Evidence-based clinical practice guidelines for inflammatory bowel disease 2020

Hiroshi Nakase 1,2 · Motoi Uchino 1 · Shinichiro Shinzaki 1 · Minoru Matsuura 1 · Katsuyoshi Matsuoka 1 · Taku Kobayashi 1 · Masayuki Saruta 1 · Fumihito Hirai 1 · Keisuke Hata 1 · Sakiko Hiraoka 1 · Motohiro Esaki 1 · Ken Sugimoto 1 · Toshimitsu Fuji 1 · Kenji Watanabe 1 · Shiro Nakamura 1 · Nagamu Inoue 1 · Toshiyuki Itoh 1 · Makoto Naganuma 1 · Tadakazu Hisamatsu 1 · Mamoru Watanabe 1 · Hiroto Miwa 1 · Nobuyuki Enomoto 1 · Tooru Shimosegawa 1 · Kazuhiko Koike 1

Received: 25 March 2021 / Accepted: 25 March 2021 / Published online: 22 April 2021 © The Author(s) 2021

Abstract Inflammatory bowel disease (IBD) is a general term for chronic or remitting-relapsing inflammatory diseases of the intestinal tract and generally refers to ulcerative colitis (UC) and Crohn’s disease (CD). Since 1950, the number of patients with IBD in Japan has been increasing. The etiology of IBD remains unclear; however, recent research data indicate that the pathophysiology of IBD involves abnormalities in disease susceptibility genes, environmental factors and intestinal bacteria. The elucidation of the mechanism of IBD has facilitated therapeutic development. UC and CD display heterogeneity in inflammatory and symptomatic burden between patients and within individuals over time. Optimal management depends on the understanding and tailoring of evidence-based interventions by physicians. In 2020, seventeen IBD experts of the Japanese Society of Gastroenterology revised the previous guidelines for IBD management published in 2016. This English version was produced and modified based on the existing updated guidelines in Japanese. The Clinical Questions (CQs) of the previous guidelines were completely revised and categorized as follows: Background Questions (BQs), CQs, and Future Research Questions (FRQs). The guideline was composed of a total of 69 questions: 39 BQs, 15 CQs, and 15 FRQs. The overall quality of the evidence for each CQ was determined by assessing it with reference to the Grading of Recommendations Assessment, Development and Evaluation (GRADE) approach, and the strength of the recommendation was determined by the Delphi consensus process. Comprehensive up-to-date guidance for on-site physicians is provided regarding indications for proceeding with the diagnosis and treatment.

Keywords Inflammatory bowel disease · Steroid · Immunomodulators · Biologics

Introduction

1. Purpose of the revised guidelines

The purpose of these practice guidelines is to improve patient outcomes by providing appropriate practice measures for health care providers and patients for the treatment of inflammatory bowel disease (IBD).

2. Basic policy

In accordance with the policy of the previous guideline, the basic concepts of the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system, which has been used in many foreign guidelines, were incorporated as much
as possible to create a medical index that emphasizes the totality of evidence from systematic reviews[1].

3. Development method
In the actual preparation of the guideline, we held a series of face-to-face preparation committee meetings and e-mail deliberations, prepared draft questions, and formulated items. The Clinical Questions (CQs) of the previous guideline were completely revised and categorized as follows:

a. Background questions (BQs): those for which conclusions are already clear, and those for which a consensus has already been reached in previous guidelines.

b. CQs: questions that affect the direction of medical treatment and for which recommendations and evidence levels can be determined by an exhaustive literature search.

c. Future research questions (FRQs): questions for which a recommendation and level of evidence cannot be determined by the current exhaustive literature search (no evidence exists.)

This guideline includes 69 questions: 39 BQs, 15 CQs, and 15 FRQs. A literature search was created for each question, and for CQs and FRQs, the search period was from 1983 to April 2019 for English articles and from 1983 to May 2019 for Japanese articles. The Japan Association of Medical Libraries was commissioned to conduct a literature search in PubMed and the Central Journal of Medicine. For the CQs, three meta-analyses were prepared and have just published. For BQs, references were manually searched by each committee member, and no search period was applied. The statement and commentary were completed. The overall quality of the evidence for each CQ was determined by assessing it with reference to the GRADE approach (Table 1). The strength of the recommendation was determined by the the drafting committee using the Delphi method (Table 2). The appropriateness of the wording of the statement was independently evaluated by 17 members of the drafting committee. A rating scale of 9 (9 = most appropriate, 1 = most inappropriate) was used, with a median rating of 9 or 8 indicating a strong recommendation and a median rating of 7 indicating a weak recommendation. As a result, a consensus recommendation (median value of 7 or higher) was obtained for all statements, but there were variations in the ratings for some statements, requiring reevaluation to reach a consensus.

The draft was submitted to the evaluation committee, and after the evaluation comments were collected, feedback was given to the drafting committee members in charge, and necessary revisions were made. This process was repeated once more, and the final draft was developed. The final draft was posted on the website of the Japanese Society of Gastroenterology from August 3 to 17, 2020, for public comment.

4. Application of guidelines
This guideline is intended to support decision-making in clinical practice by describing standard information on the disease concept, diagnosis, treatment, and follow-up of IBD. The Japanese Gastroenterological Association (JGAA) and this Guideline Development and Evaluation Committee are not responsible for the results of individual treatment. The Japanese Society of Gastroenterology and the Committee for the Preparation and Evaluation of this guideline are not responsible for the results of individual treatment. The contents of this guideline are not to be used as a legal basis for medical litigation.

5. Structure of the medical algorithm
In this guideline, the following treatment algorithm is presented in a flowchart (nine figures).

The algorithm is simplified to the maximum extent possible, although treatment may be complicated in IBD, where treatment options vary by disease state.

**Definition and pathophysiology of IBD**

**BQ 1 What is IBD?**

| Table 1 Quality of evidence | |
|-------------------------------|----------------------------------|
| A: High quality evidence    | We are confident that the true effect approximates the effect estimates |
| B: Moderate quality evidence | Moderate confidence in the effect estimates. The true effect is approximately close to the effect estimate, but it may be substantially different |
| C: Low quality evidence      | Confidence in the estimated effect is limited |
|                              | The true effect may be substantially different from the effect estimate |
| D: Very-low-quality evidence | Effect estimates are largely unreliable |
|                              | The true effect is likely to be substantially different from the effect estimate |
Statements

- IBD is a general term for chronic or remitting/remitting inflammatory diseases of the intestinal tract and generally refers to ulcerative colitis (UC) and Crohn’s disease (CD).
- UC is a diffuse, nonspecific inflammation of unknown origin that continuously damages the colonic mucosa from the rectal side, often leading to erosions and ulcers.
- CD is a chronic inflammatory disease of unknown etiology characterized by noncontiguous distributed, all-stratified granulomatous inflammation and fistulae.

These statements and supplementary were made with reference to [2, 3]

Supplementary information

Cases of enteritis that cannot be differentiated as UC or CD are referred as follows:

1. IBD unclassified (IBDU): this term is used for patients who do not have a surgical specimen available (i.e., have not undergone surgery) and whose diagnosis is difficult to make despite a combination of clinical, endoscopic and histological findings.

2. Indeterminate colitis: as a rule, a surgical specimen is used for the diagnosis of indeterminate colitis and is used in cases with characteristics of UC and CD.

BQ 2. What is the epidemiology of IBD in Japan?

Statements

- The number of IBD patients is estimated to be more than 220,000 for UC and more than 70,000 for CD, based on the current number of medical certificates issued.
- Both UC and CD occur at a relatively young age, with a high incidence in the late teens to early 30 s.

These statements were made with reference to [4, 5]

BQ 3. What are the factors that contribute to and exacerbate IBD?

Statements

- Multiple loci have been reported to be associated with the development of UC and CD.
- Although the cause of UC/CD is unknown, an association with certain dietary factors has been reported.
- Smoking and appendicectomy have been reported to be protective against UC.
- Current smoking has been reported to be a risk factor for the development of CD.
- Oral contraceptives and nonsteroidal anti-inflammatory drugs (NSAIDs) have been reported to be associated with the development of IBD.
- The pathogenesis of IBD is associated with dysbiosis.

These statements were made with reference to the following information and paper [6–15]

Diagnosis

BQ 4. How do you proceed with the diagnosis of IBD?

Statements

- The diagnosis of IBD is established by suspicion based on the information obtained in the medical interview, characteristic findings of IBD in the physical examination and the endoscopic and other imaging study results typical of IBD.
- Persistent or recurrent bloody diarrhea with abdominal pain and frequent bowel movements should lead to the suspicion of IBD, especially in young patients.
- A problem in the differentiation of IBD is infectious enteritis.
- Chronic abdominal pain, diarrhea, bloody stools, weight loss, fever, and anal lesions should lead to the suspicion of IBD, especially in young patients.

Table 2: Strength of recommendation

| Grade of recommendation | Criteria (mean Delphi score) | Interpretation |
|-------------------------|-------------------------------|---------------|
| 1. Strong recommendation | 8–9                           | Recommendation to do |
| 2. Weak recommendation  | 7                             | Suggest to do  |

These statements were made with reference to [179x702]
BQ 5. What are the diagnostic criteria for IBD?

Statement

- A diagnosis of IBD is made following the diagnostic criteria of the Ministry of Health, Labour, and Welfare’s “Research on Intractable Inflammatory Bowel Disorders”.

These statements were made with reference to [2, 3]. Please refer to Figs. 1 and 2.

Fig. 1 Diagnostic approach for ulcerative colitis. Mild to moderate active left-sided colitis type (not extending beyond the sigmoid colon) and proctitis type

Fig. 2 Diagnostic approach to Crohn’s disease
BQ 6. What is the pathology, classification, and severity of UC?

Statements

- There are two phases of UC: the active phase, in which symptoms are present, and the remission phase, in which symptoms disappear.

- UC can be divided into three types according to the extent of the lesion: "proctitis", "left-sided colitis" (up to the splenic flexure), and "total colitis".
The severity of UC is classified as "mild", "moderate", or "severe" based on clinical symptoms, signs, and blood tests (Table 5).

Depending on the disease’s clinical course of UC, the disease is classified as relapsing–remitting, chronically persistent, acutely fulminant, or first attack types. These statements and supplementary information were made with reference to [16–20]. Please refer to Table 5.

### Supplementary information

The partial Mayo score is a four-point scale that incorporates the frequency of bowel movements, rectal bleeding, and the physician’s general assessment of the patient’s condition. A score of 0–1 indicates remission, 2–4 indicates mild disease, 5–7 indicates moderate disease, and 8 or more indicates severe disease (Table 6).

### Table 5 Classification of severity of ulcerative colitis [16]

|                  | Severe | Moderate | Mild |
|------------------|--------|----------|------|
| (1) Bowel movements | ≥ 6    | ≤ 4      | (1) Bowel movements |
| (2) Blood in stools     | (+++)  | (++)~(--) | (2) Blood in stools |
| (3) Pyrexia          | ≥ 37.5 °C | Between mild and moderate | (3) Pyrexia |
| (4) Pulse            | ≥ 90/min | No      | (4) Pulse |
| (5) Anemia           | Hb ≤ 10 g/dL | No | (5) Anemia |
| (6) ESR or CRP       | ≥ 30 mm/h | Normal | (6) ESR |
|                   |         | Normal or CRP |

Patients are classified as severe if they present both (1) and (2) plus at least one of (3) or (4)

While satisfying 4 or more out of 6 features, patients with extremely severe symptoms are classified as fulminant, and further divided into acute fulminant or relapsing fulminant types. Diagnostic criteria of fulminant colitis: all of the below:

1. Satisfy criteria of severe cases
2. Bloody diarrhea 15 or more times day continuously
3. Persistent high fever ≥ 38.0 °C
4. White blood cell count ≥ 10,000/mm³
5. Severe abdominal pain

ESR, erythrocyte sedimentation rate; CRP, C-reactive protein

### Table 6 Partial Mayo score [19]

| Mayo items               | Clinical assessment            |
|--------------------------|-------------------------------|
| Stool frequency          | 0 = Normal                    |
|                          | 1 = 1–2 stools/day more than normal |
|                          | 2 = 3–4 stools/day more than normal |
|                          | 3 = > 4 stools/day more than normal |
| Rectal bleeding          | 0 = None                       |
| a                        | 1 = Visible blood with stool less than half the time |
|                          | 2 = Visible blood with stool half of the time or more |
|                          | 3 = Passing blood alone        |
| Physician rating of disease activity | 0 = Normal |
|                          | 1 = Mild                       |
|                          | 2 = Moderate                   |
|                          | 3 = Severe                     |

*aA score of 3 for bleeding required patients to have at least 50% of bowel movements accompanied by visible blood and at least one bowel movement with blood alone*
BQ 7. What is the pathology, classification, and severity of CD?

Statements

- The most common sites of CD are in the small and large intestine (especially in the ileum) and perianal region and are classified as “ileal-type”, “colonic-type”, and “ileocolonic-type”.
- It has been proposed to classify the disease pattern of CD in three ways: “non-stricturing, non-penetrating type”, “penetrating type”, and “stricturing type” (Table 7).
- The Crohn’s Disease Activity Index (CDAI) (Table 8), the International Organization for the Study of Inflammatory Bowel Disease (IOIBD) index, and the Harvey-Bradshaw index have been proposed as CD activity indicators. Nevertheless, they have not been widely used in general practice.

These statements were made with reference to [16, 21–24]. Please refer to Tables 7 and 8.

Endoscopy and other imaging modalities

BQ 8. What is the role of endoscopy in the diagnosis and treatment of UC?

Statements

- Colonoscopy is necessary to confirm the diagnosis if UC is suspected based on clinical findings.
- Colonoscopy is used to confirm the diagnosis of UC and to evaluate the severity of the disease, determine the effectiveness of treatment, and conduct surveillance for carcinogenesis.

These statements were made with reference to [2, 16, 19, 25–33].

BQ 9. What are nonendoscopic, noninvasive tests used in the diagnosis of UC?

Statement

- Noninvasive abdominal ultrasound (US), computed tomography (CT) and magnetic resonance imaging (MRI) scans are used to assess activity and confirm complications before and after treatment.

This statement was made with reference to [16, 26, 34–38].

BQ 10. What is the role of endoscopy in the diagnosis and treatment of CD?

Statements

- If CD is suspected, lower gastrointestinal (GI) endoscopy (including observation of the ileum’s distal end) and histopathological examination with biopsy should be performed.
- Upper GI endoscopy is recommended, especially if the diagnosis cannot be confirmed by lower GI endoscopy or if the patient complains of upper GI symptoms.
- Endoscopy is performed when necessary to confirm the diagnosis of CD and evaluate the severity of the disease, determine the effectiveness of treatment, and conduct surveillance for carcinogenesis.
- Balloon-assisted enteroscopy or small-bowel capsule endoscopy (SBCE) may be useful for the close examination and follow-up of small bowel lesions in CD.

These statements were made with reference to [16, 25, 39–49].

CQ 1. Is SBCE useful for the assessment of small bowel disease activity in CD?

Recommendation

- SBCE is as useful as CT enterography (CTE) and magnetic resonance enterography (MRE) for the assessment of small bowel disease activity or postoperative recurrence in patients with CD -confirmed bowel patency. [Strong recommendation, moderate-quality evidence]
Commentary

In the meta-analysis comparing the diagnostic yield of active small bowel lesions [50], SBCE was reported to demonstrate a better diagnostic yield when comparing small-bowel follow-through or enteroclysis. The yield was no different compared with that of CTE or MRE. In cross-sectional studies comparing the yields between SBCE and MRE [51, 52], SBCE has been reported to show better diagnostic yields, especially in the upper part of the small bowel [52]. However, the results should be interpreted cautiously, because the types and the severity of small-bowel lesions detected by SBCE are different from those detected by cross-sectional studies. Another meta-analysis demonstrated that SBCE, MRE and US showed favorable diagnostic yields of anastomotic recurrence in patients with CD who underwent ileocecal resection [53]; however, the difference in the definition of postoperative recurrence among the studies could cause selection bias.

The diagnostic yield of SBCE for CD reportedly varies from 20 to 86% in suspected CD [54]. Such a difference might be because the diagnosis of CD cannot be confirmed by SBCE findings alone, and because the definition of small-bowel lesions was different among the studies. SBCE findings that can be useful for the distinction of CD have been recently proposed [49]. It is necessary to determine the usefulness of SBCE for the diagnosis of CD based on certain SBCE findings. Another report concluded that SBCE is not recommended for patients with negative CTE or enteroclysis findings when considering cost-effectiveness [55].

The association of SBCE findings with clinical disease activity and biomarkers has been scarcely investigated. A cross-sectional study reported a positive association [56], while another cohort study failed to show a correlation between the severity of SBCE and of biomarkers (C-reactive protein (CRP), erythrocyte sedimentation rate (ESR), fecal calprotectin). The risk of capsule retention has been reportedly high in both established CD (5–13%) and suspected CD (4–13%), however, the risk can be minimized with the application of a patency capsule (PC) beforehand. While the confirmation of small-bowel patency by using a PC is thus recommended, we should consider the impaction of the PC itself.

BQ 11. What are imaging studies other than endoscopy are used to diagnose CD?

Statements
- Radiographic and other imaging studies are used to determine treatment strategy and to determine the extent, severity, and complications of the lesion.
- US, CT, and MRI are mainly used to evaluate patient disease activity before and after treatment and to check for complications.

These statements were made with reference to [16, 26, 35, 37, 47, 57–64].

CQ 2. Is MRI useful for the evaluation of CD disease activity?

Recommendation
- The use of MRE or magnetic resonance enterocolonography (MREC) is recommended for monitoring intestinal disease activity, evaluating mucosal healing and extraluminal disease, and evaluating treatment response. [Strong recommendation, moderate-quality evidence]

Commentary

MRE and MREC are useful in assessing for small-bowel lesions that are difficult to visualize with endoscopy. MRE and MREC are able to detect lesions in the small and large bowel with a high degree of accuracy, and are useful for diagnosis, monitoring for disease activity, and evaluating treatment response in CD [65]. The presence of edema, wall thickening of more than 3 mm, contrast enhancement, stenosis, and fistula are assessed [66]. A meta-analysis of MRE and MREC studies reported a sensitivity of > 80% and specificity of > 90% for the detection of inflammation and a sensitivity of > 90% and specificity of > 95% for the
detection of intestinal damage such as abscesses and fistulae [67].

Several scores have been developed for the assessment of disease activity, the most frequently validated of which is the Magnetic Resonance Index of Activity (MaRIA) [68]. The MaRIA was shown to correlate well with the CD Endoscopic Index of Severity (CDEIS) both before and after treatment in a prospective study, detecting endoscopic mucosal healing with a sensitivity of 85% and specificity of 78% [69]. In addition, the modified MaRIA correlated well with the Simple Endoscopic Score for Crohn’s Disease (SES-CD), and a sensitivity of 87% and specificity of 86% were reported for mucosal healing [70]. In recent years, several simpler scores have been developed, all of which show a high correlation with endoscopy and the MaRIA [71, 72]. The presence of disease activity upon MRE significantly correlates with relapse, including postoperative relapse, and surgery [73, 74]. For disease monitoring, modalities such as MRE/MREC and intestinal US are recommended, especially in cases where repeated examinations are required and in patients under 35 years of age, where radioactive exposure should be minimized [75]. MRE/MREC has been associated with problems of access and training, and a consensus statement has been issued regarding the procedure, imaging sequence and interpretation [76].

Biomarkers

BQ 12. Is fecal calprotectin testing useful for the differential diagnosis of IBD?

Statement

- Fecal calprotectin testing is useful for differentiating between organic intestinal diseases such as IBD and functional intestinal diseases such as irritable bowel syndrome (IBS).

This statement was made with reference to [65, 77–79].

BQ 13. Are fecal calprotectin tests and fecal immunochromic tests (FITs) useful for assessing disease activity in UC patients in remission?

Statement

- Fecal calprotectin tests and FITs (hemoglobin concentrations in feces measured by using an antibody for human hemoglobin) are useful for evaluating the disease activity in UC patients in the remission stage.

This statement was made with reference to [80–84].

Treatment

BQ 14. What is “Treat to Target” (T2T) in IBD treatment?

Statements

- The concept of T2T is when physicians and patients discuss treatment goals and review treatment options at appropriate intervals using a comprehensive activity index to achieve early clinical remission or low disease activity.
- Prospective observational studies are required to determine whether the treatment goals for UC and CD proposed by the Selecting Therapeutic Targets in Inflammatory Bowel Disease (STRIDE) program contribute to improved patient quality of life.

These statements and supplementary information were made with reference to [30, 31, 85–90].

Supplementary information

The STRIDE program has been implemented at the IOIBD [87]. The aim of this program is to use an evidence-based consensus to identify therapeutic targets that would be useful in the implementation of T2T therapy in clinical practice. The results of the STRIDE program are as follows: (1) T2T in UC is aimed at achieving no rectal bleeding, improving diarrhea, improving defecation habits (decrease in frequency) and improving findings on endoscopy (Mayo score 0–1), with histological remission as an adjunct goal. T2T in CD is aimed at improving abdominal pain and diarrhea, improving defecation habits (decrease in frequency) and improving ulceration findings on ileal and colonoscopy, or improving inflammatory findings on cross-sectional imaging (CT/MRI/US) in patients in whom lesions cannot be assessed by lower endoscopy up to the terminal ileum. Calprotectin levels can serve as adjunctive targets.

Current treatment strategy of IBD

- Curative medical therapy has not been established for IBD patients.
- At present, medical treatment goals are early induction of remission and long-term maintenance to prevent relapse.
- In the active stage, it is necessary to accurately diagnose the patient’s general condition and the extent of the disease and proceed with treatment based on the treatment guidelines proposed by the Ministry of
Health, Labour and Welfare Grant-in-Aid for Scientific Research on Intractable Diseases, “Research on Intractable Inflammatory Bowel Disorders”.

- In severe cases, surgery should always be considered a treatment option, and medical treatment should be carried out in close communication with the surgeon (please refer to Figs. 3, 4, 5, 6, 7, 8 and 9).
5-Aminosalicyclic acid (ASA)

**BQ 15. What are the benefits and precautions for the use of 5-ASA drugs in the treatment of IBD?**

**Statements**

- 5-ASA is effective in inducing remission in active UC and preventing the relapse of UC in remission.
- The effect of 5-ASA on CD is generally lower than that on UC, and although 5-ASA has a suppressive effect on active CD, it has not been proven to be effective in maintaining remission.
- It should be kept in mind that there are cases of 5-ASA intolerance.

These statements and supplementary information were made with reference to [91–104].

**Supplementary information**

Although 5-ASA drugs have relatively few side effects, they can cause abdominal pain, fever, joint pain, and bloody stools, making it appear as if the IBD itself is
worsening. Saito et al. reported that the drug-induced lymphocyte stimulation test (DLST) has low sensitivity but high specificity for the diagnosis of 5-ASA allergy [103]. Therefore, in addition to 5-ASA allergy, there are cases of 5-ASA intolerance without DLST positivity. 5-ASA intolerance (allergy) should be suspected when the clinical course after the start of 5-ASA administration is unnatural.

**BQ 16. Are oral and topical 5-ASA formulations useful for the induction of remission in mild to moderately active UC?**

**Statements**

- Oral and local 5-ASA formulations are useful for the induction of remission in mild to moderately active UC.
- The combination of oral and enteral 5-ASA formulations is useful when a more potent effect on mild to moderately active distal UC is needed.
These statements were made with reference to [93, 105–108].

**BQ17. Is maintenance treatment with 5-ASA drugs for UC in remission useful in maintaining clinical and endoscopic remission?**

**Statement**

- Maintenance treatment with 5-ASA drugs for UC in remission is useful in maintaining clinical and endoscopic remission.

This statement was made with reference to [92, 94].

**BQ18. Are 5-ASA suppositories useful for the induction of remission in mild to moderately active UC of the proctitis type?**

**Statement**

- 5-ASA suppositories are useful for the induction of remission in mild to moderately active UC of the proctitis type.

This statement was made with reference to [109–111].

**CQ3. What is the appropriate maintenance dose of 5-ASA for treating UC?**

**Recommendation**

- It is recommended that the dose of 5-ASA in the maintenance therapy for UC be 2 g or more. [Strong recommendation, high-quality evidence]

**Commentary**

The latest Cochrane analysis [94] shows that the administration of 2 g or more of 5-ASA is effective for maintaining remission. There was no difference in efficacy between the various 5-ASA preparations. Regarding pH-dependent mesalazine delayed-release preparation (Asacol®), it has been shown that there is no statistically significant difference in the efficacies of maintaining remission between 1.2 g/day and 2.4 g/day and between 2.4 g/day and 4.8 g/day [112, 113]. However, for patients aged less than 40 years or those with total colitis, it has been reported that the administration of 4.8 g/day of Asacol® significantly increased the remission rate and duration of remission compared with the administration of 2.4 g/day of Asacol® [113]. For time-dependent mesalazine controlled-release preparations (Pentasa®), it has been reported that there is no statistically significant difference in the efficacies of maintaining remission between 1.5 g/day and 3.0 g/day [114]. The dose range for maintaining remission in Multi Matrix System™ (MMX) mesalazine has not yet been investigated.

**CQ 4. Does 5-ASA treatment reduce the risk of UC-associated colorectal cancer (CRC)?**

**Recommendation**

- It is recommended that 5-ASA treatment reduces the risk of UC-associated CRC. [Strong recommendation, low-quality evidence]
Commentary

One of the risk factors for the development of UC-associated CRC is the length of the disease period [115], and successful treatment of UC is thought to reduce the risk of UC-associated CRC [116]. Several meta-analyses have demonstrated the protective effect of 5-ASA on IBD-associated CRC [117–119]; however, there is also a report that 5-ASA does not prevent the development of IBD-associated CRC [120]. Recently, Bonovas et al. [121] conducted a comprehensive meta-analysis of 31 studies and reported that exposure to 5-ASA reduces the risk of developing IBD-associated CRC to 43%. Twenty-one of these studies had limited data on UC but showed a 50% reduction in the risk of UC-associated CRC associated with 5-ASA use. In addition, the analysis of two reports describing doses of less than 1.2 g/day showed no significant effect. Furthermore, since it has been reported that inflammation itself is an independent risk factor for the development of CRC in UC patients [122], the anti-inflammatory effect of 5-ASA is also considered to be involved in reducing the risk of UC-associated CRC.

FRQ 1. Is the combined use of 5-ASA with biologics or immunomodulators (IMs) for CD in remission useful?

Statement

- The combination of 5-ASA with biologics or IMs may be considered as a maintenance therapy for CD in remission, but its usefulness has not been proven.

Commentary

5-ASA drugs have not been shown to be more effective than the placebo group in 2010 and 2016 meta-analyses of the maintenance of remission after induction therapy [100]. In addition, the American College of Gastroenterology (ACG) Clinical Guideline states that oral 5-ASA drugs have not been demonstrated to be effective in maintaining CD remission and are not recommended for long-term treatment [75].

The integrated analysis of nine randomized controlled trials (RCTs) in a 2011 Cochrane Review demonstrated that the effect of 5-ASA preparations on maintaining remission in CD patients after surgical treatment as an induction therapy was found to be slightly significant in suppressing recurrence compared to the effect of placebo group [101]. At present, the role of the use of 5-ASA alone is often negative for maintaining CD remission but the additive and synergistic effects of the combined use of 5-ASA with biologics or IMs for CD in remission are unclear.

Nutritional therapy

BQ 19. What are the benefits and caveats of nutritional therapy in the treatment of IBD?

Statements

- It is not clear that nutritional therapy, such as enteral nutrition and central venous nutrition alone, effectively induces remission in UC. Nevertheless, drug therapy and blood-cell removal therapy should be the mainstay of UC treatment.
- Enteral nutrition is an effective remission-inducing therapy for active CD. Enteral nutrition therapy is safe, but acceptance of the treatment can be difficult.
- Home enteral nutrition is effective in maintaining the remission of CD.

These statements were made with reference to [3, 16, 26, 123–132].

Cytapheresis

BQ 20. What are the benefits and cautions of Cytapheresis (CAP) in the treatment of IBD?

Statements

- In patients with moderate to severely active UC, blood-cell removal therapy should be considered a treatment option.
- Twice-weekly intensive therapy can reduce the time to the initiation of remission and improve the remission rate compared to once-weekly treatment in both UC and CD.
- In active CD, patients with colorectal lesions, granulocyte and monocyte adsorption apheresis (GMA) should be considered in combination with pharmacological and nutritional therapy when they are ineffective or inapplicable.

These statements and supplementary information were made with reference to [133–145].
**Supplementary information**

Although CAP is a safe treatment with few side effects, it is difficult to perform in patients in whom access to peripheral blood vessels is difficult (including patients with dehydration and anemia) because effective blood flow cannot be obtained. In addition, it is known from clinical practice that the therapeutic effect of CAP is poor in severe cases of UC, such as those with extensive ulcerations. Although it is not covered by insurance, there is a report on the effect of CAP on maintaining the remission of UC [141], and there is only one case report on the effect of CAP on maintaining the remission of CD [142].

**Immunosuppressive drugs and biologics**

**BQ 21. What are the precautions to take when using immunosuppressive drugs?**

**Statements**

- The use of overlapping immunosuppressive therapy should be done with caution, taking into account the risk of infection and other risks.
- In patients infected with hepatitis B virus (HBV) (carriers and those previously infected), the risk of developing hepatitis B due to the HBV reactivation should be considered after the initiation of immunosuppressive drugs.
- Anti-TNF-α antibody therapy should be used with caution because of the risk of tuberculosis complications.

These statements were made with reference to [146, 147].

**Corticosteroids**

**BQ 22. What is the usefulness and precautions for the use of corticosteroids in IBD treatment?**

**Statements**

- Corticosteroids have potent anti-inflammatory effects and are useful for inducing the remission of UC and CD.
- Corticosteroids are not useful for maintaining remission because of long-term administration side effects.
- After the initiation of corticosteroid therapy, it is preferable to reduce the dose to less than 10 mg/day of prednisolone (PSL) equivalent (3 mg/day for budesonide) within three months.

These statements were made with reference to [148–154].

**BQ 23. Is budesonide foam useful for UC?**

**Statement**

- Budesonide enema is useful for mild to moderate UC from the rectum to the sigmoid colon.

This statement was made with reference to [155–157].

**BQ 24 Are steroids (PSL, budesonide) useful for induction therapy for CD?**

**Statements**

- Budesonide is useful for mildly to moderately active ileocecal CD (ileum to ascending colon).
- The Administration of oral steroids (PSL 40 mg/day) is useful for moderately active ileocecal CD
- Intravenous administration of steroids (PSL 40–60 mg/day) is effective for severely active CD after excluding infections.
- The administration of steroids is involved in the development of perforating complications (abscesses and fistulas), therefore, it is generally contraindicated in such cases.

These statements were made with reference to [75, 148, 149, 153, 158–161].

**Immunomodulators**

**BQ 25. What are the usefulness and precaution for the use of immunomodulators in the treatment of IBD?**

**Statements**

- Azathioprine (AZA)/6-mercaptopurine (6-MP) treatment effectively prevents relapse in UC in remission, especially in steroid-dependent patients and those unable to maintain remission with 5-ASA.
- AZA/6-MP treatment is effective in maintaining remission for CD patients who achieve remission.
- Adverse effects of AZA/6-MP treatment include nausea and other GI symptoms, myelosuppression, alopecia, and pancreatitis.

These statements were made with reference to [162–180].
CQ 5. Is the NUDT15 gene R139C variant useful for predicting acute severe leukopenia induced by thiopurine?

Recommendation

• The NUDT15 gene R139C variant is useful for predicting acute severe leukopenia and severe alopecia induced by thiopurine, and we recommend confirming whether the NUDT15 gene R139C variant is present before the initiation of thiopurine. [Strong recommendation, high-quality evidence]

Commentary

In 2014, a study from Korea revealed that a single-nucleotide polymorphism in the nucleoside diphosphate-liked moiety X-type motif 15 (NUDT15) gene, in which C at position 415 changes to T, was strongly associated with acute severe leukopenia induced by thiopurines [174]. This polymorphism converts the 139th amino acid from arginine (Arg; R) to cysteine (Cys; C) (R139C variant). This was also confirmed in Japanese patients [176]. Acute severe leukopenia is inevitable in patients homozygous for the NUDT15 gene R139C variant. These patients also develop severe alopecia. In a large multicenter study conducted in Japan involving 1,291 patients previously treated with thiopurine (the MENDEL study), all 49 patients homozygous for the NUDT15 gene R139C variant discontinued thiopurine early due to adverse events (AEs) [181]. Among Japanese individuals, the frequency of homozygotes (Cys/Cys) for the NUDT15 gene R139C variant is approximately 1%, and the frequency of heterozygotes (Arg/Cys) is approximately 20%.

CQ 6. Is thiopurine effective for the prevention of postoperative recurrence of CD?

Recommendation

• We propose the use of thiopurines for the prevention of postoperative recurrence of CD. [Weak recommendation, moderate-quality evidence]

Commentary

Thiopurines, such as azathioprine (AZA) and 6-mercaptopurine (6-MP), are more effective than placebo in preventing postoperative clinical recurrence of patients with CD [166, 182, 183]. However, whether thiopurine is superior to placebo in preventing endoscopic recurrence after surgically induced remission in CD is controversial [166, 183]. The superiority of thiopurines over 5-ASA compounds in preventing postoperative recurrence of CD has not yet been established [183, 184]. Further studies are needed to (i) compare the efficacy between thiopurines vs. molecular-target drugs (e.g., TNF antagonist), and (ii) determine whether thiopurines on molecular-target drugs provide an additional effect, in preventing postoperative recurrence of CD.

Note: Currently, 6-MP is not covered by insurance for the treatment of IBD in Japan.

FRQ 2. Is thiopurine associated with an increased incidence of lymphoma in Asian IBD patients?

Statement

• The risk of developing lymphoma associated with thiopurine may be lower in Asian IBD patients than in Caucasian patients, but further studies are warranted.

Commentary

A large French cohort study reported that the incidence rate of lymphoproliferative disease was 0.90/1000 patient-years (95% confidence interval (CI) 0.50–1.49) in patients receiving thiopurine compared with 0.26/1000 patient-years (95% CI 0.10–0.57) in patients not receiving thiopurine. The adjusted hazard ratio was 5.28 (95% CI 2.01–13.9) [169]. No prospective cohort study has examined the incidence of lymphoma associated with thiopurine in Asian IBD patients. A Korean retrospective cohort study reported a standardized incidence ratio for lymphoma of 2.03 (95% CI 0.81–4.18) for all patients versus 5.93 (95% CI 6.16–15.18) for patients receiving thiopurine [185]. A questionnaire survey in Japan reported that 28 out of 36,939 patients had hematological malignancies, including 10 patients with lymphoma and the odds ratios for the incidence of hematological malignancies associated with thiopurine were 1.37 (95% CI 0.30–6.24) for UC and 1.86 (95% CI 0.60–5.78) for CD [172]. On the other hand, a recent Japanese study using a nationwide administrative database reported no increase in the incidence of lymphoma associated with thiopurine in patients with IBD [186].

Calcineurin inhibitors

BQ 26. What are the benefits and precautions for the use of calcineurin inhibitors in the treatment of IBD?

Statements

• Consider the use of intravenous cyclosporine (CsA) for severe UC that does not respond to steroid therapy.
• Oral tacrolimus should be considered for patients with severe UC who do not respond to steroids.

These statements were made with reference to [187–192].

FRQ 3. Is tacrolimus treatment effective for CD?

Statement

• Tacrolimus treatment is useful for patients with CD, but further research results will need to be accumulated.

Commentary

Few RCTs have reported the efficacy and AEs of tacrolimus (Tac) in patients with refractory CD, and only case studies have been reported [193, 194]. Recently, a systematic review and meta-analysis of the therapeutic effects and AEs of Tac in patients with CD was conducted [195]. Based on the case studies, systematic reviews and meta-analyses to date, Tac therapy is effective for patients with CD, and Tac therapy can be considered an option for patients with active CD. However, few RCTs have been conducted to accurately evaluate the efficacy of this therapy, and further clinical studies are needed.

Note: Currently, Tac is not covered by insurance for the treatment of CD in Japan.

Anti-TNF-α agents

BQ 27. What are the benefits and precautions for the use of anti-TNF-α agents in the treatment of IBD?

Statements

• Anti-TNF-α agents effectively induce and maintain remission in moderately to severely active UC that is either steroid-resistant or steroid-dependent.
• Anti-TNF-α agents are effective in inducing and maintaining the remission of pro-inflammatory CD.

These statements were made with reference to [2, 20, 146, 196–210].

BQ 28. Is there a difference in the effectiveness of Infliximab (originator) and biosimilars in introducing and maintaining remission?

Statement

• There is no difference in the effectiveness of Infliximab (originator) and biosimilars in inducing and maintaining remission.

This statement and supplementary information were made with reference to [211–213].

Supplementary information

While the use of CT-P13 should be considered from the viewpoint of medical cost-effectiveness, the nocebo effect that occurs when switching to a biosimilar (the effect of treatment that should have been obtained is affected due to anxiety and poor impression about the use of biosimilar on the part of patients) has become an issue [211–213]. In this regard, physicians and other healthcare professionals (nurses and pharmacists) should carefully explain the use of biosimilar when prescribing them to patients.

CQ 7. Is retreatment with the same anti-TNF-α agent effective and safe for the relapse after one discontinuation?

Recommendation

• Retreatment with the same anti-TNF-α agent for the relapse after discontinuation is recommended because it is generally effective and safe. [Weak recommendation, low-quality evidence]

Commentary

The efficacy of retreatment with the same anti-TNFα agent for the relapse after the discontinuation is generally favorable (> 80%) [214–218]. However, careful interpretation is required because these retrospective observational studies might have different efficacy criteria and most reports focused more on IFX than on other anti-TNFα agents.

There may be a concern that anti-drug antibodies could be increased due to the interval of drug suspension. However, drug suspension during remission is generally considered safe because fewer anti-drug antibodies are produced during drug suspension than with intermittent administration aimed at inducing remission [214]. On the other hand, it has also been confirmed that the presence of anti-drug antibodies is a risk factor for an infusion reaction during re-administration [216]. There is no evidence for the efficacy of switching to another anti-TNFα agent with less immunogenicity upon relapse.

CQ 8. Is concomitant use of immunomodulators and anti-TNF-α agents useful in the treatment of patients with IBD?

Recommendations

In the treatment of CD, combination therapy is recommended because it is more effective than monotherapy,
In the treatment of UC, combination therapy is suggested, because it may be more effective compared to monotherapy, which each drug is used separately. [Strong recommendation, high-quality evidence]

In the SONIC study, patients with moderate to severe CD who had not previously been treated with biologics or immunomodulatory drugs were assigned to three groups, namely, a IFX monotherapy group, an immunomodulator (IM) monotherapy group, and a group treated with the combination of both therapies. The findings showed that the combined therapy group was superior to the other two groups in terms of remission induction and endoscopic healing rates [200]. In the DIAMOND study performed in Japan, the efficacy of a combination therapy using adalimumab (ADA) with IMs in comparison with ADA monotherapy was examined, but the remission rates were not significantly different between two groups [203]. However, a subanalysis report showed that the endoscopic efficacy tended to be greater when combination therapy was used [219]. Based on these results, meta-analyses and major guidelines published in Western countries strongly recommend the concomitant use of IMs when using anti-TNF-α antibody agents in CD patients [75, 159, 220]. The additional efficacy of combination therapy is based on increasing concentrations of anti-TNF-α antibody products as a result of the suppression of the production of anti-drug antibodies [75, 220]; continuing the combination therapy for the first 6 months is considered particularly important [159].

In UC patients, the only one major research study, the UC SUCCESS trial, examined combination therapies using IFX with IMs [221]. In this trial, patients with moderate to severe UC who had not previously been treated with biologics or IMs were assigned to the following three groups: an IFX monotherapy group, an IM monotherapy group, and a group treated with a combination of both therapies. The findings showed that the combination therapy group was superior to the other two groups in terms of rates of steroid-free induction of remission and endoscopic healing. ADA and golimumab, which are less immunogenic than IFX, have not yet been shown to increase the effect of the combination therapy. For this reason, the efficacy of combination therapy is limited to IFX, and even guidelines in Western countries do not recommend it as strongly as in CD [222, 223].

Given that combination therapy can also be associated with risks, there needs to be a comprehensive assessment that includes an estimation of each individual patient’s clinical background, such as age, comorbidity, and medical history, as well as an assessment of the need for combination therapy depending on the course of treatment. Decisions for the administration of combination therapy and its duration should be based on the abovementioned assessment findings [224].

CQ 9. Do anti-TNF-α agents prevent the recurrence of CD after surgery?

Recommendation

- It is recommended that anti-TNF-α agents be administered to prevent endoscopic recurrence. [Strong recommendation, moderate-quality evidence]

Commentary

We performed a systematic review and meta-analysis of a total of 570 patients, including 254 in an intervention group and 316 in a control group [225]. The results of eight randomized control studies performed to determine efficacy of anti-TNF-α agents administered after surgery for the prevention of endoscopic recurrence [relative risk (RR) 0.34, 95% CI 0.22–0.53] showed no increase in AEs (RR 1.75, 95% CI 0.81–3.79). However, clinical recurrence was not prevented (RR 0.60, 95% CI 0.36–1.02) [225].

The meta-analysis included analyses conducted during 1–2 years of treatment with anti-TNF-α agents after surgery, without the consideration of concomitant treatment and with different definitions for outcomes. Moreover, the participants included patients who had been treated with an anti-TNF-α agent and those naïve to anti-TNF agent before surgery. The findings of efficacy may differ based on the outcome of interest, such as long-term prevention, the avoidance of further surgery, cost-effectiveness, and safety. Our results provide evidence for the efficacy as well as the safety of anti-TNF-α agents, which should be confirmed in a future nationwide observational or prospective study.

CQ 10. Is the long-term combination of anti-TNF-α agents and immunomodulators safe?

Recommendation

- It is recommended to evaluate the long-term combination of anti-TNF-α agents and immunomodulators from the viewpoint of usefulness and safety, considering the patient background, treatment course, and risk differences between Japan and Western countries. [Strong recommendation, moderate-quality evidence]
Commentary

The combination therapy of anti-TNF\(\alpha\) agents and immunomodulators (IMs) has been reported to increase the risk of opportunistic infections such as herpes zoster, lymphoma (non-Hodgkin) and skin cancer (nonmelanoma) in Western countries \[159, 169, 226–234\]. However, studies in Japan did not confirm an increase in the incidence of lymphoma associated with the combination therapies \[172, 186\]. Regarding skin cancer, a 3.39 to 4.03-fold increase in risk was reported in Japanese patients treated with IMs, and the actual prevalence was only 2.94 to 4.94 per 100,000 per year, which is very low compared with that in the Western population.

Long-term combination therapies in young to adolescent male patients have been reported to increase the risk of high-mortality hepatosplenic T-cell lymphoma in the Western countries, but not in Japan \[210, 235\]. Regarding the contribution of IMs to Epstein-Barr (EB) virus infection, it has been reported that the incidence of lymphoma increases in preinfected patients and the risk of hemophagocytic syndrome and lymphoma increases in uninfected patients when they are first infected. Therefore, it is recommended to test for EB virus serum antibodies prior to administration and to avoid the use of IMs in uninfected individuals. Screening for cervical cancer is recommended for female patients receiving long-term AZA or 6-MP therapy \[210\].

At present, there is no previous report that provides a clear answer as to how long the combination anti-TNF\(\alpha\) agents and IMs will increase risks. Despite the benefit of IMs such as the suppression of antibody production for anti-TNF\(\alpha\) agents, the indication is determined by considering the risks and benefits for each patient.

FRQ 4. Can anti-TNF-\(\alpha\) agents be stopped?

Statement

- The discontinuation of anti-TNF\(\alpha\) agents should be carefully discussed, considering the risk of relapse, safety, medical costs, and patient desire, because discontinuation is likely to increase the risk of relapse.

Commentary

The possibilities of discontinuating anti-TNF\(\alpha\) agents in IBD patients maintaining long-term remission with the maintenance treatment are increasingly discussed. However, it is necessary to take not only the advantages but also the disadvantages of long-term treatment, such as safety, medical costs, and treatment fatigue, into consideration when discussing the discontinuation.
FRQ 6. Is an anti-TNF-α agent useful for CD with GI bleeding?

Statement
• An anti-TNF-α agent is one option for CD with GI bleeding.

Commentary
Massive bleeding from the GI tract is rarely seen in CD. First, we start conservative treatment with fasting and fluid replacement, and then the intestinal tract is rested. There have been reports that steroids and IFX were effective as drug treatments [242, 243]. However, no reports have summarized a large number of cases. It has also been reported that IMs reduce the risk of bleeding from lower GI lesions [244]. Endoscopic hemostasis should be attempted if possible. There are reports that vasopressin injection and arterial embolization have been shown to be useful in angiography [245, 246], but intestinal necrosis due to intestinal ischemia has been a problem with arterial embolization. Surgical treatment is required when hemostasis is difficult to achieve with medical treatment [247]. The surgical rate for initial massive bleeding has been reported to be 20–90%, and the surgical rate for rebleeding after conservative treatment has been reported to be 30–35% [248, 249].

Ustekinumab

BQ 29. Is ustekinumab useful for treating CD?

Statement
• Ustekinumab is useful as an induction/maintenance therapy for moderate to severe CD.

These statements were made with reference to [250–255].

FRQ 7. Is the concomitant use of an immunomodulator with ustekinumab more useful than ustekinumab monotherapy as an induction therapy for CD?

Statement
• Based on the current evidence, we cannot state that the concomitant use of an immunomodulator with ustekinumab is more useful than ustekinumab monotherapy as an induction therapy for CD.

Commentary
At the time of the survey, there have been no RCTs directly comparing the effectiveness of ustekinumab (UST) monotherapy and UST plus an immunomodulator (IM) as an induction therapy for CD. Six studies that analyzed induction therapy with UST for CD patients were extracted [250, 252, 256–259], and a meta-analysis was performed. The concomitant use of IM was significantly more effective than UST monotherapy (OR: 1.35, 95% CI: 1.06–1.71) [260]. However, the risk of bias was considered to be high and the level of evidence was judged to be “weak”. There is currently no clear evidence of AEs with the addition of an IM to UST [261], however, the risks of infection and malignancy should be considered as with the risks of using IMs in combination with other biological agents. Collectively, the current evidence in the published papers does not indicate that the concomitant use of an IM with UST is more useful than UST monotherapy as an induction therapy for CD, and accumulating evidence is necessary in the future.

FRQ 8. Is ustekinumab useful for preventing postoperative recurrence in CD?

Statement
• There have been no reports investigating the prevention of postoperative recurrence of CD by ustekinumab.

Commentary
There have been many reports on the usefulness of anti-TNF-α agents for the prevention of postoperative recurrence in CD [262, 263]. UST was approved for CD worldwide from 2016 to 2017, and there has been no report investigating its effect for the prevention of postoperative recurrence at the time of the survey (April 2020). Data will be collected in the future regarding the selection of drugs suitable for preventing postoperative recurrence.

FRQ 9. Is ustekinumab useful for perianal lesions of CD?

Statement
• Ustekinumab may be useful for perianal lesions in CD, and further evidence needs to be accumulated.
Commentary

As of April 2020, there have been no prospective studies investigating the efficacy of UST for perianal lesions in CD. According to the summarized subanalysis limited to patients with active penetrating lesions in placebo-controlled RCTs, 39 of 150 patients (26%) showed improvement in penetrating lesions 8 weeks after UST treatment, and the improvement rate was higher than that for placebo at 8, 22, and 44 weeks [264]. In addition, a meta-analysis of these subanalyses showed a risk ratio of UST to improvement of penetrating lesions to placebo of 1.77 (95% CI: 0.93–3.37), suggesting a nonsignificant but improving effect of UST [238]. The real-world data after approval are almost in line with the results of clinical trials [256, 258, 265].

Based on these results, UST may be useful for perianal lesions. However, the subanalysis of the clinical trials includes all penetrating lesions other than perianal disease, and there remain many unclear points, such as in what kind of anal lesions is UST effective and how many bionaive cases are included. Detailed analysis focusing on perianal lesions is required in the future.

FRQ 10. Is ustekinumab safe for pregnant women with CD?

Statement

- The safety of ustekinumab to pregnant women with CD has not been established.

Commentary

In approximately 10 cases of CD reported so far, serious problems were not reported in either mothers or infants, except for one case of miscarriage at 8 weeks gestation [266]. UST, a human monoclonal IgG1 antibody, can transfer to the placenta in late pregnancy and to breast milk after parturition [267]. In two CD patients who used UST until late pregnancy, cord-blood drug levels were higher than maternal serum levels, but there was no problem with the condition of the children in either case [268, 269]. In addition, the transfer of UST to breast milk has been reported [269], but there has been no report that there is a major problem in the growth process of the infants. However, the safety of UST for pregnant women has not been established due to the limited data, and the possibility of continuing administration should be carefully examined in each individual case.

Vedolizumab

BQ 30. Is vedolizumab effective for UC?

Statement

- Vedolizumab is effective in inducing and maintaining remission in moderate to severe UC.

This statement was made with reference to [111, 270–280].

BQ 31. Is vedolizumab effective for CD?

Statement

- Vedolizumab is effective in inducing and maintaining remission in moderate to severe CD.

This statement was made with reference to [275, 276, 281–287].

BQ 32. Is vedolizumab effective for IBD patients refractory to anti-TNFα agents?

Statements

- Vedolizumab is useful for both UC and CD patients refractory to anti-TNFα agents.
- Vedolizumab is particularly useful for maintaining remission in UC and CD patients refractory to anti-TNF-α agents.

These statements were made with reference to [273, 274, 277, 283, 285, 288–293].

BQ 33. What should we consider in the safety of vedolizumab?

Statements

- We need to pay attention to respiratory tract infections (especially upper respiratory tract infections) and enteric infections (e.g., C. difficile) during vedolizumab treatment.
- No significant association of vedolizumab with the development of progressive multifocal leukoencephalopathy (PML) or malignancy has been reported to date.
- The safety of vedolizumab for pregnant women, lactating women, women attempting to conceive, and children has not been sufficiently established.

These statements were made with reference to [276, 294–304].
FRQ 11. Are anti-TNFα agents effective for patients refractory to vedolizumab?

Statement
- Evidence regarding the efficacy of anti-TNFα agents in IBD patients in whom vedolizumab failed has not been accumulated, and this issue remains to be investigated.

Commentary
A post-hoc analysis of the clinical trials of vedolizumab (VDZ) demonstrated that VDZ was less effective in patients in whom anti-TNFα antibody agents failed than in those who had not received anti-TNFα agents in both UC and CD [273, 282]. On the other hand, we did not find any studies investigating the efficacy of anti-TNFα agents in patients in whom VDZ had failed.

FRQ 12. Should vedolizumab be used with immunomodulators in IBD patients?

Statement
- The currently available evidence does not suggest a benefit for the concomitant use of immunomodulators with vedolizumab, but further studies are warranted.

Commentary
There have been no RCT to examine the efficacy of the concomitant use of VDZ with IMs. A subgroup analysis of the GEMINI 1 study, a phase III study of VDZ in UC patients, reported that the rates of clinical remission and mucosal healing at weeks 6 and 52 were higher in patients receiving VDZ than in those receiving placebo regardless of use of IMs at baseline [305]. Most observational studies have reported that concomitant use of IMs did not affect the effectiveness of VDZ in patients with UC or CD [277, 293], while a small observational study reported that concomitant use of IMs was a predictor of clinical remission and clinical response at week 54 in patients with CD (odds ratio 8.33, 95% CI 2.15 -32.26) [306]. It has been reported that concomitant use of IMs does not affect serum trough concentrations of VDZ or the development of anti-VDZ antibodies [294, 307]. In terms of safety, an integrated analysis of the clinical trials of VDZ reported that use of IMs at baseline was not associated with serious infections [294]. On the other hand, a retrospective study demonstrated that concomitant use of immunosuppressive therapy (IMs or steroids) was associated with infections (odds ratio 1.72, 95% CI 1.20 -2.46) [308].

Tofacitinib

BQ 34. Is tofacitinib useful for refractory patients with moderate to severely active UC?

Statement
- Tofacitinib is useful for refractory patients with moderate to severely active UC.

This statement was made with reference to [309, 310].

BQ 35. What are the precautions to take when using tofacitinib in UC treatment?

Statements
- Tofacitinib therapy should be used with caution for the risk of infection complications, especially herpes zoster infections.
- Elderly patients with cardiovascular disease and rheumatoid arthritis treated with tofacitinib (predominantly 10 mg twice daily) have an increased risk of pulmonary embolism.

These statements were made with reference to [311–315].

CQ 11: Is tofacitinib effective for UC patients refractory to anti-TNFα agents?

Recommendation
- It is recommended to use tofacitinib for UC patients refractory to anti-TNFα agents [Weak recommendation, moderate-quality evidence]

Commentary
Tofacitinib 10 mg twice daily was shown to be more effective as induction therapy in a phase III, randomized, double-blind, placebo-controlled trial; remission at 8 weeks occurred in 12.6% of the patients in the tofacitinib group versus 1.5% in the placebo group (P < 0.01) in the OCTAVE 1 trial and in 12.0% versus 0.0% (P < 0.01) in the OCTAVE 2 trial [309]. In the maintenance OCTAVE Sustain trial, week 52 remission rates were 36.6% in the tofacitinib group versus 12.0% in the placebo group(p < 0.0001). A network meta-analysis comparing tofacitinib with other drugs also demonstrated its usefulness in patients who had failed to respond to anti-TNFα therapy [316].
Endoscopic treatment for CD

CQ 12. Can endoscopic balloon dilation (EBD) for intestinal stenosis in CD avoid surgical intervention?

Recommendation

• EBD is recommended for indicated intestinal stenosis in CD because it can avoid surgical intervention at least in the short-term. [Strong recommendation, moderate-quality evidence]

Commentary

Indications for EBD for intestinal stenosis in CD include (a) intestinal stricture length of 5 cm or less, (b) no fistula or abscess in stricture site, (c) no deep ulcer in stricture site, and (d) no severe curvature and strong adhesion in strictured part [317]. It is important to take into carefully consider the abovementioned indications when performing EBD in CD patients. In several meta-analyses with regard to EBD for intestinal stenosis in CD focusing on lesions of the lower GI tract (small intestine, ileocolonic anastomosis, large intestine), the short-term technical success rate was 86–94%, and the short-term clinical symptom improvement rate was 58–87% [318–321]. For upper GI (gastro-duodenal) lesions, a meta-analysis and a prospective observational study reported a short-term technical success rate of 93–100% and a short-term clinical symptom improvement rate of 87% [322, 323]. Since complications associated with EBD for intestinal stenosis in CD are reported in 2 to 6% and perforation is also observed in 1 to 3%, strict intraoperative and postoperative monitoring and management are needed [318–323].

Surgical treatment and colitic cancer

BQ 36. What are the indications and precautions for the surgical treatment of IBD?

Statements

• In severe cases of IBD and those with cancer or dysplasia, surgical treatment is expected to improve life expectancy. In addition, the quality of life can be improved in patients with symptoms caused by the primary disease that do not improve with medical treatment, side effects of medication, and extraintestinal complications (especially pyoderma gangrenosum).
• Postoperative complications such as suture failure and intestinal obstruction, ileocolitis in UC, and short bowel syndrome in CD can occur.
• Complications such as significant bleeding and toxic megacolon are likely to occur in elderly patients with IBD due to the delay in surgery.

These statements were made with reference to [16, 26, 111, 324–338].

CQ 13: Who should receive surveillance colonoscopy for detecting CRC in UC?

Recommendation

• Surveillance colonoscopy is useful and recommended for patients with total and left-sided UC starting at 8 years after the onset of UC. [Strong recommendation, low-quality evidence]

Commentary

Long-standing UC is a well-known risk factor for the development of colorectal cancer (CRC). A Cochrane Review integrating four observational studies showed that surveillance colonoscopy is associated with a reduction in CRC development and death [339]. This review included both CD and UC cases, and the CRC death rates were 7.7% and 22.3% in the surveillance and the nonsurveillance group, respectively. A multicenter retrospective study from Japan investigating surgically resected cases also demonstrated a survival benefit of the surveillance colonoscopy [33]. Western guidelines recommend that patients with total and left-sided colitis receive surveillance colonoscopy starting at eight years after the onset of UC [223, 340]. However, approximately 20% of CRC developed within eight years of UC duration [33, 341]. Especially in late-onset UC cases (> 40 or > 50 years old), the concomitant CRC is high even within 8 years of UC duration [33, 342]. Patients with primary sclerosing cholangitis (PSC) should receive surveillance colonoscopy from the onset of the disease due to its ambiguous disease onset and high CRC risk [223, 340]. The optimal interval of surveillance colonoscopy is not established. Several studies have showed that those who received surveillance within two or three years had a better survival [33, 343]. Although European Crohn’s and Colitis Organisation (ECCO) guidelines set a different interval from 1 to 5 years according to the risk stratification, its evidence level is not high [340]. Analyses from surgically resected cases showed that 12% of CRC cases were already staged III or IV at the time of surgery among patients who received surveillance within two years [33]. A determination of the
optimal surveillance interval incorporating both the CRC risk and progression speed is warranted.

CQ 14. What kind of biopsy method is recommended for UC-associated CRC surveillance?

Recommendation

- Targeted biopsy is recommended for UC-associated CRC surveillance. [Strong recommendation, moderate-quality evidence]

Commentary

An RCT demonstrated that targeted biopsy is comparable to random biopsy in terms of the neoplasia detection rate in UC-associated cancer surveillance [344]. Therefore, a targeted biopsy is recommended for patients with quiescent disease as enrolled in the RCT. On the other hand, there are cases with invisible dysplastic lesions that only random biopsy could detect [33, 345]. An observational study investigating 1000 consecutive cases of UC and CD showed that random biopsy increased the dysplasia detection rate by 15% [345]. The study mentioned that patients with the lead pipe appearance, PSC, and past dysplasia merit random biopsy. Random biopsy takes four samples every 10 cm, however, it is time-consuming and not realistic in clinical settings. Analyses from surgically resected specimens clarified the rectum and the sigmoid colon to be hotspots of UC-associated CRC in Japan [33]. Therefore, it is vital to carefully observe these hotspot regions and take biopsies from areas with subtle changes such as reddish (or pale) lesions and surface pattern changes in addition to targeted biopsy from suspicious lesions.

A meta-analytic study showed the benefit of chromoendoscopy over white-light endoscopy in UC-associated CRC surveillance [346]. When analyzing RCTs alone, although chromoendoscopy was superior to white-light endoscopy in the non-high-definition (HD) endoscopy setting, it was not evident in the HD endoscopy setting. Different RCTs adopted different concentrations of methylene blue or indigo carmine. In terms of narrow-band imaging (NBI), although a meta-analytic study denied the benefit of NBI over chromoendoscopy [347], a recent study using HD endoscopy showed a comparable neoplasia detection rate between NBI and white-light endoscopy in UC [348]. In real-world settings, it is vital to combine available modalities at each institution to increase the neoplasia detection rate.

FRQ 13. How is cancer surveillance performed for CD?

Statement

- Cancer surveillance is recommended for CD patients. However, proper methods remain unclear.

Commentary

European guidelines recommend surveillance colonoscopy for patients with CD, as the incidence of CRC is significantly higher in those than in the general population [349–352]. We performed a systematic review, and those results also confirmed that the standardized incidence ratios of colorectal and small-bowel cancer were significantly high in association with CD, although the incidence of anorectal lesion cancers was found to be significantly higher than that of either colonic or small bowel lesion cancers in Asian countries [353]. Therefore, we were not able to draw a conclusion regarding the usefulness of or provide a recommendation for surveillance colonoscopy for CD. Although efficacy has not been proven, an annual examination under anesthesia, as well as MRI and examinations by an expert IBD surgeon, are recommended for patients in Japan in the Japanese guidelines for intractable disease affiliated with the Japan Ministry of Health, Labor and Welfare.

FRQ 14. Is endoscopic treatment recommended for colitis-associated cancer/dysplasia in patients with IBD?

Statement

- We cannot make a recommendation regarding endoscopic treatment for colitis-associated cancer/dysplasia.

Commentary

Endoscopic treatment is recommended only for patients with a polypoid dysplastic lesion with a clearly visible boundary line, while close repeated follow-up surveillance colonoscopy examinations may also be conditionally needed [354]. However, long-term safety factors including recurrence, mortality, and morbidity related to endoscopic treatment have not been sufficiently investigated. In addition, the characteristics of metachronous or synchronous colitis-associated cancer have not been well elucidated. At present, we cannot recommend endoscopic treatment for colitis-associated cancer/dysplasia.
FRQ 15. Are anti-TNF-α agents efficacious for pouchitis?

Statement

• Evidence for the efficacy of anti-TNF-α agents for pouchitis is limited.

Commentary

The first-line treatment for pouchitis includes ciprofloxacin and metronidazole. Second-line treatment for those with antibiotic-resistant or dependent pouchitis has not been established. In real-world settings, drugs effective for UC are empirically used for pouchitis. One RCT investigated the efficacy of anti-TNF-α agents, comparing ADA with placebo [355]. This RCT failed to accrue 24 initially planned patients with pouchitis resistant to >4 weeks of antibiotics, and enrolled only 13 patients. Thus, the study was underpowered to show the efficacy of ADA for antibiotic-resistant pouchitis. A meta-analytic study [356] integrated retrospective observational studies using anti-TNF-α agents, mainly one-arm studies consisting of a few cases. In this meta-analysis, the short-term (at ~8 weeks) clinical response rates of IFX and ADA were 56% (95% CI 36–75%) and 38% (95% CI 8–72%), respectively, while the long-term (at ~12 months) response rates of IFX and ADA were 59% (95% CI 45–72%) and 30% (95% CI 15–46%), respectively. This study also showed that those with CD of the pouch had a higher response rate to anti-TNF-α agents. Although anti-TNF-alpha agents seem useful for antibiotic-dependent or resistant-pouchitis based on the results of several observational studies, further studies with controls are warranted.

Special situations

BQ 37. How do you deal with elderly IBD patients?

Statements

• The treatment of elderly patients with IBD is essentially the same as that of patients of average age. Nevertheless, it is essential to determine the appropriate surgery timing in severe cases, bearing in mind that delays in diagnosis and surgery can have a prognostic impact.
• It is essential to consult a specialist as soon as possible in refractory cases resistant to immunosuppressive therapy.

These statements and supplementary information were made with reference to [16, 336, 337, 357–359].

Supplementary information

There is no absolute definition of an elderly UC patient, but for convenience, an elderly UC patients is often defined as those “over 60” or “over 65”. The surgery rate remains the same in elderly UC patients as in nonelderly UC patients. However, it has been reported that the surgery rate of elderly-onset UC is higher than that of non-elderly-onset UC [359]. Therefore, it is necessary to clearly distinguish between elderly-onset UC with a short disease duration and young-onset UC with a long disease duration (aged UC).

BQ 38. How do you deal with patients with IBD during pregnancy and lactation?

Statements

• Patients and their physicians should discuss and select a treatment for patients with IBD during pregnancy and lactation, taking into account each case’s treatment benefits and harms.
• In many cases, the benefits of treatment outweigh medication harm; therefore, treatment should be continued during pregnancy.

These statements and supplementary information were made with reference to [360–376].

Supplementary information

Although AZA, CsA, and Tac are contraindicated for administration to pregnant women in the Japanese package insert, no clinically significant teratogenicity or fetal toxicity has been demonstrated. Therefore, the above three drugs, with colchicine, can be administered with informed consent, even during pregnancy, under certain circumstances.

In clinical practice, anti-TNFα agents are administered to patients with moderate to severe IBD, and the continuation of administration is often necessary. When anti-TNFα agents are used beyond 22 weeks of gestation, live vaccines such as the Bacillus Calmette-Guérin vaccine (BCG) (usually administered at 5 to 7 months) should be avoided before the child reaches 6 months of age (until the disappearance of administered antibodies) [360].

BQ 39. What are the extraintestinal complications observed in patients with IBD?

Statements

• Extraintestinal complications associated with IBD are mainly skin lesions and arthritis.
Erythema nodosum and pyoderma gangrenosum are two of the most common skin lesions associated with IBD. Characteristically, they are often painful and can be relieved by controlling the inflammation of the intestinal tract.

Arthritis associated with IBD includes ankylosing spondylitis and peripheral arthritis, both of which are negative for rheumatoid factors.

These statements were made with reference to [16, 377–382].

CQ 15. Is thromboembolism prophylaxis necessary for hospitalized IBD patients?

Recommendation

- We propose that thromboprophylaxis in hospitalized IBD patients should be considered with an understanding of the increased risk of bleeding associated with the intervention. [Weak recommendation, low-quality evidence]

Commentary

The risk of venous thrombosis in patients with UC and CD is reported to be approximately twice that of the non-IBD individuals [383] and is particularly higher during flares, the chronic active phase, and hospitalization periods [384]. Comorbidities and a history of steroid use are also associated with an increased risk of thrombosis [385]. Anticoagulant thromboprophylaxis is recommended for hospitalized IBD patients without severe GI bleeding, especially in moderate to severe cases [386]. Mechanical prophylaxis (e.g., intermittent pneumatic compression) is also recommended in IBD cases with severe bleeding [386]. However, further epidemiological studies are needed to determine the contribution of thromboprophylaxis to physical prognosis and social resources in hospitalized IBD patients. The implementation of thromboprophylaxis in hospitalized IBD patients should be determined considering other risk factors (obesity, steroid use, abdominal surgery, etc.) and the increased risk of bleeding from the GI tract and other organs associated with the intervention [387].

Postscript

Trends in the diagnosis and treatment of IBD are constantly evolving. With the accumulation of new evidence and the approval of new therapeutic agents, the treatment system for IBD has changed dramatically. In the future, diagnosis using artificial intelligence will be applied to daily clinical practice. Therefore, by the time the next guideline is published, it may be necessary to supplement the guideline with an annual Review and other documents that provide a high level of evidence and new treatments that should be known in clinical practice.

The preparation of these guidelines required a great deal of time and effort, from the process of narrowing down the literature to the preparation of statements and commentaries. This was a truly arduous task. I would like to take this opportunity to express my sincere gratitude to the members of the creation committee and the evaluation committee.

Acknowledgements

This article was supported by a Grant-in-Aid from JSGE. The authors thank the investigators and supporters for participating in the studies. The authors express special appreciation to Mr. Yuji Tatsugami and Miss. Ayari Sada (Nankodo) for their help in creating these guidelines.

Funding

This work was supported by Health and Labour Sciences Research Grants for research on intractable diseases from the Ministry of Health, Labour and Welfare of Japan (Investigation and Research for intractable Inflammatory Bowel Disease).

Compliance with ethical standards

Conflict of interest

Any financial relationship with enterprises, businesses or academic institutions in the subject matter or materials discussed in the manuscript are listed as follows; (1) those from which the authors, the spouse, partner or immediate relatives of authors, have received individually any income, honoraria or any other types of remuneration; Abbvie, Kissei Pharmaceutical, Kyorin Pharmaceutical, Mitsubishi Tanabe Pharma, Janssen Pharmaceutical, Takeda Pharmaceutical, Pfizer, Celgene, EA Pharma, Zeria Pharmaceutical, Moehida Pharmaceutical, Nippon Kayaku, Daiichi Sankyo, JIMRO, Alphresa Pharma, Gilead Sciences, and (2) those from which the authors have received scholarship/research grant; Nippon Kayaku, Takeda Pharmaceutical, Otsuka Pharmaceutical, Eisai, EA Pharma, Hoya Group Pentax Medical, Abbvie, Mitsubishi Tanabe Pharma, Moehida Pharmaceutical, Janssen Pharmaceutical, EA Pharma, JIMRO, Zeria Pharmaceutical, Pfizer, Alphresa Pharma, Boehringer Ingelheim, Celgene, Eli Lilly and Company, Gilead Sciences, Kissei Pharmaceutical, and (3) those from which the authors have received individually endowed chair; Mitsubishi Tanabe Pharma, EA Pharma, AbbVie, Janssen Pharmaceutical, Moehida Pharmaceutical, Takeda Pharmaceutical, Kyorin Holdings, JIMRO, Zeria Pharmaceutical, Otsuka Pharmaceutical.

Appendix

The members of the Guidelines Committee who created and evaluated the JSGE “Evidence-based clinical practice guidelines for inflammatory bowel disease” are listed below.
37. Horsthuis K, Bipat S, Bennink RJ, et al. Inflammatory bowel disease diagnosed with US, MR, scintigraphy, and CT: meta-analysis of prospective studies. Radiology. 2008;247:64–79.

38. Hollerbach S, Geissler A, Schieg H, et al. The accuracy of abdominal ultrasound in the assessment of bowel disorders. Scand J Gastroenterol. 1998;33:1201–8.

39. Mary JY, Modigliani R. Development and validation of an endoscopic index of the severity for Crohn’s disease: a prospective multicentre study. Groupe d’Etudes Therapeutiques des Affections Inflammatoires du Tube Digestif (GETAID). Gut. 1989;30:983–9.

40. Daperno M, D’Haens G, Van Assche G, et al. Development and validation of a new, simplified endoscopic activity score for Crohn’s disease: the SES-CD. Gastrointest Endosc. 2004;60:505–12.

41. Denis M-A, Reenaers C, Fontaine F, et al. Assessment of endoscopic activity index and biological inflammatory markers in clinically active Crohn’s disease with normal C-reactive protein serum level. Inflamm Bowel Dis. 2007;13:1100–5.

42. Pera A, Bellando P, Caldera D, et al. Colonscopy in inflammatory bowel disease. Diagnostic accuracy and proposal of an endoscopic score. Gastroenterology. 1987;92:181–5.

43. Witte AM, Veenendaal RA, Van Hogezaand RA, et al. Crohn’s disease of the upper gastrointestinal tract: the value of endoscopic examination. Scand J Gastroenterol Suppl. 1998;225:100–5.

44. Schmitz-Moormann P, Schäg M. Histology of the intestinal tract in Crohn’s disease of children and adolescents: multicentric paediatric Crohn’s disease study. J Clin Pathol. 2007;60:1268–78.

45. Dionisio PM, Gurudu SR, Leighton JA, et al. Capsule endoscopy has a significantly higher diagnostic yield in patients with suspected and established small-bowel Crohn’s disease: a meta-analysis. Am J Gastroenterol. 2010;105:1240–8 (quiz 1249).

46. Bourreille A, Ignatovic A, Aabakken L, et al. Role of small-bowel endoscopy in the management of patients with inflammatory bowel disease: an international OMEDECCO consensus. Endoscopy. 2009;41:618–37.

47. Ozaki R, Kobayashi T, Okuyama T, et al. Capsule endoscopy findings for the diagnosis of Crohn’s disease: a nationwide case-control study. J Gastroenterol. 2019;54:249–60.

48. Choi M, Lim S, Choi M-G, et al. Effectiveness of capsule endoscopy compared with other diagnostic modalities in patients with small bowel crohn’s disease: a meta-analysis. Gut Liver. 2017;11:62–72.

49. González-Suárez B, Rodriguez S, Ricart E, et al. Comparison of capsule endoscopy and magnetic resonance enterography for the assessment of small bowel lesions in Crohn’s disease. Inflamm Bowel Dis. 2018;24:775–80.

50. Göldner SK, Schreyer AG, Endlicher E, et al. Comparison of capsule endoscopy and magnetic resonance (MR) enteroclysis in suspected small bowel disease. Int J Colorectal Dis. 2006;21:97–104.

51. Yung DE, Har-Nevo O, Tham YS, et al. Capsule endoscopy, magnetic resonance enterography, and small bowel ultrasound for evaluation of postoperative recurrence in Crohn’s disease: systematic review and meta-analysis. Inflamm Bowel Dis. 2017;23:93–100.
55. Levesque BG, Cipriano LE, Chang SL, et al. Cost effectiveness of alternative imaging strategies for the diagnosis of small-bowel Crohn’s disease. Clin Gastroenterol Hepatol. 2010;8(261–7):267.e1-4.

56. Morita E, Murano M, Kojima Y. Role of capsule endoscopy in the management of patients with Crohn’s disease. Stomatol Intest. 2010;45:1689–95.

57. Yao T, Matsui T, Hiwatashi N. Crohn’s disease in Japan: diagnostic criteria and epidemiology. Dis Colon Rectum. 2000;43:S85-93.

58. Hisabe T, Hirai F, Matsui T, et al. Evaluation of diagnostic criteria for Crohn’s disease in Japan. J Gastroenterol. 2014;49:93-9.

59. Van Assche G, Dignass A, Panes J, et al. The second European evidence-based consensus on the diagnosis and management of Crohn’s disease: definitions and diagnosis. J Crohns Colitis. 2010;4:7–27.

60. Panés J, Bouzas R, Chaparro M, et al. Systematic review: the use of ultrasonography, computed tomography and magnetic resonance imaging for the diagnosis, assessment of activity and abdominal complications of Crohn’s disease. Aliment Pharmacol Ther. 2011;34:125–45.

61. Solem CA, Loftus EV, Fletcher JG, et al. Small-bowel imaging for diagnostic and therapeutic targets in inflammatory bowel disease. Gastrointest Endosc. 2010;68:255–66.

62. Giles E, Barclay AR, Chippington S, et al. Systematic review: MRI enterography for assessment of small bowel involvement in paediatric Crohn’s disease. Aliment Pharmacol Ther. 2013;37:1121-31.

63. Takenaka K, Ohtsuka K, Kitazume Y, et al. Correlation of the detection of inflammation and intestinal damage in Crohn’s disease. Gastroenterology. 2011;140(1817–1826):e2.

64. Ministry of Health, Labour and Welfare, Grant-in-Aid for Scientific Research on Intractable Diseases, ‘‘Research on Intractable Inflammatory Bowel Disorders’’ (Suzuki Group), 2017, Diagnostic Criteria and Treatment Guidelines for Ulcerative Colitis and Crohn’s Disease (revised version). http://www.ibdpjapan.org/. Accessed 11 Dec 2017.

65. Maaser C, Sturm A, Vavricka SR, et al. ECCO-ESGAR guideline for diagnostic assessment in IBD Part 1: Initial diagnosis, monitoring of known IBD, detection of complications. J Crohns Colitis. 2019;13:144–64.

66. Sturm A, Maaser C, Calabrrese E, et al. ECCO-ESGAR guideline for diagnostic assessment in IBD Part 2: IBD scores and general principles and technical aspects. J Crohns Colitis. 2019;13:273–84.

67. Church PC, Turner D, Feldman BM, et al. Systematic review with meta-analysis: magnetic resonance enterography signs for the detection of inflammation and intestinal damage in Crohn’s disease. Aliment Pharmacol Ther. 2015;41:153–66.

68. Rimola J, Ordás I, Rodríguez S, et al. Magnetic resonance imaging for evaluation of Crohn’s disease: validation of parameters of severity and quantitative index of activity. Inflamm Bowel Dis. 2011;17:1759–68.

69. Ordás I, Rimola J, Rodríguez S, et al. Accuracy of magnetic resonance enterography in assessing response to therapy and mucosal healing in patients with Crohn’s disease. Gastroenterology. 2014;146:374-382.e1.

70. Takenaka K, Ohtsuka K, Kitazume Y, et al. Correlation of the endoscopic and magnetic resonance scoring systems in the deep small intestine in Crohn’s disease. Inflamm Bowel Dis. 2015;21:1832–8.

71. Kitazume Y, Fujioka T, Takenaka K, et al. Crohn disease: a 5-point MR enterocolonoigraphy classification using endoscopic findings. Am J Roentgenol. 2019;212:67–76.

72. Ordás I, Rimola J, Alfaro I, et al. Development and Validation of a Simplified Magnetic Resonance Index of activity for Crohn’s disease. Gastroenterology. 2019;157(432–439):e1.

73. Koilakou S, Sailer J, Peloschek P, et al. Endoscopy and MR enteroclysis: equivalent tools in predicting clinical recurrence in patients with Crohn’s disease after ileocolic resection. Inflamm Bowel Dis. 2010;16:198–203.

74. Takenaka K, Ohtsuka K, Kitazume Y, et al. Utility of magnetic resonance enterography for small bowel endoscopic healing in patients with Crohn’s disease. Am J Gastroenterol. 2018;113:283–94.

75. Lichtenstein GR, Loftus EV, Isaacs KL, et al. ACG clinical guideline: management of Crohn’s disease in adults. Am J Gastroenterol. 2018;113:481–517.

76. Bruining DH, Zimmermann EM, Loftus EV, et al. Consensus recommendations for evaluation, interpretation, and utilization of computed tomography and magnetic resonance enterography in patients with small bowel Crohn’s disease. Gastroenterology. 2018;154:1172–94.

77. Lewis JD. The utility of biomarkers in the diagnosis and therapy of inflammatory bowel disease. Gastroenterology. 2011;140(1817–1826):e2.

78. Tibble JA, Sigthorsson G, Foster R, et al. Use of surrogate markers of inflammation and Rome criteria to distinguish organic from nonorganic intestinal disease. Gastroenterology. 2002;123:450–60.

79. Rheenen PF, van, Vijver EV de, Füller V. Faecal calprotectin for screening of patients with suspected inflammatory bowel disease: diagnostic meta-analysis. Br Med J Publ Group. 2010;341:c3369.

80. Takashima S, Kato J, Hiraoka S, et al. Evaluation of mucosal healing in ulcerative colitis by fecal calprotectin vs. fecal immunochromatographic test. Am J Gastroenterol. 2015;110:873–80.

81. Naganuma M, Kobayashi T, Nasuno M, et al. Significance of conducting 2 types of fecal tests in patients with ulcerative colitis. Clin Gastroenterol Hepatol. 2020;18(1102–1111):e5.

82. Dai C, Jiang M, Sun M-J, et al. Fecal immunochromatographic test for predicting mucosal healing in ulcerative colitis patients: a systematic review and meta-analysis. J Gastroenterol Hepatol. 2018;33:990–7.

83. Mosli MH, Zou G, Garg SK, et al. C-reactive protein, fecal calprotectin, and stool lactoferrin for detection of endoscopic activity in symptomatic inflammatory bowel disease patients: a systematic review and meta-analysis. Am J Gastroenterol. 2015;110:802–19.

84. Hiraoka S, Kato J, Nakarai A, et al. Consecutive measurements by faecal immunochromatographic test in quiescent ulcerative colitis patients can detect clinical relapse. J Crohns Colitis. 2016;10:687–94.

85. Laharie D, Filippi J, Robin X, et al. Impact of mucosal healing on long-term outcomes in ulcerative colitis treated with infliximab: a multicenter experience. Aliment Pharmacol Ther. 2013;37:998–1004.

86. Bryant RV, Winer S, Travis SPL, et al. Systematic review: histological remission in inflammatory bowel disease. Is “complete” remission the new treatment paradigm? An IOIBD initiative. J Crohns Colitis. 2014;8:1582–97.

87. Peyrin-Biroulet L, Sandborn W, Sands BE, et al. Selecting therapeutic targets in inflammatory bowel disease (STRIDE): determining therapeutic goals for treat-to-target. Am J Gastroenterol. 2015;110:1324–38.

88. Shah SC, Colombel J-F, Sands BE, et al. Systematic review with meta-analysis: mucosal healing is associated with improved long-term outcomes in Crohn’s disease. Aliment Pharmacol Ther. 2016;43:317–33.
89. Shah SC, Colombel J-F, Sands BE, et al. Mucosal healing is associated with improved long-term outcomes of patients with ulcerative colitis: a systematic review and meta-analysis. Clin Gastroenterol Hepatol. 2016;14(1245–1255):e8.

90. Zenlea T, Yee EU, Rosenberg L, et al. Histology grade is independently associated with relapse risk in patients with ulcerative colitis in clinical remission: a prospective study. Am J Gastroenterol. 2016;111:685–90.

91. Ford AC, Achkar J-P, Khan KJ, et al. Efficacy of 5-aminosalicylates in ulcerative colitis: systematic review and meta-analysis. Am J Gastroenterol. 2011;106:601–16.

92. Marshall JK, Thabane M, Steinhart AH, et al. Rectal 5-aminosalicylic acid for maintenance of remission in ulcerative colitis. Cochrane Database Syst Rev. 2012;11:CD004118.

93. Feagan BG, Macdonald JK. Oral 5-aminosalicylic acid for induction of remission in ulcerative colitis. Cochrane Database Syst Rev. 2012;10:CD00543.

94. Wang Y, Parker CE, Feagan BG, et al. Oral 5-aminosalicylic acid for maintenance of remission in ulcerative colitis. Cochrane Database Syst Rev. 2016;2016:CD005544.

95. Hanauer SB, Sandborn WJ, Kornbluth A, et al. Delayed-release oral mesalamine at 4.8 g/day (800 mg tablet) for the treatment of moderately active ulcerative colitis: The ASCEND II trial. Am J Gastroenterol. 2005;100:2478–85.

96. D’Haens G, Hommes D, Engels L, et al. Once daily MMX mesalazine for the treatment of mild-to-moderate ulcerative colitis: a phase II, dose-ranging study. Aliment Pharmacol Ther. 2006;24:1087–97.

97. Lim W-C, Hanauer S. Aminosalicylates for induction of remission or response in Crohn’s disease. Cochrane Database Syst Rev. 2010;12:CD008870.

98. Hanauer SB, Strömberg U. Oral Pentasa in the treatment of active Crohn’s disease: a meta-analysis of double-blind, placebo-controlled trials. Clin Gastroenterol Hepatol. 2004;2:379–88.

99. Coward S, Kuenzg ME, Hazlewood G, et al. Comparative effectiveness of mesalazine, sulfasalazine, corticosteroids, and budesonide for the induction of remission in Crohn’s disease: a Bayesian network meta-analysis. Inflamm Bowel Dis. 2017;23:461–72.

100. Akobeng AK, Zhang D, Gordon M, et al. Oral 5-aminosalicylic acid for maintenance of medically-induced remission in Crohn’s disease. Cochrane Database Syst Rev. 2005;100:1345–53.

101. Gordon M, Naidoo K, Thomas AG, et al. Oral 5-aminosalicylic acid for maintenance of surgically-induced remission in Crohn’s disease. Cochrane Database Syst Rev. 2011;1:CD008414.

102. Ogata H, Yokoyama T, Mizushima S, et al. Comparison of efficacy of once daily multimatrix mesalazine 2.4 g/day and 4.8 g/day with other 5-aminosalicylic acid preparation in active ulcerative colitis: a randomized, double-blind study. Intest Res. 2018;16:255–66.

103. Ogata H, Aoyama N, Mizushima S, et al. Comparison of efficacy of multimatrix mesalazine 4.8 g/day once-daily with other high-dose mesalazine in active ulcerative colitis: a randomized, double-blind study. Intest Res. 2017;15:368–79.

104. Saito D, Hayashi M, Sato T, et al. Evaluation of the drug-induced lymphocyte stimulation test for diagnosing mesalazine allergy. Intest Res. 2018;16:273–81.

105. Nguyen NH, Fumery M, Dulai PS, et al. Comparative efficacy and tolerability of pharmacological agents for management of mild to moderate ulcerative colitis: a systematic review and network meta-analyses. Lancet Gastroenterol Hepatol. 2018;3:742–53.

106. Marteau P, Probert CS, Lindgren S, et al. Combined oral and enema treatment with Pentasa (mesalazine) is superior to oral therapy alone in patients with extensive mild/moderate active ulcerative colitis: a randomised, double blind, placebocontrolled study. Gut. 2005;54:960–5.

107. Ford AC, Khan KJ, Achkar J-P, et al. Efficacy of oral vs. topical, or combined oral and topical 5-aminosalicylates, in Ulcerative Colitis: systematic review and meta-analysis. Am J Gastroenterol. 2012;107:167–76 (author reply 177).

108. Safdi M, DeMicco M, Sninsky C, et al. A double-blind comparison of oral versus rectal mesalamine versus combination therapy in the treatment of distal ulcerative colitis. Am J Gastroenterol. 1997;92:1867–71.

109. Marshall JK, Thabane M, Steinhart AH, et al. Rectal 5-aminosalicylic acid for induction of remission in ulcerative colitis. Cochrane Database Syst Rev. 2010;1:CD004115.

110. Watanabe M, Nishino H, Sameshima Y, et al. Randomised clinical trial: evaluation of the efficacy of mesalazine (mesalamine) suppositories in patients with ulcerative colitis and active rectal inflammation—a placebo-controlled study. Aliment Pharmacol Ther. 2013;38:264–73.

111. Hambor D, Eliakim R, Bettenworth D, et al. Third European evidence-based consensus on diagnosis and management of ulcerative colitis. part 2: current management. J Crohns Colitis. 2017;11:769–84.

112. Paoluzi OA, Iacopini F, Pica R, et al. Comparison of two different daily dosages (2.4 vs. 1.2 g) of oral mesalazine in maintenance of remission in ulcerative colitis patients: 1-year follow-up study. Aliment Pharmacol Ther. 2005;21:1119–6.

113. Pica R, Cassieri C, Cocco A, et al. A randomized trial comparing 4.8 vs. 2.4 g/day of oral mesalazine for maintenance of remission in ulcerative colitis. Dig Liver Dis. 2015;47:933–7.

114. Fockens P, Mulder CI, Tytgat GN, et al. Comparison of the efficacy and safety of 1.5 compared with 3.0 g oral slow-release mesalazine (Pentasa) in the maintenance treatment of ulcerative colitis. Dutch Pentasa Study Group. Eur J Gastroenterol Hepatol. 1995;7:1025–30.

115. Eaden JA, Mayberry JF. Colorectal cancer complicating ulcerative colitis: a review. Am J Gastroenterol. 2000;95:2710–9.

116. Carter MJ, Lobo AJ, Travis S, et al. Guidelines for the management of inflammatory bowel disease in adults. Gut. 2004;53(Suppl 5):V1–16.

117. Velasquez FS, Terdiman JP, Walsh JM. Effect of 5-aminosalicylate use on colorectal cancer and dysplasia risk: a systematic review and meta-analysis of observational studies. Am J Gastroenterol. 2005;100:1345–53.

118. O’Connor A, Packey CD, Akhari M, et al. Mesalamine, but not sulfasalazine, reduces the risk of colorectal neoplasia in patients with inflammatory bowel disease: an agent-specific systematic review and meta-analysis. Inflamm Bowel Dis. 2015;21:2562–9.

119. Zhao L-N, Li J-Y, Yu T, et al. 5-Aminosalicylates reduce the risk of colorectal neoplasia in patients with ulcerative colitis: an updated meta-analysis. PLoS ONE. 2014;9:e94208.

120. Nguyen GC, Gulamhusein A, Bernstein CN. 5-aminosalicylic acid is not protective against colorectal cancer in inflammatory bowel disease: a meta-analysis of non-referral populations. Am J Gastroenterol. 2012;107:1298–304.

121. Bonovas S, Fiorino G, Lytras T, et al. Systematic review with meta-analysis: use of 5-aminosalicylates and risk of colorectal neoplasia in patients with inflammatory bowel disease. Aliment Pharmacol Ther. 2017;45:1179–92.

122. Rubin DT, Huo D, Kimnucan JA, et al. Inflammation is an independent risk factor for colonic neoplasia in patients with ulcerative colitis: a case-control study. Clin Gastroenterol Hepatol. 2013;11:1601–8.e1–4.

123. Lohse H, Dejong C, Hammarqvist F, et al. ESPEN guidelines on enteral nutrition: gastroenterology. Clin Nutr Edinb Scotl. 2006;25:260–74.
124. Jowett SL, Seal CI, Phillips E, et al. Dietary beliefs of people with ulcerative colitis and their effect on relapse and nutrient intake. Clin Nutr Edinb Scotl. 2004;23:161–70.

125. Zachos M, Tondeur M, Griffiths AM. Enteral nutritional therapy for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2007;1:CD000584.

126. Okada M, Yao T, Yamamoto T, et al. Controlled trial comparing an elemental diet with prednisolone in the treatment of active Crohn’s disease. Hepatogastroenterology. 1990;37:72–80.

127. Messori A, Trallori G, D’Albasio G, et al. Defined-formula diets versus steroids in the treatment of active Crohn’s disease: a meta-analysis. Scand J Gastroenterol. 1996;31:267–72.

128. Takagi S, Utsunomiya K, Kuriyama S, et al. Effectiveness of an “half elemental diet” as maintenance therapy for Crohn’s disease: a randomized-controlled trial. Aliment Pharmacol Ther. 2006;24:1333–40.

129. Verma S, Kirkwood B, Brown S, et al. Oral nutritional supplementation is effective in the maintenance of remission in Crohn’s disease. Dig Liver Dis. 2000;32:769–74.

130. Akobeng AK, Thomas AG. Enteral nutrition for maintenance of remission in Crohn’s disease. Cochrane Database Syst Rev. 2007;3:CD005984.

131. Matsui T, Ueki M, Yamada M, et al. Indications and options of nutritional treatment for Crohn’s disease. A comparison of elemental and polymeric diets. J Gastroenterol. 1995;30(Suppl 8):95–7.

132. Hisamatsu T, Kunisaki R, Nakamura S, et al. Effect of elemental diet combined with infliximab dose escalation in patients with Crohn’s disease with loss of response to infliximab: CERISIER trial. Intest Res. 2018;16:494–8.

133. Sands BE, Sandborn WJ, Feagan B, et al. A randomized, double-blind, sham-controlled study of granulocyte/monocyteapheresis for active ulcerative colitis. Gastroenterology. 2008;135:400–9.

134. Zhu M, Xu X, Nie F, et al. The efficacy and safety of selective leukocytapheresis in the treatment of ulcerative colitis: a meta-analysis. Int J Colorectal Dis. 2011;26:999–1007.

135. Yoshino T, Nakase H, Minami N, et al. Efficacy and safety of granulocyte and monocyte adsorption apheresis for ulcerative colitis: a meta-analysis. Dig Liver Dis. 2014;46:219–26.

136. Sigurðbjörnsson FT, Bjarnason I. Leukocytapheresis for the maintenance of remission in Crohn’s disease. Cochrane Database Syst Rev. 2007;3:CD000296.

137. Sakuraba A, Motoya S, Watanabe K, et al. An open-label prospective randomized multicenter study shows very rapid remission of ulcerative colitis by intensive granulocyte and monocyte adsorptive apheresis as compared with routine weekly treatment. Am J Gastroenterol. 2009;104:2990–5.

138. Naganuma M, Yokoyama Y, Motoyama T, et al. Efficacy of apheresis as maintenance therapy for patients with ulcerative colitis in an open-label prospective multicenter randomised controlled trial. J Gastroenterol. 2020;55:390–400.

139. Fukuoka Y, Matsui T, Suzuki Y, et al. Adsorptive granulocyte and monocyte apheresis for refractory Crohn’s disease: an open multicenter prospective study. J Gastroenterol. 2004;39:1158–64.

140. Yoshimura N, Yokoyama Y, Matsuoka K, et al. An open-label prospective randomized multicenter study of intensive versus weekly granulocyte and monocyte apheresis in active crohn’s disease. BMC Gastroenterol. 2015;15:163.

141. Fukanaga K, Yokoyama Y, Kamokozuru K, et al. Adsorptive granulocyte/monocyte apheresis for the maintenance of remission in patients with ulcerative colitis: a prospective randomised, double blind, Sham-Controlled Clinical Trial. Gut Liver. 2012;6:427–33.

142. Tate D, Cairnes V, Valori R, et al. First successful use of leukocyte apheresis as maintenance therapy for Crohn’s disease in the United Kingdom. J Clin Apheresis. 2014;29:181–2.

143. Yokoyama Y, Matsuoka K, Kobayashi T, et al. A large-scale, prospective, observational study of leukocytapheresis for ulcerative colitis: treatment outcomes of 847 patients in clinical practice. J Crohns Colitis. 2014;8:981–91.

144. Sands BE, Katz S, Wolf DC, et al. A randomised, double-blind, sham-controlled study of granulocyte/monocyte apheresis for moderate to severe Crohn’s disease. Gut. 2013;62:1288–94.

145. Seow CH, Benchimol EI, Griffiths AM, et al. Traditional corticosteroids for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2008;2008:CD000792.

146. Steinhart AH, Ewe K, Griffiths AM, et al. Corticosteroids for maintenance of remission in Crohn’s disease. Cochrane Database Syst Rev. 2003;CD00301.

147. Tokuda H, Harigai M, Kameda H, et al. Consensus statements for medical practice: biological agents and lung disease. Respir Investig. 2017;55:229–51.

148. Ford AC, Bernstein CN, Khan KJ, et al. Glucocorticosteroid therapy in inflammatory bowel disease: systematic review and meta-analysis. Am J Gastroenterol. 2011;106:590–9 (quiz 600).

149. Benchimol EI, Seow CH, Steinhart AH, et al. Traditional corticosteroids for induction of remission in Crohn's disease. Cochrane Database Syst Rev. 2008;2008:CD000296.

150. Seow CH, Benchimol EI, Griffiths AM, et al. Budesonide for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2008;3:CD000296.

151. Benchimol EI, Seow CH, Otley AR, et al. Budesonide for maintenance of remission in Crohn’s disease. Cochrane Database Syst Rev. 2009;1:CD002913.

152. Seow CH, Benchimol EI, Griffiths AM, et al. Budesonide for medical practice: biological agents and lung disease. Respir Investig. 1979;77:847–69.
162. Khan KJ, Dubinsky MC, Ford AC, et al. Efficacy of immunosuppressive therapy for inflammatory bowel disease: a systematic review and meta-analysis. Am J Gastroenterol. 2011;106:630–42.

163. Timmer A, Patton PH, Chande N, et al. Azathioprine and 6-mercaptopurine for maintenance of remission in ulcerative colitis. Cochrane Database Syst Rev. 2016;2016:CD000478.

164. Prefontaine E, Sutherland LR, MacDonald JK, et al. Azathioprine and 6-mercaptopurine for maintenance of remission in Crohn’s disease. Cochrane Database Syst Rev. 2009;1:CD000067.

165. Chatu S, Subramanian V, Saxena S, et al. The role of thiopurines in reducing the need for surgical resection in Crohn’s disease: a systematic review and meta-analysis. Am J Gastroenterol. 2014;109:23–34.

166. Peyrin-Biroulet L, Deltenre P, Ardizzone S, et al. Azathioprine and 6-mercaptopurine for the prevention of postoperative recurrence in Crohn’s disease: a meta-analysis. Am J Gastroenterol. 2009;104:2089–96.

167. Chande N, Tsoulis DJ, MacDonald JK. Azathioprine or 6-mercaptopurine for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2013;4:CD000545.

168. Kandel A, Fraser AG, Korelitz BI, et al. Increased risk of lymphoma among inflammatory bowel disease patients treated with azathioprine and 6-mercaptopurine. Gut. 2005;54:1121–5.

169. Beaumerie L, Brousse N, Bouvier AM, et al. Lymphoproliferative disorders in patients receiving thiopurines for inflammatory bowel disease: a prospective observational cohort study. Lancet. 2009;374:1617–25.

170. Khan N, Abbass AM, Lichtenstein GR, et al. Risk of lymphoma in patients with ulcerative colitis treated with thiopurines: a nationwide retrospective cohort study. Gastroenterology. 2013;145(1007–1015):e3.

171. Magro F, Peyrin-Biroulet L, Sokol H, et al. Extra-intestinal malignancies in inflammatory bowel disease: results of the 3rd ECCO Pathogenesis Scientific Workshop (III). J Crohns Colitis. 2014;8:31–44.

172. Fukata N, Okazaki K, Omiya M, et al. Hematologic malignancies in the Japanese patients with inflammatory bowel disease. J Gastroenterol. 2014;49:1299–306.

173. Gisbert JP, Gomollon F. Thiopurine-induced myelotoxicity in patients with inflammatory bowel disease: a review. Am J Gastroenterol. 2008;103:1783–800.

174. Yang S-K, Hong M, Bae C, et al. A common missense variant in NUDT15 confers susceptibility to thiopurine-induced leukopenia. Nat Genet. 2014;46:1017–20.

175. Yang JJ, Landier W, Brezis M, et al. Cyclosporine A induces and maintains clinical remission in patients with severe ulcerative colitis refractory to intravenous steroids: a parallel, open-label randomised controlled trial. Lancet. 2012;380:1909–15.

176. Baumgart DC, MacDonald JK, Feagan B. Tacrolimus (FK506) for induction of remission in refractory ulcerative colitis. Cochrane Database Syst Rev. 2008;3:CD007216.

177. Ogata H, Matsu T, Nakamura M, et al. A randomised dose finding study of oral tacrolimus (FK506) therapy in refractory ulcerative colitis. Gut. 2006;55:1235–62.

178. Yamanoto S, Nakase H, Mikami S, et al. Long-term effect of tacrolimus therapy in patients with refractory ulcerative colitis. Aliment Pharmacol Ther. 2008;28:589–97.

179. Tamaki H, Nakase H, Matsuura M, et al. The effect of tacrolimus (FK-506) on Japanese patients with refractory Crohn’s disease. J Gastroenterol. 2008;43:774–9.

180. Fukuda A, Nakase H, Seno H, et al. Refractory enterovesical and duodenocolic fistulas in Crohn’s disease successfully managed with tacrolimus. J Gastroenterol. 2005;40:433–5.

181. Iida T, Nojima M, Nakase H. Therapeutic efficacy and adverse events of tacrolimus in patients with Crohn’s disease: systematic review and meta-analysis. Dig Dis Sci. 2019;64:2945–54.

182. Ford AC, Sandborn WJ, Khan KJ, et al. Efficacy of biological therapies in inflammatory bowel disease: systematic review and meta-analysis. Am J Gastroenterol. 2011;106:644–59 (quiz 660).

183. Sandborn WJ, van Assche G, Reinisch W, et al. Adalimumab induces and maintains clinical remission in patients with moderate-to-severe ulcerative colitis. Gastroenterology. 2012;142:257–65, e1–3.

184. Ma C, Huang V, Fedorak DK, et al. Outpatient ulcerative colitis primary anti-TNF responders receiving adalimumab or infliximab maintenance therapy have similar rates of secondary loss of response. J Clin Gastroenterol. 2015;49:675–82.
199. Sandborn WJ, Rutgeerts P, Eens R, et al. Adalimumab induction therapy for Crohn disease previously treated with infliximab: a randomized trial. Ann Intern Med. 2007;146:829–38.

200. Colombel JF, Sandborn WJ, Reinisch W, et al. Infliximab, azathioprine, or combination therapy for Crohn’s disease. N Engl J Med. 2010;362:1383–95.

201. Gecse KB, Bemelman W, Kamm MA, et al. A global consensus on the classification, diagnosis and multidisciplinary treatment of perianal fistulising Crohn’s disease. Gut. 2014;63:1381–92.

202. Louis E, Mary J-Y, Vernier-Massouille G, et al. Maintenance of remission among patients with Crohn’s disease on antimetabolite therapy after infliximab therapy is stopped. Gastroenterology. 2012;142:63–70.e5 (quiz e31).

203. Matsumoto T, Motoya S, Watanabe K, et al. Adalimumab monotherapy and a combination with azathioprine for Crohn’s disease: a prospective, randomized trial. J Crohns Colitis. 2016;10:1259–66.

204. Askling J, Fored CM, Brandt L, et al. Risk and case characteristics of tuberculosis in rheumatoid arthritis associated with tumor necrosis factor antagonists in Sweden. Arthritis Rheum. 2005;52:1986–92.

205. Ryu HH, Lee EY, Shin K, et al. Hepatitis B virus reactivation in rheumatoid arthritis and ankylosing spondylitis patients treated with anti-TNFα agents: a retrospective analysis of 49 cases. Clin Rheumatol. 2012;31:931–6.

206. Cleynen I, Van Moerkercke W, Billiet T, et al. Characteristics of skin lesions associated with anti-tumor necrosis factor therapy in patients with inflammatory Bowel disease: a cohort study. Ann Intern Med. 2016;164:10–22.

207. Lichtenstein GR, Feagan BG, Cohen RD, et al. Serious infections and mortality in association with therapies for Crohn’s disease: TREAT registry. Clin Gastroenterol Hepatol. 2006;4:621–30.

208. Lichtenstein GR, Feagan BG, Cohen RD, et al. Drug therapies and the risk of malignancy in Crohn’s disease: results from the TREAT Registry. Am J Gastroenterol. 2014;109:212–23.

209. Siegel CA, Marden SM, Persing SM, et al. Risk of lymphoma associated with combination anti-tumor necrosis factor and immunomodulator therapy for the treatment of Crohn’s disease: a meta-analysis. Clin Gastroenterol Hepatol. 2009;7:874–81.

210. Kotlyar DS, Osterman MT, Diamond RH, et al. A systematic review of factors that contribute to hepatosplenic T-cell lymphoma in patients with inflammatory bowel disease. Clin Gastroenterol Hepatol. 2011;9(36–41):e1.

211. Jørgensen KK, Olsen IC, Goll GL, et al. Switching from originator infliximab to biosimilar CT-P13 compared with main-

212. Ye BD, Pesegova M, Alexeeva O, 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. 2019;393:1699–707.

213. Pouillon L, Danese S, Hart A, et al. Consensus report: clinical guidelines for the prevention and management of the coeliac effect in biosimilar-treated IBD patients. Aliment Pharmacol Ther. 2019;49:1181–7.

214. Papamichael K, Vermeire S. Withdrawal of anti-tumour necrosis factor α therapy in inflammatory bowel disease. World J Gastroenterol. 2015;21:4773–8.

215. Fiorino G, Cortes PN, Ellul P, et al. Discontinuation of infliximab in patients with ulcerative colitis is associated with increased risk of relapse: a multinational retrospective cohort study. Clin Gastroenterol Hepatol. 2016;14:1426-1432.e1.

216. Baert F, Drobne D, Gils A, et al. Early trough levels and antibodies to infliximab predict safety and success of reinitiation of infliximab therapy. Clin Gastroenterol Hepatol. 2014;12:1474–81.e2 (quiz e91).

217. Casanova MJ, Chaparro M, García-Sánchez V, et al. Evolution after anti-TNF discontinuation in patients with inflammatory Bowel disease: a multicenter long-term follow-up study. Am J Gastroenterol. 2017;112:120–31.

218. Molander P, Färkkilä M, Salminen K, et al. Outcome after discontinuation of TNFα-blocking therapy in patients with inflammatory bowel disease in deep remission. Inflamm Bowel Dis. 2014;20:1021–8.

219. Watanabe K, Matsumoto T, Hisamatsu T, et al. Clinical and pharmacokinetic factors associated with adalimumab-induced mucosal healing in patients with Crohn’s disease. Clin Gastroenterol Hepatol. 2018;16:542-49.e1.

220. Jørgensen KK, Olsen IC, Goll GL, et al. Switching from originator infliximab to biosimilar CT-P13 compared with main-

221. Rubin DT, Ananthakrishnan AN, Siegel CA, et al. ACG clinical guideline: ulcerative colitis in adults. Am J Gastroenterol. 2019;114:384–413.

222. Sturm A, Mauser C, Mendall M, et al. European Crohn’s and colitis organisation topical review on IBD in the elderly. J Crohns Colitis. 2017;11:263–73.

223. Uchino M, Ikeuchi H, Hata K, et al. Does anti-tumor necrosis factor alpha prevent the recurrence of Crohn’s disease? Systematic review and meta-analysis. J Gastroenterol Hepatol. 2020. https://doi.org/10.1111/jgh.15288.

224. Sturm A, Mauser C, Mendall M, et al. European Crohn’s and colitis organisation topical review on IBD in the elderly. J Crohns Colitis. 2017;11:263–73.

225. Rubin DT, Ananthakrishnan AN, Siegel CA, et al. ACG clinical guideline: ulcerative colitis in adults. Am J Gastroenterol. 2019;114:384–413.

226. Baert F, Drobne D, Gils A, et al. Early trough levels and anti-

227. Rubin DT, Ananthakrishnan AN, Siegel CA, et al. ACG clinical guideline: ulcerative colitis in adults. Am J Gastroenterol. 2019;114:384–413.

228. Chalhoub JM, Rimmani HH, Gumaste VV, et al. Systematic review and meta-analysis: adalimumab monotherapy versus combination therapy with immunomodulators for induction and maintenance of remission and response in patients with Crohn’s disease. Inflamm Bowel Dis. 2017;23:1316–27.

229. D’Haens G, Colombel J-F, Panaccione R, et al. Five-year safety data from ENCORE, a European observational safety registry for adults with Crohn’s disease treated with infliximab [Remicade®] or conventional therapy. J Crohns Colitis. 2017;11:680–9.

230. Colombel J-F, Sandborn WJ, Reinisch W, et al. Long-term safety of adalimumab in clinical trials in adult patients with Crohn’s disease or ulcerative colitis. Aliment Pharmacol Ther. 2018;47:219–28.

231. D’Haens G, Colombel J-F, Panaccione R, et al. Lymphoma risk and overall safety profile of adalimumab in patients with Crohn’s disease with up to 6 years of follow-up in the pyramid registry. Am J Gastroenterol. 2018;113:872–82.

232. Lemaître M, Kirchgesner J, Rudnichi A, et al. Association between use of thiopurines or tumor necrosis factor antagonists
alone or in combination and risk of lymphoma in patients with inflammatory bowel disease. JAMA. 2017:318:1679–86.

233. Lichtenstein GR, Rutgeerts P, Sandborn WJ, et al. A pooled analysis of infections, malignancy, and mortality in infliximab and immunomodulator-treated adult patients with inflammatory bowel disease. Am J Gastroenterol. 2012;107:1051–63.

234. Herrinton LJ, Liu L, Weng X, et al. Role of thiopurine and anti-TNF therapy in lymphoma in inflammatory bowel disease. Am J Gastroenterol. 2011;106:2146–53.

235. Hanauer SB, Sandborn WJ, Lichtenstein GR. Evolving considerations for thiopurine therapy for inflammatory bowel diseases: a clinical practice update: commentary. Gastroenterology. 2019;156:36–42.

236. Gisbert JP, Marín AC, Chaparro M. Systematic review: factors associated with relapse of inflammatory bowel disease after discontinuation of anti-TNF therapy. Aliment Pharmacol Ther. 2015;42:391–405.

237. Kennedy NA, Warner B, Johnston EL, et al. Relapse after withdrawal from anti-TNF therapy for inflammatory bowel disease: an observational study, plus systematic review and meta-analysis. Aliment Pharmacol Ther. 2016;43:910–23.

238. Lee MJ, Parker CE, Taylor SR, et al. Efficacy of medical therapies for fistulizing Crohn’s disease: systematic review and meta-analysis. Clin Gastroenterol Hepatol. 2018;16:1879–92.

239. Gionchetti P, Dignass A, Danese S, et al. 3rd European evidence-based consensus on the diagnosis and management of Crohn’s disease 2016: part 2: surgical management and special situations. J Crohns Colitis. 2017;11:135–49.

240. Sands BE, Anderson FH, Bernstein CN, et al. Infliximab maintenance therapy for fistulizing Crohn’s disease. N Engl J Med. 2004;350:876–85.

241. Kobayashi T, Hishida A, Tanaka H, et al. Real-world experience of anti-tumor necrosis factor alpha therapy for internal fistulas in Crohn’s disease: a retrospective multicenter cohort study. Inflamm Bowel Dis. 2017;23:2245–51.

242. Belaiche J, Louis E. Severe lower gastrointestinal bleeding in Crohn’s disease: successful control with infliximab. Am J Gastroenterol. 2002;97:3210–1.

243. Aniwan S, Eakpongpaisit S, Imraporn B, et al. Infliximab stopped severe gastrointestinal bleeding in Crohn’s disease. World J Gastroenterol. 2012;18:2730–4.

244. Kim K-J, Han BJ, Yang S-K, et al. Risk factors and outcome of acute severe lower gastrointestinal bleeding in Crohn’s disease. Dig Liver Dis. 2012;44:723–8.

245. Homan WP, Tang CK, Thorbjarnarson B. Acute massive hemorrhage from intestinal Crohn disease. Report of seven cases and review of the literature. Arch Surg Chic Ill 1960. 1976;111:901–5.

246. Alla VMB, Ojili V, Gorthi J, et al. Revisiting the past: intraarterial vasopressin for severe gastrointestinal bleeding in Crohn’s disease. J Crohns Colitis. 2010;4:479–82.

247. Kim E, Kang Y, Lee MJ, et al. Life-threatening lower gastrointestinal hemorrhage in pediatric Crohn’s disease. Pediatr Gastroenterol Hepatol Nutr. 2013;16:53–60.

248. Robert JR, Sachar DB, Greenstein AJ. Severe gastrointestinal hemorrhage in Crohn’s disease. Am Surg. 1991;213:207–11.

249. Belaiche J, Louis E, D’Haens G, et al. Acute lower gastrointestinal bleeding in Crohn’s disease: characteristics of a unique series of 34 patients. Belgian IBD Research Group. Am J Gastroenterol. 1999;94:2177–81.

250. Feagan BG, Sandborn WJ, Gasink C, et al. Ustekinumab as induction and maintenance therapy for Crohn’s disease. N Engl J Med. 2016;375:1946–60.

251. Hibi T, Imai Y, Murata Y, et al. Efficacy and safety of ustekinumab in Japanese patients with moderately to severely active Crohn’s disease: a subpopulation analysis of phase 3 induction and maintenance studies. Intest Res. 2017;15:475–86.

252. Sandborn WJ, Gasink C, Gao L-L, et al. Ustekinumab induction and maintenance therapy in refractory Crohn’s disease. N Engl J Med. 2012;367:1519–28.

253. MacDonald JK, Nguyen TM, Khanna R, et al. Anti-IL-12/23p40 antibodies for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2016;11:CD0007572.

254. Ma C, Fedorak RN, Kaplan GG, et al. Clinical, endoscopic and radiographic outcomes with ustekinumab in medically-refractory Crohn’s disease: real world experience from a multicentre cohort. Aliment Pharmacol Ther. 2017;45:1232–43.

255. Robert JR, Sachar DB, Greenstein AJ. Severe gastrointestinal hemorrhage in Japanese patients with moderately to severely active Crohn’s disease: a subpopulation analysis of phase 3 induction and maintenance studies. Intest Res. 2017;15:475–86.

256. Sandborn WJ, Gasink C, Gao L-L, et al. Ustekinumab induction and maintenance therapy in refractory Crohn’s disease. N Engl J Med. 2012;367:1519–28.

257. MacDonald JK, Nguyen TM, Khanna R, et al. Anti-IL-12/23p40 antibodies for induction of remission in Crohn’s disease. Cochrane Database Syst Rev. 2016;11:CD0007572.

258. Ma C, Fedorak RN, Kaplan GG, et al. Clinical, endoscopic and radiographic outcomes with ustekinumab in medically-refractory Crohn’s disease: real world experience from a multicentre cohort. Aliment Pharmacol Ther. 2017;45:1232–43.

259. Dayan JR, Dolinger M, Benkow K, et al. Real world experience with ustekinumab in children and young adults at a tertiary care pediatric inflammatory bowel disease center. J Pediatr Gastroenterol Nutr. 2019;69:61–7.

260. Biemans VBC, van der Meulen-de Jong AE, van der Woude CJ, et al. Ustekinumab for Crohn’s disease: results of the ICC Registry, a Nationwide Prospective Observational Cohort Study. J Crohns Colitis. 2020;14:33–45.

261. Greenup A-J, Rosenfeld G, Bressler B. Ustekinumab use in Crohn’s disease: a Canadian tertiary care centre experience. Scand J Gastroenterol. 2017;52:1354–9.

262. Vils P, Bouhnik Y, Michetti P, et al. Subcutaneous ustekinumab provides clinical benefit for two-thirds of patients with crohn’s disease refractory to anti-tumor necrosis factor agents. Clin Gastroenterol Hepatol. 2016;14:242–50.e1–2.

263. Yoshihara T, Shinizaki S, Amano T, et al. Concomitant use of an immunomodulator with ustekinumab as an induction therapy for Crohn’s disease: a systematic review and meta-analysis. J Gastroenterol Hepatol. 2021 (Online ahead of print).

264. Verstockt B, Deelenheer B, Van Assche G, et al. A safety assessment of biological therapies targeting the IL-23/IL-17 axis in inflammatory bowel diseases. Expert Opin Drug Saf. 2017;16:809–21.

265. Regueiro M, Schraut W, Baidoo L, et al. Infliximab prevents Crohn’s disease recurrence after ileal resection. Gastroenterology. 2009;136:441–450.e1 (quiz 716).

266. Zhao Y, Ma F, Chen Y-F, et al. Biologics for the prevention of postoperative Crohn’s disease recurrence: a systematic review and meta-analysis. Clin Gastroenterol Hepatol. 2015;39:637–49.

267. Sands B, Gasink C, Jacobstein D, et al. Fistula healing in pivotal studies of ustekinumab in Crohn’s disease. Gastroenterology. 2017;152:S185.

268. Vils P, Bouhnik Y, Michetti P, et al. Long-term efficacy and safety of ustekinumab in 122 refractory Crohn’s disease patients: a multicentre experience. Aliment Pharmacol Ther. 2018;47:588–95.

269. Venturin C, Nancey S, Danion P, et al. Fetal death in utero and miscarriage in a patient with Crohn’s disease under therapy with ustekinumab: case-report and review of the literature. BMC Gastroenterol. 2017;17:80.

270. Martin PL, Sachs C, Imai N, et al. Development in the cynomolgus macaque following administration of ustekinumab, a human anti-IL-12/23p40 monoclonal antibody, during pregnancy and lactation. Birth Defects Res B Dev Reprod Toxicol. 2010;89:351–63.

271. Rowan CR, Cullen G, Mulcahy HE, et al. Ustekinumab drug levels in maternal and cord blood in a woman with Crohn’s disease treated until 33 weeks of gestation. J Crohns Colitis. 2018;12:376–8.
J Gastroenterol (2021) 56:489–526

341. Lutgens MWMD, Vleggaar FP, Schipper MEI, et al. High frequency of early colorectal cancer in inflammatory bowel disease. Gut. 2008;57:1246–51.

342. Winther KV, Jess T, Langholz E, et al. Long-term risk of cancer in ulcerative colitis: a population-based cohort study from Copenhagen County. Clin Gastroenterol Hepatol. 2004;2:1088–95.

343. Ananthakrishnan AN, Cagan A, Cai T, et al. Colonoscopy is associated with a reduced risk for colon cancer and mortality in patients with inflammatory bowel diseases. Clin Gastroenterol Hepatol. 2015;13:322-329.e1.

344. Feuerstein JD, Rakowsky S, Sattler L, et al. Meta-analysis of dye-based chromendooscopy compared with standard- and high-definition white-light endoscopy in patients with inflammatory bowel disease at increased risk of colon cancer. Gastrointest Endosc. 2019;90:186-195.e1.

345. Moussata D, Allez M, Cazals-Hatem D, et al. Are random biopsies still useful for the detection of neoplasia in patients with IBD undergoing surveillance colonoscopy with chromendooscopy? Gut. 2018;67:616–24.

346. Bessissow T, Dulai PS, Restellini S, et al. Comparison of endoscopic dysplasia detection techniques in patients with ulcerative colitis: a systematic review and network meta-analysis. Inflamm Bowel Dis. 2018;24:2518–26.

347. Bisschops R, Bessissow T, Joseph JA, et al. Chromendooscopy versus narrow band imaging in UC: a prospective randomised controlled trial. Gut. 2018;67:1087–94.

348. Torres J, Caprioli F, Katsanos KH, et al. Predicting outcomes to optimize disease management in IBD patients with colorectal cancer. J Crohns Colitis. 2016;10:1385–94.

349. Eaden JA, Mayberry JF, British Society for Gastroenterology, et al. Guidelines for screening and surveillance of asymptomatic colorectal cancer in patients with inflammatory bowel disease. Gut. 2002;51(Suppl 5):V10–2.

350. Itzkowitz SH, Present DH, Crohn’s and Colitis Foundation of America Colon Cancer in IBD Study Group. Consensus conference: Colorectal cancer screening and surveillance in inflammatory bowel disease. Inflamm Bowel Dis. 2005;11:34–21.

351. Bye WA, Nguyen TM, Parker CE, et al. Strategies for detecting colon cancer in patients with inflammatory bowel disease. Cochrane Database Syst Rev. 2017;9:CD000279.

352. Uchino M, Ikeuchi H, Hata K, et al. Intestinal cancer in patients with inflammatory bowel disease at increased risk of colorectal cancer. Gastroenterology. 2011;148:639-51.e28.

353. Ananthakrishnan AN, Cagan A, Cai T, et al. Colonoscopy is recommended for surveillance of ulcerative colitis-associated colorectal cancer. Gastroenterology. 2016;151:1122–30.

354. Winther KV, Jess T, Langholz E, et al. Long-term risk of cancer in patients given anti-tumor necrosis factor therapy for inflammatory bowel disease. Clin Gastroenterol Hepatol. 2011;9:30–5.

355. van der Woude CJ, Ardizzzone S, Bengtson MB, et al. The second European evidenced-based consensus on reproduction and pregnancy in inflammatory bowel disease. J Crohns Colitis. 2015;9:107–24.

356. Waljee A, Waljee J, Morris AM, et al. Threefold increased risk of infertility: a meta-analysis of infertlity after ileal pouch anastomosis in ulcerative colitis. Gut. 2006;55:1575–80.

357. O’Morain C, Smethurst P, Doré CJ, et al. Reversible male infertility due to sulphasalazine: studies in man and rat. Gut. 1984;25:1078–84.

358. Bortoli A, Pedersen N, Duricova D, et al. Pregnancy outcome in inflammatory bowel disease: prospective European case-control ECCO-EpiCom study, 2003–2006. Aliment Pharmacol Ther. 2011;34:724–34.

359. Volkman K, Van der Woude CJ, Martens T, et al. Pregnancy and outcome in women with inflammatory bowel disease: a multicentre study from Japan. J Crohns Colitis. 2011;5:217–23.

360. Nielsen OH, Maxwell C, Hendel J. IBD medications during pregnancy and lactation. Nat Rev Gastroenterol Hepatol. 2014;11:116–27.

361. Tavernier N, Fumery M, Peyrin-Biroulet L, et al. Systematic review: fertility in non-surgically treated inflammatory bowel disease. Aliment Pharmacol Ther. 2013;38:847–53.

362. Sato A, Naganuma M, Asakura K, et al. Conception outcomes and opinions about pregnancy for men with inflammatory bowel disease. J Crohns Colitis. 2010;4:183–8.

363. Ujjhara M, Ando T, Ishiguro K, et al. Importance of appropriate pharmaceutical management in pregnant women with ulcerative colitis. BMC Res Notes. 2013;6:210.

364. Briggs GG, Freeman RR, Yaffe SJ. Drugs in pregnancy and lactation. 9th ed. Philadelphia: Lippincott Williams & Wilkins; 2011.

365. Akbari M, Shah S, Velayos FS, et al. Systematic review and meta-analysis on the effects of thiopurines on birth outcomes from female and male patients with inflammatory bowel disease. Inflamm Bowel Dis. 2013;19:15–22.

366. Lisby G, Chaparro M. Safety of anti-TNF agents during pregnancy and breastfeeding in women with inflammatory bowel disease. Am J Gastroenterol. 2013;108:1426–38.

367. Ito S. Drug therapy for breast-feeding women. N Engl J Med. 2000;343:118–26.

368. Cheent K, Nolan J, Shariq S, et al. Case Report: Fatal case of disseminated BCG infection in an infant born to a mother taking infliximab for Crohn’s disease. J Crohns Colitis. 2010;4:603–5.

369. Flint J, Panchal S, Hurrell A, et al. BSR and BHPR guideline on prescribing drugs in pregnancy and breastfeeding -Part I: standard and biologic disease modifying anti-rheumatic drugs and corticosteroids. Rheumatology. 2016;55:1693–7.

370. Briggs GG, Freeman RR, Yaffe SJ. Drugs in pregnancy and breastfeeding—Part I: standard and biologic disease modifying anti-rheumatic drugs and corticosteroids. Rheumatology. 2011;50:1186–95.

371. van der Woude CJ, Martens T, et al. The second European evidenced-based consensus on reproduction and pregnancy in inflammatory bowel disease. J Crohns Colitis. 2015;9:107–24.

372. Waljee A, Waljee J, Morris AM, et al. Threefold increased risk of infertility: a meta-analysis of infertility after ileal pouch anal anastomosis in ulcerative colitis. Gut. 2006;55:1575–80.

373. O’Morain C, Smethurst P, Doré CJ, et al. Reversible male infertility due to sulphasalazine: studies in man and rat. Gut. 1984;25:1078–84.

374. Bortoli A, Pedersen N, Duricova D, et al. Pregnancy outcome in inflammatory bowel disease: prospective European case-control ECCO-EpiCom study, 2003–2006. Aliment Pharmacol Ther. 2011;34:724–34.

375. Volkman K, Van der Woude CJ, Martens T, et al. Pregnancy and outcome in women with inflammatory bowel disease: a multicentre study from Japan. J Crohns Colitis. 2011;5:217–23.

376. Tavernier N, Fumery M, Peyrin-Biroulet L, et al. Systematic review: fertility in non-surgically treated inflammatory bowel disease. Aliment Pharmacol Ther. 2013;38:847–53.

377. Sato A, Naganuma M, Asakura K, et al. Conception outcomes and opinions about pregnancy for men with inflammatory bowel disease. J Crohns Colitis. 2010;4:183–8.

378. Ujjhara M, Ando T, Ishiguro K, et al. Importance of appropriate pharmaceutical management in pregnant women with ulcerative colitis. BMC Res Notes. 2013;6:210.

379. Briggs GG, Freeman RR, Yaffe SJ. Drugs in pregnancy and lactation. 9th ed. Philadelphia: Lippincott Williams & Wilkins; 2011.

380. Akbari M, Shah S, Velayos FS, et al. Systematic review and meta-analysis on the effects of thiopurines on birth outcomes from female and male patients with inflammatory bowel disease. Inflamm Bowel Dis. 2013;19:15–22.

381. Lisby G, Chaparro M. Safety of anti-TNF agents during pregnancy and breastfeeding in women with inflammatory bowel disease. Am J Gastroenterol. 2013;108:1426–38.

382. Ito S. Drug therapy for breast-feeding women. N Engl J Med. 2000;343:118–26.

383. Cheent K, Nolan J, Shariq S, et al. Case Report: Fatal case of disseminated BCG infection in an infant born to a mother taking infliximab for Crohn’s disease. J Crohns Colitis. 2010;4:603–5.

384. Flint J, Panchal S, Hurrell A, et al. BSR and BHPR guideline on prescribing drugs in pregnancy and breastfeeding -Part I: standard and biologic disease modifying anti-rheumatic drugs and corticosteroids. Rheumatology. 2016;55:1693–7.

385. Briggs GG, Freeman RR, Yaffe SJ. Drugs in pregnancy and breastfeeding—Part I: standard and biologic disease modifying anti-rheumatic drugs and corticosteroids. Rheumatology. 2011;50:1186–95.

386. van der Woude CJ, Martens T, et al. The second European evidenced-based consensus on reproduction and pregnancy in inflammatory bowel disease. J Crohns Colitis. 2015;9:107–24.
378. Orchard TR. Chapter 43, Extraintestinal manifestations; skin, joints and mucocutaneous manifestations. In: Satsangi J, Sutherland LR, editors. Inflammatory Bowel disease. Philadelphia: Elsevier; 2003. p. 669–97.

379. Polcz M, Gu J, Florin T. Pyoderma gangrenosum in inflammatory bowel disease: the experience at Mater Health Services’ Adult Hospital 1998–2009. J Crohns Colitis. 2011;5:148–51.

380. Ali M, Duerksen DR. Ulcerative colitis and Sweet’s syndrome: a case report and review of the literature. Can J Gastroenterol. 2008;22:296–8.

381. Iida T, Hida T, Matsuura M, et al. Current clinical issue of skin lesions in patients with inflammatory bowel disease. Clin J Gastroenterol. 2019;12:501–10.

382. Orchard TR. Management of arthritis in patients with inflammatory Bowel disease. Gastroenterol Hepatol. 2012;8:327–9.

383. Yuhara H, Steinmaus C, Corley D, et al. Meta-analysis: the risk of venous thromboembolism in patients with inflammatory bowel disease. Aliment Pharmacol Ther. 2013;37:953–62.

384. Grainge MJ, West J, Card TR. Venous thromboembolism during active disease and remission in inflammatory bowel disease: a cohort study. Lancet. 2010;375:657–63.

385. Ananthakrishnan AN, Cagan A, Gainer VS, et al. Thromboprophylaxis is associated with reduced post-hospitalization venous thromboembolic events in patients with inflammatory bowel diseases. Clin Gastroenterol Hepatol. 2014;12:1905–10.

386. Nguyen GC, Bernstein CN, Bitton A, et al. Consensus statements on the risk, prevention, and treatment of venous thromboembolism in inflammatory bowel disease: Canadian Association of Gastroenterology. Gastroenterology. 2014;146(835–848):e6.

387. Ra G, Thanabal R, Ratneswaran S, et al. Predictors and safety of venous thromboembolism prophylaxis among hospitalized inflammatory bowel disease patients. J Crohns Colitis. 2013;7:e479–85.

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