2005).
19. Vande Casteele, N. et al. The relationship between infliximab concentrations, antibodies to infliximab and disease activity in Crohn’s disease. Gut 64, 1539–1545 (2015).
20. Steyerberg, E. W. et al. Internal validation of predictive models: Efficiency of some procedures for logistic regression analysis. J. Clin. Epidemiol. 54, 774–781 (2001).
21. Snapinn, S. M. & Jiang, Q. Illustrating the impact of a time-varying covariate with an extended Kaplan–Meier estimator. Am. Stat. 59, 301–307 (2005).
22. Levesque, B. G. et al. Converging goals of treatment of inflammatory bowel disease from clinical trials and practice. Gastroenterology 148, 37–51. e1 (2015).
23. Kang, B. et al. Mucosal healing in paediatric patients with moderate-to-severe luminal Crohn’s disease under combined immunosuppression: Escalation versus early treatment. J. Crohns Colitis 10, 1279–1286 (2016).

24. Gupta, A. et al. Treat to target: The role of histologic healing in inflammatory bowel diseases: A systematic review and meta-analysis. Clin. Gastroenterol. Hepatol. 19, 1800–1813. e4 (2021).

25. Peyrin-Biroulet, L. et al. Selecting therapeutic targets in inflammatory bowel disease (STRIDE): Determining therapeutic goals for treat-to-target. Am. J. Gastroenterol. 110, 1324–1338 (2015).

26. Choe, Y. H. J. & Medicine, F. Precision medicine for pediatric inflammatory bowel disease: A perspective. Precision Future Med. 4, 43–52 (2020).

27. Torres, J. et al. Systematic review of effects of withdrawal of immunomodulators or biologic agents from patients with inflammatory bowel disease. Gastroenterology 149, 1716–1730 (2015).

28. Reenaers, C. et al. Outcomes 7 years after infliximab withdrawal for patients with Crohn’s disease in sustained remission. Clin. Gastroenterol. Hepatol. 16, 234–2432 (2018).

29. Brooks, A. et al. Outcome of elective withdrawal of anti-tumour necrosis factor-α therapy in patients with Crohn’s disease in established remission. J. Crohns Colitis. 11, 1456–1462 (2017).

30. Song, J. H. et al. Long-term outcomes after the discontinuation of anti-tumor necrosis factor-α therapy in patients with inflammatory bowel disease under clinical remission: A Korean Association for the Study of Intestinal Disease Multicenter Study. Gut Liver 15, 752 (2021).

31. Scarallo L, Bolasco G, Barp J, et al. Anti-tumor necrosis factor-alpha withdrawal in children with inflammatory bowel disease in endoscopic and histologic remission. Inflamm. Bowel Dis. 2020.

32. Oussalah, A. et al. Predictors of infliximab failure after azathioprine withdrawal in Crohn’s disease treated with combination therapy. Am. J. Gastroenterol. 105, 1142–1149 (2010).

33. Roblin, X. et al. Azathioprine dose reduction in inflammatory bowel disease patients on combination therapy: An open-label, prospective and randomised clinical trial. Aliment. Pharmacol. Ther. 46, 142–149 (2017).

34. Doherty, G. et al. European Crohn’s and Colitis Organisation topical review on treatment withdrawal ‘exit strategies’ in inflammatory bowel disease. J Crohns Colitis 12, 17–31 (2018).

35. Drobne, D. et al. Withdrawal of immunomodulators after co-treatment does not reduce trough level of infliximab in patients with Crohn’s disease. Clin. Gastroenterol. Hepatol. 13, 514–5214 (2015).

36. Maser, E. A. et al. Association of trough serum infliximab to clinical outcome after scheduled maintenance treatment for Crohn’s disease. Clin. Gastroenterol. Hepatol. 4, 1248–1254 (2006).

37. Baert, F. et al. Influence of immunogenicity on the long-term efficacy of infliximab in Crohn’s disease. N. Engl. J. Med. 348, 601–608 (2003).

38. Kang, B. et al. Infliximab trough levels are associated with mucosal healing during maintenance treatment with infliximab in paediatric Crohn’s disease. J. Crohns Colitis 13, 189–197 (2019).

39. Papamichael, K. et al. Long-term outcome of patients with Crohn’s disease who discontinued infliximab therapy after clinical remission. Clin. Gastroenterol. Hepatol. 13, 1103–1110 (2015).

40. Louis, E. et al. Maintenance of remission among patients with Crohn’s disease on antimetabolite therapy after infliximab therapy is stepped. Gastroenterology 142, 63–705 (2012).

41. Vermeire, S. et al. Immunogenicity of biologics in inflammatory bowel disease. Therap. Adv. Gastroenterol. 11, 1756283X17750355 (2018).

42. Carman N, Mack DR, Benchimol ELJ. Therapeutic drug monitoring in pediatric inflammatory bowel disease. Curr Gastroenterol Rep 2018;20:1–13.

43. Ordás, I. et al. Anti-TNF monoclonal antibodies in inflammatory bowel disease: Pharmacokinetics-based dosing paradigms. Clin. Pharmacol. Ther. 91, 635–646 (2012).

44. Rosen, M. J. et al. Applying pharmacokinetics to optimise dosing of anti-TNF biologics in acute severe ulcerative colitis. Aliment Pharmacol. Ther. 41, 1094–1103 (2015).

45. Ben-Horin, S. & Chowers, Y. Loss of response to anti-TNF treatments in Crohn’s disease. Aliment Pharmacol. Ther. 33, 987–995 (2011).

46. Kansen, H. M. et al. Less anti-infliximab antibody formation in paediatric Crohn patients on concomitant immunomodulators. Gastroenterol. Nutr. 65, 425–429 (2017).

47. Colombel, J.-F. et al. Combination therapy with infliximab and azathioprine improves infliximab pharmacokinetic features and efficacy: A post hoc analysis. Clin. Gastroenterol. Hepatol. 17, 1525–15321 (2019).

48. Yarar, A. J. et al. Concentrations of 6-thioguanine nucleotide correlate with trough levels of infliximab in patients with inflammatory bowel disease on combination therapy. Clin. Gastroenterol. Hepatol. 13, 1118-1124.e3 (2015).

49. Mogensen, D. V. et al. A role for thiopurine metabolites in the synergism between thiopurines and infliximab in inflammatory bowel disease. J Crohns Colitis 12, 298–305 (2018).

50. van Walraven, C. et al. Time-dependent bias was common in survival analyses published in leading clinical journals. J Clin Epidemiol 57, 672–682 (2004).

Author contributions Guarantor of article: Eun Sil Kim. TJJ and ESK contributed to the conception and design of the study, acquisition, analysis, and interpretation of data, drafting of the initial manuscript, and critical revision of the manuscript for important intellectual contribution. Y-YK contributed to the analysis and interpretation of data. S-WK contributed to the study design, statistical analyses and interpretation of data, and critical revision of the manuscript for important intellectual contribution. SWS contributed to design of the study and interpretation of data. YHC and MJK contributed to the conception and design of the study, analysis and interpretation of data, and critical revision of the manuscript for important intellectual contribution. All authors approved the final version of the manuscript, had full access to all the data in the study, and had final responsibility for the decision to submit for publication.

Funding This work was supported by the National Research Foundation of Korea (NRF) Grant funded by the Korea government (MSIT) (No. 2020R1A2C2007192).

Competing interests The authors declare no competing interests.
Additional information
Correspondence and requests for materials should be addressed to Y.H.C. or M.J.K.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022