Evaluation of the severity of ulcerative colitis using endoscopic dual red imaging targeting deep vessels

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ABSTRACT
Background and study aims Colonoscopies can predict long-term prognoses in patients with ulcerative colitis (UC). Recently, a new imaging technology has been developed that uses 3 types of illumination with center wavelengths of 540 nm, 600 nm, and 630 nm. The use of both the 600-nm and 630-nm lights (Dual red imaging; DRI) is critical for identifying blood vessels in deeper tissue. The aim of this study was to evaluate the usefulness of DRI for assessing the severity of inflammation in patients with UC.

Patients and methods A total of 43 UC patients were retrospectively enrolled to evaluate the endoscopic severity of 112 colon segments, and Mayo endoscopic scores, DRI scores and the severity of inflammation on a visual analogue scale (VAS) were compared. The Mayo endoscopic scores, DRI scores, and histologic scores were evaluated, and the interobserver agreement on DRI scores among 5 investigators was also assessed. The usefulness of DRI scores for predicting prognoses was also assessed in patients with clinical remission.

Results The DRI scores were closely correlated with the VAS for the severity of colonic inflammation (r = 0.96) and the histologic scores (r = 0.72–0.8). The DRI scores had a higher rate of interobserver agreement (κ values = 0.63–0.88) than the Mayo endoscopic scores (κ values = 0.44–0.59). Inter-observer agreement between 4 non-experts was also excellent (mean κ value = 0.76, range 0.63–0.82). The expected time until recurrence was significantly longer in patients with lower DRI scores (P < 0.01).

Conclusion DRI can be used in patients with mild to moderate endoscopic severity because it targets the deep vascular pattern. The prognosis of UC can be predicted by assessing deep vessels using DRI.

Introduction
Narrow-band imaging (NBI) was developed to enhance the contrast of the capillary pattern in the superficial layer of the gastrointestinal tract [1–5]. NBI at 415 nm is a useful method to support endoscopic observations of early stage cancers, such as esophageal cancer [3]. The combination of the NBI technique with magnification endoscopy is useful for detecting early gastrointestinal cancers and for estimating the depth of cancer invasion by assessing vascular and surface patterns [4, 5]. In active ulcerative colitis (UC), however, the vascular pattern seen using NBI is not as clear (Supplemental Fig. 1).

Recently, a new imaging technology has been introduced that uses 3 wavelengths for illumination: 540 nm, 600 nm and 630 nm. The 600-nm and 630-nm lights are critical for identifying blood vessels in the submucosal tissue and bleeding points.

Thus, dual red imaging (DRI) could be a promising way to avoid severe bleeding during endoscopic submucosal resection for the treatment of early stage gastrointestinal tumors [6, 7]. With respect to the identification of blood vessels in the submucosal tissue, DRI may detect deeper vessels in UC patients (Supplemental Fig. 1c). Because DRI can primarily enhance vessel patterns, we hypothesized that, by using DRI, differences among investigators’ assessments of the severity of inflammation would be minimal. However, no study has evaluated the usefulness of DRI in UC patients.

The aim of this study was to evaluate the usefulness of DRI for assessing the severity of inflammation in patients with active UC. The assessment of UC lesions in patients with mild or moderate disease is important for assessing the severity of inflammation, and discrepancies among investigators have been
observed in UC. Therefore, we evaluated the usefulness of DRI in UC patients with mild to moderate endoscopic severity.

Patients and methods

Patients

From March 2012 to April 2013, a total of 43 patients previously diagnosed with UC at the outpatient department of Keio University Hospital were enrolled in this study. All UC patients were diagnosed according to the criteria of the Research Committee on Inflammatory Bowel Disease in Japan [8]. Patients with fulminant disease were excluded because they required immediate intensive medical therapeutics, hospitalization, or surgery. This study was retrospectively conducted, and the clinical indices were prospectively collected during regular visits.

Dual-red imaging

This new imaging technology uses illumination at 3 wavelengths: 540 nm, 600 nm and 630 nm. With respect to the identification of blood vessels in the submucosal tissue, the 600-nm and 630-nm lights can penetrate the deeper tissue because the scattering properties of these lights are minimal. Furthermore, light absorption by blood vessels is much stronger at 600 nm than at 630 nm. Therefore, the intensity of light reflected by blood vessels at 630 nm is greater than that at 600 nm. This difference allows detection of blood vessels in the deeper tissue using the intensity balance of each wavelength of reflected light rather than white-light imaging (WLI) and DRI. The mechanism underlying how blood flow and bleeding can be more easily identified using DRI compared with conventional imaging methods is being actively studied.

Colonoscopy

Total ileocolonoscopies (ICS) were performed on all 43 patients. All patients were required to drink 2000 mL to 3000 mL of polyethylene glycol (Ajinomoto Farma, Co, Ltd, Tokyo, Japan) before the procedure. Pethidine hydrochloride (Tanabe-Mitsubishi, Tokyo, Japan) was administered immediately prior to ICS to reduce patient burden. All colonoscopy images were recorded on a digital versatile disc. Physicians could instantaneously switch from WLI to DRI by pressing a button on the control handle of the colonoscope. WLI and DRI recordings were taken in the ascending colons, sigmoid colons, and rectums of all 43 patients. To reduce the risk of bleeding caused by biopsies, we evaluated the severity of inflammation for only 3 colonic segments. In all cases, at each segment, the WLI was recorded first, and then the DRI was recorded at the same lesions. A total of 129 videos were collected, and 17 videos were excluded due to insufficient bowel cleaning. Finally, 112 videos (37 from the ascending colon, 35 from the sigmoid colon, 40 from the rectum) were collected to evaluate the endoscopic severity of inflammation using WLI and DRI.

Scoring the severity of inflammation

Five endoscopists independently evaluated the severity of mucosal inflammation using the Mayo endoscopic score [9] for the WLI videos of the 112 segments of the colon. One was an expert in diagnosis of UC (approximately 400 colonoscopies for UC annually), and the other 4 investigators were non-experts (50 cases annually). The endoscopic severity of inflammation seen using DRI was also evaluated using a novel DRI score. The Mayo score was classified as follows: below 0, normal; 1, erythema, decreased vascular pattern, mild friability; 2, marked erythema, absent vascular pattern, friability, erosion; 3, spontaneous bleeding, ulcerations. The DRI scores were classified as DRI 1, 2, 3, and 4. DRI1 was defined as a normal vascular pattern in both the superficial (brown) and deep (green) vessels. DRI2 was defined as the patchy or complete obliteration of the brown vessels with clear reorganization of the green vessels. DRI3 was defined as patchy obliteration of the green vessels, and DRI4 was defined as the complete obliteration of the green vessels. The overall severity was also assessed on a visual analogue scale (VAS). VAS scores ranged from 0 (completely normal) to 100 (most severe).

Histologic severity of inflammation was scored by a pathologist using the Geboes score [10] as follows: grade 0, structural and architectural changes; grade 1, chronic inflammatory infiltrate; grade 2, neutrophils and eosinophils in the lamina propria; grade 3, neutrophils in the epithelium; grade 4, crypt destruction; grade 5, erosions or ulceration. The pathologist was blinded to the clinical information.

Assessment of clinical prognosis in patients with various DRI scores

The usefulness of DRI scores for predicting clinical outcomes was assessed. Partial Mayo scores were recorded on the patients’ medical charts at each visit to the clinic. Clinical remission was defined as a partial Mayo score ≤ 1. The rate of clinical recurrence was analyzed for patients with each DRI score (from DRI1 to DRI4). Clinical recurrence was defined as a partial Mayo score ≥ 3.

Data collection and statistical assessment

Our main endpoints in this study were as follows: 1) assessment of the correlation between the DRI score and the Mayo endoscopic score and between the DRI score and the histological grade, 2) inter-observer agreement on the DRI score and the Mayo endoscopic score, and 3) assessment of the correlation between the DRI score and the time until clinical recurrence in patients with clinical remission.

We evaluated the correlation between the DRI score, the Mayo endoscopic score and the histopathological grade using Spearman’s rank correlation coefficient. To validate the DRI score, interobserver agreement between 2 endoscopists was examined, and the kappa value was calculated. Clinical information, such as age, gender, duration of disease, disease type (first attack, relapsing/remission or chronic continuous) and the use of medical treatments (at the time of the colonoscopy and at each visit to the clinic), was collected. Although we did not schedule visits to the clinic, most patients visited our hospital every 8 to 12 weeks.

In patients with clinical remission, the association between the DRI score and the time until clinical recurrence was assessed using the Kaplan-Meier method, and the difference be-
The clinical features of the patients and their medications for UC are listed in Table 1. Forty-three patients with UC were included; of them, 36 (84%) and 5 (12%) were taking oral 5-aminosalicylates and thiopurine, respectively, whereas 7 patients did not take any medication for UC. The mean partial Mayo score of this cohort was 1.2. Serum CRP levels ranged from 0.01 to 1.35 mg/dL, and CRP levels were within the normal range (less than 0.35 mg/dL in our institution) in 84% of patients.

Correlation between DRI score and VAS score for the severity of inflammation

Fig. 1 indicates the typical findings for each DRI score in patients with UC. A total of 112 segments were assessed for the severity of inflammation using the Mayo endoscopic score, the DRI score and the VAS score. The mean DRI and VAS scores were 2.9 ±1.2 (range: 1–4) and 25.0 ±18.1 (range 4.5–65.5), respectively. The distribution of severity scores using the VAS (0–100 points) with DRI is presented in Fig. 2. The VAS score was significantly correlated with the DRI score (r =0.957, P < 0.001). The median VAS score in patients scored as DRI1, 2, 3, and 4 was 6.5 (interquartile range, IQR =2.6), 18.0 (IQR = 5.0), 35.0 (IQR =7.0), and 52.5 (IQR = 8.3), respectively.

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**Results**

**Patient characteristics at baseline**

The clinical features of the patients and their medications for UC are listed in Table 1. Forty-three patients with UC were included; of them, 36 (84%) and 5 (12%) were taking oral 5-aminosalicylates and thiopurine, respectively, whereas 7 patients did not take any medication for UC. The mean partial Mayo score of this cohort was 1.2. Serum CRP levels ranged from 0.01 to 1.35 mg/dL, and CRP levels were within the normal range (less than 0.35 mg/dL in our institution) in 84% of patients.

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Correlation between DRI score, endoscopic score and histopathology

Next, we assessed the relationship between the DRI score and the Mayo endoscopic score. As Table 2 indicates, the DRI score was significantly correlated with the Mayo endoscopic score in analyses by both experts and non-experts ($r = 0.695$–$0.778; P < 0.001$ in each investigator). The relationship between the DRI score and the histological grade was significantly similar for each investigator (Table 3).

Interobserver agreement was better for the DRI score than for the Mayo endoscopic score

The $\kappa$ value validating the interobserver agreement on the DRI score was moderate to excellent (mean $\kappa$ value $= 0.76$, range $0.63 – 0.88$), as Table 4a indicates. Interestingly, interobserver agreement was even excellent among the analyses from 4 non-experts (mean $\kappa$ value $= 0.76$, range $0.63 – 0.82$). However, relatively lower interobserver agreement among the 5 investigators was observed for the Mayo endoscopic score assessments, with a mean $\kappa$ value of $0.53$ (Table 4b), and this agreement was significantly lower than the agreement on DRI scores ($P < 0.001$). Intra-observer agreement values for the Mayo endoscopic score and the DRI score were comparable (the $\kappa$ values were $0.89$ and $0.91$, respectively).

DRI score and the prediction of prognosis in patients with clinical remission

We next assessed whether endoscopic severity assessed using DRI at baseline was associated with an increased risk of clinical recurrence among 35 patients in clinical remission. In this analysis, the highest DRI score (the most severe score) among the segments was used. Over a mean follow-up of $25.3 \pm 9.3$ months, 9 patients ($43\%$) experienced clinical recurrence, whereas the other 26 patients remained in clinical remission. No recurrence was observed in patients scored as DRI1. The recurrence rates in patients scored as DRI2, 3, or 4 were $15\%$
in the lower DRI group and 20.7 months (95% CI 14.3 – 27.1) in the higher DRI group.

Fig. 3 Non-recurrence rates in patients with various DRI scores that were in clinical remission at baseline. The expected time until recurrence was significantly longer in patients with lower DRI scores (log-rank test, Breslow test $P = 0.002$).

(2/13), 50% (5/10), and 67% (2/3), respectively. The expected time until recurrence was significantly longer in patients with lower DRI scores (Fig. 3, log-rank test, Breslow test $P = 0.002$). When patients were divided into a lower DRI group (DRI 1 – 2) and a higher DRI group (DRI 3 – 4), the mean expected time until recurrence was 33.2 months (95% CI 30.7 – 35.7) in the lower DRI group and 20.7 months (95% CI 14.3 – 27.1) in the higher DRI group.

Discussion

UC is a chronic inflammatory bowel disease characterized by blood in the stool, diarrhea, abdominal pain, and extra-intestinal symptoms. Endoscopy plays a pivotal role in the assessment of the severity of UC, and endoscopic assessment is a critical endpoint for assessing the efficacy of medications. Endoscopic remission gives an improved prognosis in patients with clinical remission without mucosal healing than in those with Mayo endoscopic scores of 0 – 1 than in those with Mayo endoscopic scores of 2 – 3. Furthermore, significant differences between Mayo endoscopic scores of 0 and 1 that were detected 8 weeks after starting on IFX could predict clinical remission at week 30. These results suggest that endoscopy is critical for assessing the severity of inflammation and can predict medium- to long-term prognoses in UC patients after the onset of remission.

Although original and modified Baron scores [15, 16] and the Mayo score for proctosigmoidoscopy [17] have been widely used and are easy for investigators to use, recent studies have demonstrated that the rate of agreement is only 27% for endoscopic remission (Baron score 0) and 37% for moderate activity (Baron 2). Inter-observer disagreement on the severity of inflammation has been reported for UC [18]. Thus, the development of new methods/techniques to assess severity is warranted. More recently, the Ulcerative Colitis Endoscopic Index of Severity (UCEIS) and the Ulcerative Colitis Colonoscopic Index of Severity (UCCIS) were developed to solve the problem of low interobserver agreement on endoscopic scores [18 – 20]. An assessment of vascular patterns including both endoscopic indices revealed excellent interobserver agreement for the vascular pattern ($\kappa$ value = 0.74 – 0.86), and the intraobserver agreement for the vascular pattern ($\kappa$ value = 0.87) was better than that for bleeding ($\kappa$ value = 0.47).

DRI was developed to easily identify blood vessels in deeper tissue. In this study, the usefulness of DRI for assessing the severity of UC was investigated by focusing on only the mucosal and submucosal vessels. Usually, the vascular pattern in a conventional colonoscopy is partially identifiable or is absent in cases of mild to moderate UC inflammation. DRI can primarily enhance the vessel pattern and the identification of deep vessels. Importantly, deep vessels can be detected even when the vascular pattern is partially visible or has disappeared using WLI. Therefore, by targeting the deep vascular pattern, DRI can be used in patients with mild to moderate endoscopic severity. Because DRI enhances the vascular pattern, inter-observer disagreement might be minimal. In fact, our study indicated that the $\kappa$ values for the DRI scores between 2 investigators were better than those for the previously used endoscopic score. Furthermore, our study also indicated that the clinical prognosis is better in patients with deep vessels observed using DRI (DRI = 2) than in patients with partially visible vessels (DRI = 3). Thus, DRI is useful for predicting the middle- to long-term prognosis of UC patients. Furthermore, DRI could be used to evaluate colonic inflammation after beginning intensive medical treatment (e.g., anti-TNF agents, steroids) when the vessel pattern is absent using conventional WLI.

There are some limitations to this study. The investigators observed videos of WLI before DRI in all cases. Thus, assessment of severity based on DRI was affected by the WLI. In this study, we did not use high-definition endoscopy. However, our recent preliminary study using high-definition endoscopy indicated that the interobserver agreement on Mayo endoscopic scores was not as high (data not shown, $\kappa$ value = 0.66) compared with that using DRI. Furthermore, a pathologist assessed the histopathological severity in this study, therefore, inter-observer validation of the histopathological assessment was not considered. The usefulness of DRI for UC should be assessed in a larger study. Finally, in this study, we did not assess the endoscopic usefulness of white light imaging in predicting clinical
outcomes using endoscopic indices such as the Mayo score. A previous study indicated that a Mayo endoscopic score of 0 (complete endoscopic remission) predicted a better clinical outcome, and significant differences have been identified between a Mayo endoscopic score of 0 and 1 (mild endoscopic severity). However, a Mayo endoscopic score of 1 was observed in almost 70% of the patients in our cohort. Thus, we did not confirm the significant differences that have been identified between a MES of 0 and 1 in this study. Importantly, DRI could separate patients with the same Mayo endoscopic score into groups with DRI scores of 2 or DRI scores of 3 – 4.

Conclusion

In conclusion, DRI is a novel, simple tool for assessing the severity of inflammation in UC patients. The DRI score was correlated with the VAS score for the severity of colonic inflammation and exhibited a high rate of inter-observer agreement. The DRI score was closely correlated with both the endoscopic and the histological grades. The prognosis of UC can be predicted by assessing deep vessels using DRI.

Competing Interests

Colonoscopy with DRI capability was provided by the Olympus company. However, the company representative did not contribute either to the performance or the planning of this study, and did not write or help to write the manuscript.

References

**Supplemental Fig. 1** Rectal mucosa in patients with mild inflammation. 

- **a** The vessel pattern was obliterated using white light imaging, and vessel information was not observed using narrow-band imaging.
- **b** Dual-red imaging indicated that the superficial vasculature was patchy (brown), and deep vessels (green) were also observed.