The timing of colonoscopy in patients with active ulcerative colitis (UC) lacks coherence. The published guidelines and recommendations advocate time-bound colonoscopy in patients with active UC to assess for mucosal healing. However, the practice of performing colonoscopies at fixed time frames lacks reasoning. The time to achieve mucosal healing in UC is not uniform across the patient populations and is influenced by the disease severity and efficacy and time to therapeutic response of the drugs being used. Additionally, with the availability of sensitive noninvasive inflammatory biomarkers such as fecal calprotectin, that parallel the disease activity and correlate with mucosal healing, the notion of performing colonoscopy at fixed intervals sounds unjustifiable. The authors express their view that a response-guided colonoscopy (driven by normalization of clinical symptoms and inflammatory biomarkers), rather than a time-bound colonoscopy, would be more logical, apart from being cost-effective and patient-friendly. (Intest Res 2022;20:297-302)

Key Words: Colitis, ulcerative; Colonoscopy; Fecal calprotectin

INTRODUCTION

Endoscopy (colonoscopy or sigmoidoscopy) has a key role in the management of ulcerative colitis (UC). It provides crucial information for the diagnosis (and differential diagnosis), assessment of disease extent, activity and severity, evaluation of infections during relapse or development of new unexplained symptoms, surveillance for dysplasia/malignancy and documenting response to therapy. With the availability of novel therapies for UC, the therapeutic targets have shifted from resolution of symptoms to achievement of mucosal healing (endoscopic or histologic). Achievement of mucosal healing is associated with a favorable disease course with reduced probability of relapse and lower risk of development of colorectal cancer. Various practice guidelines for management of UC, therefore, suggest periodic endoscopic examinations to look for endoscopic and histologic disease activity.

For indications such as suspected relapse or surveillance for malignancy, endoscopic assessment is considered gold standard. However, ambiguity surrounds the timing and frequency of doing endoscopy for assessment of disease activity and mucosal healing after initiation of therapy. The Selecting Therapeutic Targets in Inflammatory Bowel Disease (STRIDE I and STRIDE II) programs initiated by the International Organization for the Study of Inflammatory Bowel Diseases (IOIBD) suggest endoscopic mucosal assessment every 3–6 months after initiation of therapy. The European Crohn's and Colitis Organisation (ECCO) and the European Society of Gastrointestinal and Abdominal Radiology (ESGAR) guidelines also recommend assessing clinical responders by endoscopy every 3–6 months, though the option of being guided by fecal calprotectin (FC) has also been propounded. This time-bound repeated endoscopic assessment for mucosal healing is likely to result in very frequent invasive procedures which may be un-
comfortable to patients. Additionally, frequent interventions add to the cost of the therapy. In real-world scenarios, therefore, following recommendations of time-bound endoscopy do not seem practical. There is hence a need to rethink and reposition endoscopy, performed for assessment for mucosal healing, in the management algorithm for patients with UC.

**IS ENDOSCOPIC ASSESSMENT A REQUISITE FOR MONITORING DISEASE ACTIVITY IN UC?**

Conventionally, evaluation of clinical symptoms and endoscopic assessment have been the 2 parameters to monitor disease activity in patients with UC. Resolution of clinical symptoms (increased frequency of stools, urgency and rectal bleeding) is the first therapeutic target. A parallelism between clinical symptoms and endoscopic disease activity has been described.\(^2\)\(^1\) Partial Mayo Clinic score (including only the clinical components of disease activity) correlates well with the total Mayo Clinic score (including both clinical and endoscopic components of disease activity).\(^1\)\(^1\)\(^ -\)\(^1\)\(^2\) A careful assessment of the clinical symptoms can, therefore, correlate with the severity of inflammatory endoscopic lesions. However, discordance between clinical symptoms and disease activity may exist. Even with complete abatement of symptoms (i.e., symptomatic remission), there could be endoscopically and histologically active disease.\(^1\)\(^3\)\(^ -\)\(^1\)\(^5\) Nearly half of the patients in symptomatic remission have been reported to have evidence of active disease on endoscopy.\(^1\)\(^6\)\(^ -\)\(^1\)\(^8\) Conversely, a proportion of patients with endoscopic mucosal healing may have persistent symptoms due to superimposed irritable bowel syndrome (IBS).\(^1\)\(^9\)\(^,\)\(^2\)\(^0\) Nevertheless, it is important to rule out active inflammation in such patients who have supposedly functional symptoms.

The disagreement between clinical symptoms and degree of inflammation in a subgroup of patients led to the development of noninvasive biomarkers of inflammation such as erythrocyte sedimentation rate, C-reactive protein, and FC. Of these, FC has been demonstrated to correlate with clinical disease activity (partial Mayo Clinic score and endoscopic disease activity (endoscopic Mayo Clinic score and Ulcerative Colitis Endoscopic Index of Severity)).\(^2\)\(^1\) FC also has moderate correlation with the histological indices of disease activity.\(^1\)\(^4\)\(^,\)\(^1\)\(^6\)\(^,\)\(^1\)\(^7\)\(^,\)\(^2\)\(^2\) It has been hypothesized that endoscopic and histological resolution of inflammation will result in reduction in the amount of neutrophil migration into the gut lumen and therefore low FC values.\(^2\)\(^3\)\(^,\)\(^2\)\(^4\) Thus, FC appears to be a reliable marker to monitor treatment response in UC.

Various concentration thresholds of FC have been proposed across correlation studies to predict mucosal healing. The FC cutoff values depend upon the commercial kit used, type of assay and the population to which the test is applied. Theede et al.\(^2\)\(^5\) demonstrated that a cutoff level of FC of 192 μg/g correlated with endoscopic evidence of mucosal healing, while a cutoff of 171 μg/g identified patients with histologic evidence of mucosal healing. FC values < 150 μg/g have been shown to correlate with endoscopic Mayo Clinic score of 0.\(^1\)\(^9\) In another retrospective analysis, an FC level of ≤ 60 μg/g predicted endoscopic Mayo Clinic score 0/1 and Nancy score ≤ 1.\(^2\)\(^5\) A systematic review and meta-analysis proposed a cutoff of ≤ 50 μg/g to be predictive of mucosal (both endoscopic and histologic) healing.\(^2\)\(^2\) On the other hand, FC > 250 μg/g was associated with mucosal/histologic activity in a majority of the patients.\(^2\)\(^6\) Basis the existing literature, FC values less than 50–150 μg/g in patients with UC may suggest endoscopic (endoscopic Mayo Clinic score 0 or 1) and/or histological mucosal healing, though a single cutoff value to discriminate between active and inactive UC, using endoscopy as the reference, has not been validated as yet. Performing colonoscopy in patients with high FC (> 150 μg/g), at the suggested 3–6 months’ time frame, is unlikely to reveal mucosal healing which, as a matter of fact, is the very purpose of doing colonoscopy.

FC can also identify patients in clinical remission who are at risk for an impending relapse. A high FC (> 50–150 μg/g) in patients in clinical remission correlates with an increased probability of relapse over the next 2–3 months.\(^2\)\(^7\)\(^ -\)\(^2\)\(^1\) It has been reported that patients who relapse, have high FC values 4–6 months before the apparent clinical relapse. Consecutive normal FC values, on the other hand, are associated with a high probability of maintaining remission over the next couple of months.\(^2\)\(^8\)\(^,\)\(^2\)\(^2\) Additionally, FC values at 3 months after initiation of therapy in patients with new-onset UC have been documented to predict the subsequent disease course.\(^3\)\(^3\) Serial testing of FC can hence aid in therapeutic decision making regarding timing of the endoscopy. Two consecutive normal FC values, done at an interval of 2–3 months, are likely to correlate with absence of inflammatory activity on endoscopy.

The FC values can also differentiate between IBS and inflammatory bowel disease. In patients with symptoms attributed to IBS, a normal FC would suggest against endoscopic activity and prevent an unwanted endoscopy. As is evident, evaluation of clinical symptoms and inflammatory biomarkers can provide adequate information about disease activity. Performing endoscopy upfront, for assessment...
of mucosal healing in all patients, therefore, is not an obligation and the decision to perform endoscopy has to be individualized based on the response to therapy.

WHEN TO DO ENDOSCOPIC ASSESSMENT FOR MUCOSAL HEALING?

An endoscopic assessment performed too early or delayed for long would not meet the objective(s) of performing endoscopy. As is evident in various induction trials in UC, early endoscopic assessment (within 12–16 weeks) yields mucosal healing rates varying between 25% and 50% only. Similarly, delaying endoscopy for too long may result in suboptimal therapy. Therefore, it is important to perform endoscopy for mucosal assessment at the right time.

An important determinant of the right time to perform en-

Fig. 1. Proposed approach to monitoring of a patient after initiation of therapy. Symptoms include increased frequency of stools, rectal bleeding and urgency; Assessment of symptoms to be done between 4 and 24 weeks depending on the expected time to therapeutic response of the drug being used.
Endoscopic assessment in UC is the "time to therapeutic response" for a particular therapeutic agent. The initial response to therapy is resolution of clinical symptoms (rectal bleeding, frequent stools, and urgency). This generally occurs as early as 3–5 days with intravenous corticosteroids, takes around 2–8 weeks with most of the other therapeutic agents (5-aminosalicylates, oral corticosteroids, infliximab, adalimumab, and golimumab), and may even be prolonged up to 10–20 weeks, as with certolizumab and vedolizumab. Immunomodulators like thiopurines might take up to 3–4 months for reaching their maximal therapeutic efficacy, and endoscopic resolution may take even longer than that. Therefore, repeating endoscopy before the expected time to therapeutic response may not be rational. With the availability of sensitive noninvasive inflammatory biomarkers like FC that correlate with the clinical, endoscopic and histological disease activity, the timing of endoscopic assessment, for assessment of mucosal healing, can be guided by serial FC values, in combination with the clinical symptoms.

In light of the published evidence, it may be put forward that patients who achieve symptomatic remission should be tested for biomarker remission using FC every 3 months, and when 2 consecutive values of FC are <50–150 μg/g, endoscopy should be considered to look for mucosal healing. If mucosal healing is achieved, the therapeutic agent is continued to sustain remission and further endoscopic assessments are not needed, unless there is clinical relapse (worsening/new symptoms and elevated FC), suspected complications or need for surveillance for colorectal cancer. Patients who do not achieve endoscopic mucosal healing should be evaluated for optimization of therapy (Fig. 1).

IS ENDOSCOPY REQUIRED FOR PATIENTS WITH ACTIVE DISEASE?

In the authors’ opinion, the indications of performing endoscopy in patients with active disease include evaluation of reasons for nonresponse; for example, infections like cytomegalovirus, Epstein-Barr virus, etc., misdiagnosis of Crohn’s disease as UC, presence of colorectal cancer or true drug resistance.

As majority of the patients with UC have rectal involvement and the maximal disease severity is seen in the distal colon, a limited unprepared sigmoidoscopy can suffice. However, in patients with persistent symptoms and a normal/near normal distal colon, full length colonoscopy should be performed.

CONCLUSION

The appropriate intervals for endoscopic assessment of mucosal healing after initiation of therapy are not clearly defined. "Time bound" endoscopic assessment of all patients every 3–6 months, as recommended by STRIDE/ECCO-ESGAR guidelines, may neither be practical nor useful in real-world scenarios. Frequent endoscopies are also likely to encounter resistance from patients. Endoscopic assessments for mucosal healing should, therefore, be considered once symptomatic and biomarker remissions have been achieved (with the aim to document mucosal healing), rather than as a predetermined time-bound drill. We must not allow the clock and the calendar to blind us to the fact that decision about the timing of endoscopy needs to be individualized. A “response guided” stepwise approach of attaining symptomatic remission, biomarker remission and then evaluating for endoscopic (and histological) remission may be more rational, cost-effective and patient-friendly.

ADDITIONAL INFORMATION

Funding Source
The authors received no financial support for the research, authorship, and/or publication of this article.

Conflict of Interest
Sood A is an editorial board member of the journal but was not involved in the peer reviewer selection, evaluation, or decision process of this article. No other potential conflicts of interest relevant to this article were reported.

Data Availability Statement
Not applicable.

Author Contribution
Conceptualization: Sood A. Project administration: Sood A, Midha V. Supervision: Sood A. Visualization: Mahajan R, Singh A. Midha V. Writing - original draft: Mahajan R. Writing - review & editing: Sood A, Mahajan R, Singh A, Mehta V. Approval of final manuscript: all authors.

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