How disease extent can be included in the endoscopic activity index of ulcerative colitis: the panMayo score, a promising scoring system

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Abstract

Background: Colonoscopy plays crucial role in the establishment of the diagnosis, management and follow-up of ulcerative colitis (UC). None of the currently widely used endoscopic scores consider disease extent, and therefore do not correlate with the real severity of UC. Our aim was to assess the accuracy of a new score, the Pancolonic Modified Mayo Score that can reflect not only the severity, but the extent of active UC.

Methods: One hundred and four UC patients were enrolled in this prospective study. The Endoscopic Mayo Scores of the involved area of the five colorectal segments were added; furthermore, the sum was multiplied by 3 in case of eMayo ≥ 2 (range 0 [normal] to 45 [most severe]) to obtain the Pancolonic Modified Mayo Score (panMayo) in order to clearly distinguish the active and inactive disease. We analysed the correlation of panMayo Score with eMayo and Ulcerative Colitis Endoscopic Index of Severity (UCEIS) and complicated disease outcome. We compared the endoscopic indices with serum and faecal inflammatory parameters and Riley Score.

Results: The panMayo Score correlated with eMayo and UCEIS. Every endoscopic score showed correlation with Riley Score, CRP, haemoglobin, haematocrit, serum iron, faecal MMP-9 and calprotectin and also predicted a complicated disease outcome. Only panMayo score correlated exclusively with the extent of UC.

Conclusions: We suggest that this new score gives additional information about disease extent besides disease activity with a strong correlation with laboratory parameters of inflammation and with the other widely used endoscopic indices.

Keywords: Ulcerative colitis, Endoscopic Mayo subscore, UCEIS, Endoscopy, Colonoscopy

Background

Ulcerative colitis (UC) is a chronic relapsing inflammatory disease, therefore monitoring disease activity has a great importance in optimising the therapy based on relevant signs and symptoms. Several different scoring systems are available to assess endoscopic disease activity in UC; however, multiple existing indices indicate the lack of an optimal one. On the other hand, mucosal healing becomes one of the desired therapeutic endpoints of inflammatory bowel diseases nowadays [1]. Thus colonoscopy plays a crucial role not only in the establishment of diagnosis, but also in the management and therapeutic decision-making of UC. An exact scoring system for the evaluation of the mucosa is absolutely mandatory for the comparable, standard care of patients with UC all over the world [2].

Truelove and Witts were the first who described a scoring system to evaluate mucosal appearance as a measure of activity for UC [3]. Several scoring indices have evolved during the last decades. The most widely used ones are the Mayo Score, and the validated [4] Ulcerative Colitis Endoscopic Index of Severity (UCEIS) [1]. Both of these scores assess vascular pattern, presence of erythema, friability, erosions, ulcerations and bleeding [5, 6]. Some research confirmed that mucosal healing is associated with...
better disease outcome [7]. Histologic assessment and scoring is therefore important for the evaluation of UC activity predicting remission rates and risk of malignancy or surgery. Currently, one of the most frequently used histopathological scores is Riley Score [8]. Riley Score incorporates the following histological features: (a) presence of acute inflammatory cell infiltrate (neutrophils in the lamina propria), (b) crypt abscesses, (c) mucin depletion, (d) surface epithelial integrity, (e) chronic inflammatory cell infiltrate (round cells in the lamina propria), and (f) crypt architectural irregularities [7, 8].

Based on a study from Japan, colonoscopy is preferred over sigmoidoscopy for the evaluation of inflammation in UC [9]. Disease extension has a significant implication for prognosis due to the association between extensive UC and colectomy and colorectal cancer; nevertheless, disease localisation and extent determine treatment choice, as well [10]. However, none of the currently widely used endoscopic scores consider disease extent, and therefore they do not correlate with the real severity of UC outcome. Our aim was to assess the accuracy of a new score (Pancolonic Modified Mayo Score) that as a matter of fact is a modified Endoscopic Mayo Subscore (eMayo) in order to reflect not only the activity, but the extent of UC.

Methods
Study population
This single-centre, prospective study was carried out on consecutive patients with confirmed diagnosis of UC between January 2011 and October 2015. Diagnosis was based on the European evidence-based Consensus on the diagnosis and management of UC (ECCO) [11]. Disease phenotype and extension were determined according to the Montreal Classification [12] and based on the affected anatomical segments (rectum, sigmoid colon, descending, transverse and cecum/ascending colon). Subjects with UC of varying disease activities and extent were enrolled. Patients with Crohn’s disease or cases with incomplete colonoscopy were excluded. One hundred and four patients were included finally, who were examined by complete colonoscopy which performed by one of three experienced endoscopists (T.M., Z.SZ. and F.N.) in accordance with protocols. All patients had given their written informed consent to the procedure. Observers, who carried out the colonoscopy, rated mucosal lesions (vascular pattern, granularity, friability, ulceration) and the severity of inflammation for each segment and the maximum extension of involvement. Thus, panMayo Score, eMayo and UCEIS were determined in every patient. Biopsy specimens were taken from the most severely affected area for histological examination and scoring. Riley Score was determined for each patient by an experienced gastropathologist.

Laboratory and faecal markers
We collected blood and faecal samples from every patient to determine specific laboratory parameters for inflammation, namely C-reactive protein (CRP), white blood cell count (WBC), faecal calprotectin and matrix-metalloproteinase (MMP)-9, in addition to haemoglobin, haematocrit, thrombocytes and serum iron level. Faecal MMP-9 concentrations were measured using ELISA (Quantikine MMP9 assay, R&D System, UK). One gram of the sample was diluted, homogenised and centrifuged twice; the final supernatant was filtered and stored at −20 °C until analysis that was in accordance with the manufacturer’s instructions [13]. Faecal specimens were stored at −20 °C, thawed and prepared for a calprotectin quantitative lateral flow assay as described by the manufacturer (Quantum Blue, Bühlmann Laboritories, Switzerland).

Pancolonic modified Mayo score
Pancolonic Modified Mayo Score (panMayo) was calculated with the combination of disease extension and severity (Table 1). The eMayo Score of the five colorectal segments (ascendending, transverse, descendending, sigmoid colon and rectum) was determined separately and added afterwards. Finally, the sum was multiplied by the Inflammatory Constant if eMayo was ≥2 at least in one segment to clearly distinguish between the active and inactive disease. Inflammatory constant (IC = 3) was defined at the beginning of the study as the smallest number that can equilibrate the difference between the sum of the inactive and active cases

| Involvement (points) | eMayo (points) |
|---------------------|---------------|
| 0                   | 0             |
| 1                   | 1             |
| 2                   | 2 × 3 = 6     |
| 3                   | 3 × 3 = 9     |
| 4                   | 4 × 3 = 12    |
| 5                   | 5 × 3 = 15    |

Table 1: Calculation of panMayo score
Explanation:
Involvement: rectum: 1 point, rectum-sigmoid: 2 points, descending colon: 3 points, transverse colon: 4 points, cecum/ascending colon: 5 points
If no Inflammation (eMayo 0–1 points): eMayo points x involvement [1–5]
If active inflammation is presented in any segment: eMayo points x involvement [1–5] x 3
resulting a broad range from 0 [normal] to 45 [most severe, pancolitis with eMayo 3 in each segment]). Inclusion of IC is important, because calculation without IC results the same points in some cases, thus it would be impossible to distinguish different extent and activity by panMayo Score. The most important unmet need for widely used scoring systems in clinical practice that the score simultaneously reflect on the severity and the extension of the disease. The aim of panMayo score was to compensate this gap. If you heard that the eMayo score is 3 points, you do not know that what is the extension of disease and what is therefore exactly the clinical severity of the disease. In contrary, panMayo 45 points is a pancolitis with severe inflammation in all segments.

We analysed the correlation of panMayo Score with the endoscopic indices and the complicated disease outcome. The panMayo Score was compared with Endoscopic Mayo Score and UCEIS to assess the utility of our new score. Complicated disease outcome was defined as the need of hospitalization and/or colectomy during the follow-up.

Statistical analysis

Correlation of the panMayo Score with clinical, endoscopic and histological indices (partial Mayo Subscore, eMayo Subscore, UCEIS, Riley Score), laboratory measures of disease activity and anaemia (C-reactive protein, faecal calprotectin, faecal MMP-9, serum iron, haemoglobin, haematocrit etc.) were estimated using the multiple correlation analysis with Spearman method. Logrank test was used to determine connection between assessed scores and hospitalisation/colectomy rates. Receiver operating characteristic (ROC) curve analysis was performed to determine the predictive power of the panMayo Score level. Area under ROC curve (AUC), confidence interval for AUC, sensitivity and specificity were calculated. The cut-off level was determined by the maximum value of Youden’s index (sensitivity + specificity-1). The statistical analysis was performed using R (version 3.2.0) statistical software. $P < 0.05$ was considered as statistically significant.

Results

Patient characteristics

Out of 104 outpatients, 58 were female and 46 were male; the mean age at enrolment was $43.9 \pm 14.7$ years. The mean duration of the disease was $8.3 \pm 0.7$ years. Left-sided colitis was the most common extent (Table 2). On the basis of Mayo Score, 18 patients were in remission, 33 showed mild, 30 moderate and 8 severe disease activity. Nine patients did not require any treatment; however, 65 patients were on 5-ASA, 39 on corticosteroid, 37 on thiopurine, 15 on infliximab and one on cyclosporine-A therapy. Thirty-nine patients were also treated with topically administered drugs. Mean CRP was $21.7 \pm 41.4$ mg/l, and faecal calprotectin and MMP-9 were also elevated ($632.8 \pm 874.8$ μg/g; $6.8 \pm 8.3$ ng/l) in the enrolled patients. Strong correlation was shown between the inflammatory markers and the assessed endoscopic scores (see below). Serum and faecal parameters of enrolled patients are presented in Table 3. The mean duration of patient follow-up was 20.7 months (range, 0.3–54.5 months). During the follow-up period, 21 patients required hospitalisation due to relapse (mean: 0.7 times; range 0–15 times) and in 18 cases because of other causes not related to UC (mean: 0.2

Table 2 Characteristics of enrolled patients

| Parameter                          | Value       |
|------------------------------------|-------------|
| No. of enrolled patients           | 104         |
| Male/female                        | 46/58       |
| Mean age (±SD, years)              | 43.9 (±14.7) |
| Mean age at diagnosis(±SD, years)  | 35.3 (±14.7) |
| Mean disease duration (±SD, years) | 8.3 ± 0.7   |
| Localisation of disease:           |             |
| Rectum                             | 23          |
| Sigmoid colon                      | 28          |
| Descending colon                   | 18          |
| Transverse colon                   | 12          |
| Ascending colon (±cecum)           | 23          |
| Disease activity                   |             |
| The mean of eMayo Score (mean ± SD, points) | 1.8 ± 1.2 |
| The mean of UCEIS (mean ± SD, points) | 6.3 ± 2.4  |
| The mean of panMayo Score (mean ± SD, points) | 14.2 ± 13.7 |
| Therapy:                           |             |
| 5-ASA                              | 65          |
| Corticosteroids                    | 39          |
| Thiopurines                        | 37          |
| Cyclosporine-A                     | 1           |
| Biological therapy: infliximab     | 15          |
| Topical                            | 39          |
| No treatment                       | 9           |

Table 3 Laboratory markers of enrolled patients

| Laboratory parameter | value       |
|----------------------|-------------|
| CRP (mean ± SD, mg/l)| 21.7 ± 41.4 |
| WBC (mean ± SD, G/l) | 7.8 ± 3.3   |
| Thrombocytes (mean ± SD, G/l) | 299.3 ± 99.9 |
| Haematocrit (mean ± SD, %) | 39.1 ± 5.1   |
| Haemoglobin (mean ± SD, g/l) | 130.6 ± 19.7 |
| Serum iron (mean ± SD, umol/l) | 12.3 ± 6.2   |
| Faecal calprotectin (mean ± SD, μg/g) | 632.8 ± 874.8 |
| Faecal MMP-9 (mean ± SD, ng/ml) | 68 ± 8.3    |
Comparison between indices

PanMayo score

The mean panMayo Score was 14.2 ± 13.7 points (range, 0–45 points). Inactive and active disease was demonstrated in 45 and 59 patients. The ROC analysis revealed an AUC of 0.884 (p < 0.001, 95% CI: 0.822–0.946). The sensitivity and specificity of panMayo in the establishment of disease activity was 100% and 69.2%.

This new score correlated significantly with disease extent (R = 0.3; p < 0.001) and partial Mayo Score (R = 0.7; p < 0.001). In addition, it showed a strong association with the histological assessment, namely Riley Score (R = 0.7; p < 0.001). We found panMayo Score to correlate with CRP (R = 0.5; p < 0.001), faecal MMP-9 (R = 0.5; p < 0.001), calprotectin (R = 0.35; p < 0.004), leucocyte (R = 0.3; p = 0.009), thrombocyte levels (R = 0.3; p = 0.003), and also with haemoglobin (R = -0.3; p = 0.001), haematocrit (R = -0.3; p = 0.004) and serum iron level (R = -0.5; p < 0.001) (Table 4). Significant association was shown between panMayo Score and hospitalisation rate (p = 0.043). After reviewing our data, we observed a relationship between panMayo Score as a linear variable and colectomies, although this could not be substantiated by statistical analysis (p = 0.6) (Fig. 1).

Endoscopic Mayo score

The mean eMayo Score was 1.8 ± 1.2 points (range, 0–3 points) in this study. Inactive and active disease was presented in 41 and 63 patients. eMayo Subscore showed high correlation with partial Mayo Score (R = 0.7; p < 0.001), Riley Score (R = 0.7; p < 0.001), CRP (R = 0.3; p = 0.004) and faecal MMP-9 (R = 0.5; p < 0.001), but also with faecal calprotectin (R = 0.4; p = 0.019), serum iron (R = -0.4; p = 0.004) and leucocyte level (R = 0.3; p = 0.007), haemoglobin (R = -0.3; p = 0.02) and haematocrit (R = -0.2; p = 0.02). We did not find association with disease extent (R = -0.04; p = 0.7) and thrombocytes (R = 0.2; p = 0.1) (Table 4). We found that eMayo is strongly connected with hospitalisation (p = 0.028) and colectomies (p = 0.008).

UCEIS

The mean UCEIS was 6.3 ± 2.4 points (range, 3–11 points). Inactive and active disease was shown in 41 and 63 patients. UCEIS correlated with partial Mayo Score (R = 0.7; p < 0.001), Riley Score (R = 0.7; p < 0.001), faecal MMP-9 (R = 0.5; p < 0.001), faecal calprotectin (R = 0.35; p = 0.043), CRP (R = 0.5; p < 0.001), leucocyte count (R = 0.3; p = 0.009), thrombocytes (R = 0.3; p = 0.003), haemoglobin (R = -0.3; p = 0.001), haematocrit (R = -0.3; p = 0.004), and serum iron level (R = -0.5; p < 0.001), but not with disease extent (R = -0.04; p = 0.7) (Table 4). UCEIS showed a significant connection with hospitalisations (p < 0.001) and colectomies (p < 0.001).

Correlation between indices

The panMayo Score correlated with eMayo (R = 0.853; p < 0.001) and UCEIS (R = 0.783; p < 0.001), and eMayo with UCEIS as well (R = 0.932; p < 0.001).

Discussion

In this study we assessed the diagnostic value of a new, modified endoscopic score. The panMayo Score revealed a significant correlation with laboratory parameters of inflammation and anaemia, disease extension and histological activity of UC. Furthermore, despite the modification it showed 100% sensitivity with the “mother” endoscopic disease activity eMayo Score.

Use of endoscopic scoring systems is recommended for the evaluation of the prognosis and efficacy of treatment. The ideal score would be easy to use, reproducible, reliable, responsive to changes and validated [14]. Several endoscopic scoring systems are available for the assessment of disease severity, although only UCEIS and ulcerative colitis colonoscopy index of severity (UCCIS) [15] received formal validation [16]. UCEIS with a total score of 3 to 11 (or modified: 0–8) points was created based on vascular pattern, bleeding, and erosion/ulceration [17]. The study of Corte et al. showed that UCEIS has a predictive value also, because it was associated with the outcome of acute severe UC [18]. The Mayo Score takes into account four variables; these are stool frequency, rectal bleeding, the physician’s global assessments and findings at endoscopy, namely the eMayo Score [17]. eMayo Score differentiates between endoscopically inactive, mild, moderate and severe disease. Every endoscopic score indicates the severity of inflammation, but not the extent of the
inflamed colon which is a critically important detail for the optimisation of the therapy [10, 19]. The panMayo Score describes the bowel mucosa similarly to eMayo, but it also includes disease extent. We found strong correlation between panMayo Score and UCEIS, and eMayo as well. In addition, panMayo showed significant association with serum and faecal inflammatory markers. Therefore, this new score can correctly demonstrate the activity of UC and revealed unique correlation with disease extent hereby indicating one of the significant factors of disease outcome. Patients with initial diagnosis of pancolitis tend to present with more frequent complications, extraintestinal manifestations, occurrence of colorectal cancer is higher among them, and they are more likely to require immunosuppressive or surgical therapy [20, 21].

Despite general principles that treatment decisions should be based on disease activity, pattern of the disease and its distribution, colonoscopy and its evaluation may also play a crucial role in the therapy [19]. The panMayo Score correlates with inflammatory markers, and with parameters of anaemia as well, that is the most common systemic complication or extraintestinal manifestation of UC [22]. Anaemia is a disease activity related condition (triggered by blood loss and inflammation) in UC, moreover sometimes it is the only sign of an ongoing inflammation [23]. Furthermore, in another study, anaemia was shown to be more common in patients with UC requiring immunosuppressive therapy [24]. Nevertheless, anaemia in a population-based IBD cohort was associated with the extent of UC [25] and in our previous study the need for blood transfusion was a significant predictor of a subsequent colectomy [26]. It should be noted that disease location and extension has a great influence on the prognosis: approximately one third of patients with extensive UC will require colectomy during the disease course [27]. Nine of our patients had colectomy during the follow-up period. The panMayo Score as well as eMayo Score, and UCEIS have a predictable value on outcome in our study (Fig. 1).

To improve the accuracy of the endoscopic Mayo score, another work group had also created an alteration on the eMayo Score. The Modified Endoscopic Mayo Score which, similarly to the panMayo Score, considers the distribution of UC, showed good correlation with clinical, biological and histological activity of UC. However, calculation seems to be slightly difficult: after counting Modified Score and Extended Modified Score, the Modified Mayo Endoscopic Score [28] was obtained by dividing the Extended Modified Score with the number of segments with active inflammation.
The limitation of panMayo Score is that it can be calculated after a total colonoscopy which is not recommended in extraordinarily severe cases. The use of IC makes the calculation of panMayo Score slightly complicated, however; it is necessary for the clear differentiation of active from inactive disease. On the other hand, the general problem with every scoring system is that the reproducibility of endoscopic scores remains suboptimal as a study from Italy reported; assessment depends on the expertise of the gastroenterologist [8]. The key of consistent evaluation is suggested to be a standardized system of description and endoscopists’ training [29].

Summarizing our data, panMayo Score is a new, disease extent-related endoscopic score (created with a slight modification of eMayo) for the assessment of disease activity in UC, that showed excellent sensitivity and a good correlation with the parameters of inflammation and anaemia, Riley, eMayo Score and UCEIS. Nevertheless, it can be more favourable than the above mentioned endoscopic scores because panMayo Score gives additional information about disease location besides disease activity with a strong correlation with laboratory parameters of inflammation and anaemia.

Conclusion
There are many indices available to assess the endoscopic activity of ulcerative colitis, but all of them have some limitations. The panMayo Score provides additional information about the extent of the disease besides the disease activity.

Abbreviations
CRP: C-reactive protein; eMayo: Endoscopic Mayo Subscore; MMP-9: Matrix-metalloproteinase-9; panMayo: Pancolonic Modified Mayo Score; UC: Ulcerative colitis; UCEIS: Ulcerative Colitis Endoscopic Index of Severity; WBC: White blood cell

Acknowledgements
This paper was supported by the János Bolyai Research Scholarship of the Hungarian Academy of Sciences (BO/00632/14/5) and OTKA PD 105948 (Pt: Dr. Kádina Farkas).

Availability of data and materials
The datasets generated during and/or analysed during the current study are not publicly available due to respect participants’ rights to privacy and to protect their identity but are available from the corresponding author on reasonable request.

Authors’ contributions
Study design, supervision of patient selection, data collection and manuscript preparation: AB, BF, AF, TM; statistical analysis, tables and manuscript preparation: MSz, AB, BF, TM; patient selection, data collection: AB, BF, FN, AF, MR, ZSz, ZG, RB, AM, TM; performing colonoscopy: TM, ZSz; FN; histopathology: LT; supervision of the patient selection and manuscript preparation: TM. All authors read and approved the final manuscript.

Ethics approval and consent to participate
The study was approved by the Regional and Institutional Human Medical Biological Research Ethics Committee of the University of Szeged (Q26/2015-SZTE). All study participants, or their legal guardian, provided written informed consent prior to study enrollment.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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Received: 13 September 2016 Accepted: 12 December 2017
Published online: 08 January 2018

References
1. Walsh A, Palmer R, Travis S. Mucosal healing as a target of therapy for colonic inflammatory bowel disease and methods to score disease activity. Gastrointest Endosc Clin N Am. 2014;24:367–78.
2. Cooney RM, Warren BF, Altman DG, Abrue MT, Travis SPL. Outcome measurement in clinical trials for ulcerative colitis: towards standardisation. Trials. 2007;8:–17.
3. Truelove SC, Wtlts LJ. Cortisone in ulcerative colitis; final report on a therapeutic trial. BMJ. 1955;2:1041–8.
4. Travis SPL, et al. Reliability and initial validation of the ulcerative colitis endoscopic index of severity. Gastroenterology. 2013;145:987–95.
5. Schroeder KW, Tremaine WI, Estrup DM. Coated oral 5-aminosalicylic acid therapy for mildly to moderately active ulcerative colitis. A randomized study. N Engl J Med. 1987;317:1625–9.
6. Neurath MF, Travis SPL. Mucosal healing in inflammatory bowel diseases: a systematic review. Gut. 2012;61:1619–30.
7. Mosli MH, et al. Histologic evaluation of ulcerative colitis: a systematic review of disease activity indices. Inflamm Bowel Dis. 2014;20:564–75.
8. Riley SA, Mani V, Goodman MJ, Dutt S, Herd ME. Microscopic activity in ulcerative colitis: what does it mean? Gut. 1991;32:174–8.
9. Kato J, Kuriyama M, Hiraoka S, Yamamoto K. Is sigmoidoscopy sufficient for evaluating inflammatory status of ulcerative colitis patients? J Gastroenterol Hepatol. 2011;26:683–7.
10. D’Hanens G, et al. A review of activity indices and efficacy end points for clinical trials of medical therapy in adults with ulcerative colitis. Gastroenterology. 2007;132:763–86.
11. Stange EF, et al. European evidence-based consensus on the diagnosis and management of ulcerative colitis: definitions and diagnosis. J Crohns Colitis. 2008;2:1–23.
12. Satans J, Silverberg MS, Vermeire S, Colombel JF. The Montreal classification of inflammatory bowel disease: controversies, consensus, and implications. Gut. 2000;45:749–53.
13. Annahzad A, et al. Fecal MPP-9: a new noninvasive differential diagnostic and activity marker in ulcerative colitis. Inflamm Bowel Dis. 2013;19:316–20.
14. Tontini GE, Bispchofs R, Neumann H. Endoscopic scoring systems for inflammatory bowel disease: pros and cons. Expert Rev Gastroenterol Hepatol. 2014;8:543–54.
15. Samuel S, et al. Validation of the ulcerative colitis Colonoscopy index of severity and its correlation with disease activity measures. Clin Gastroenterol Hepatol. 2013;11:149–54.e1.
16. Annese V, et al. European evidence based consensus for endoscopy in inflammatory bowel disease. J Crohns Colitis. 2013;7(12):1982–1018.
17. Samana MA, et al. A systematic review of the measurement of endoscopic healing in ulcerative colitis clinical trials: recommendations and implications for future research. Inflamm Bowel Dis. 2014;20:1465–71.
18. Corte CJ, et al. Association between the ulcerative colitis endoscopic index of severity (UCEIS) and outcomes in acute severe ulcerative colitis. J Crohns Colitis. 2015. https://doi.org/10.1093/ecco-jcc/jjv047.
19. Dgnn A, et al. Second European evidence-based consensus on the diagnosis and management of ulcerative colitis part 2: current management. J Crohns Colitis. 2012;6(10):991–1030.
20. Roda G, et al. Systematic review with meta-analysis: proximal disease extension in limited ulcerative colitis. Aliment Pharmacol Ther. 2017; https://doi.org/10.1111/apt.14063.

21. Monstad I, Hovde Ø, Solberg KC, Mourn A. B. Clinical course and prognosis in ulcerative colitis: results from population-based and observational studies. Ann Gastroenterol Q Publ Hell Soc Gastroenterol. 2014;27:95–104.

22. Gasche C. Anemia in IBD: the overlooked villain. Inflamm Bowel Dis. 2002;6:142–50.

23. Dignass AU, et al. European consensus on the diagnosis and management of iron deficiency and anaemia in inflammatory bowel diseases. J Crohns Colitis. 2015;9:211–22.

24. Stallmach A, et al. Parameters of a severe disease course in ulcerative colitis. World J Gastroenterol WJG. 2014;20:12574–80.

25. Sjoberg D, et al. Anemia in a population-based IBD cohort (ICURE): still high prevalence after 1 year, especially among pediatric patients. Inflamm Bowel Dis. 2014;20:2266–70.

26. Farkas K, et al. Risk factors for colectomy in ulcerative colitis patients with severe flare-up: some of the few things comparable between the US and Central Europe. Inflamm Bowel Dis. 2011;17:E11–2.

27. Faubion WAJ, Loftus EVJ, Harmsen WS, Zinsmeister AR, Sandborn WJ. The natural history of corticosteroid therapy for inflammatory bowel disease: a population-based study. Gastroenterology. 2001;121:255–60.

28. Lobaton Omega T, Bessisso T, De Hertogh G, Van Assche G, Vermeire S, Rutgeerts P, Bisschops R, Bitton A, Aff W, Ferrante M. The modified Mayo endoscopic score (MMES): a new score for the assessment of extent and severity of endoscopic activity in ulcerative colitis (UC) patients. J Crohns Colitis. 2015;9:837–45.

29. Daperno M, Comberlato M, Bossa F, Brancone L, Bonanomi AG, Cassinotti A, Cosentino R, Lombardi G, Mangiarotti R, Papa A, Pica R, Rizzello F, D’Inca R, Orlando A. Inter-observer agreement in endoscopic scoring systems: preliminary report of an ongoing study from the Italian Group for Inflammatory Bowel Disease (IG-IBD). Dig Liver Dis. 2014;46:629–73.