Mucosal healing and deep remission: what does it mean?

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Mucosal healing and Deep Remission: What does it mean?

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Key words: Inflammatory bowel disease, mucosal healing, deep remission, treatment targets, clinical activity
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Competing Interest

**GR:** has received in the last 2 years consultant fees from Abbott Switzerland and Abbott International, Tillotts International, FALK Germany, Essex/MSD Switzerland, Novartis, Roche, and Vifor Switzerland; GR has received speaker’s honoraria from Abbott, FALK, MSD, Phadia, Tillotts, UCB, and Vifor; GR has received educational grants and research grants from Abbott, Ardeypharm, Essex/MSD, FALK, Flamentera, Novartis, Tillotts, UCB and Zeller.

**SV:** has received in the last 2 years consultant fees from Abbvie, MSD, and UCB, and Tillotts. SV has received speaker’s honoraria from Abbott, MSD, Tillotts, UCB, and Vifor; SV has received educational grants and research grants from Abbot/Abbvie, Essex/MSD and UCB.

**AS:** has received in the last 2 years consultant fees from Abbvie, MSD, UCB, and Tillotts. AS has received speaker’s honoraria from Abbvie; AS has received educational grants and research grants from Essex/MSD, UCB, and Tillotts.

**LPL:** has received in the last 2 years consultant and lecture fees from Abbott/Abbvie, MSD-Hungary, and Ferring
Abbreviations: CD: Crohn’s disease; IBD: inflammatory bowel disease; UC: ulcerative colitis

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Abstract

The use of specific terms under different meanings and varying definitions has always been a source of confusion in science. When we point our efforts towards an evidence based medicine for inflammatory bowel diseases (IBD) the same is true: Terms such as “mucosal healing” or “deep remission” as endpoints in clinical trials or treatment goals in daily patient care may contribute to misconceptions if meanings change over time or definitions are altered.

It appears to be useful to first have a look at the development of terms and their definitions, to assess their intrinsic and context-independent problems and then to analyze the different relevance in present-day clinical studies and trials. The purpose of such an attempt would be to gain clearer insights into the true impact of the clinical findings behind the terms. It may also lead to a better defined use of those terms for future studies.

The terms „mucosal healing“ and „deep remission“ have been introduced in recent years as new therapeutic targets in the treatment of IBD patients. Several clinical trials, cohort studies or inception cohorts provided data that the long term disease course is better, when mucosal healing is achieved. However, it is still unclear whether continued or increased therapeutic measures will aid or improve mucosal healing for patients in clinical remission. Clinical trials are under way to answer this question. Attention should be paid to clearly address what levels of IBD activity are looked at. In the present review article authors aim to summarize the current evidence available on mucosal healing and deep remission and try to highlight their value and position in the everyday decision making for gastroenterologists.
**Key words** Inflammatory bowel disease, mucosal healing, deep remission, treatment targets, clinical activity

**Core tip:**

“Mucosal healing” and “deep remission” have been discussed heavily as “new” treatment goals in IBD patients in recent years. This was based on evidence that the long term disease behaviour appears to be better, when mucosal healing is achieved. Unfortunately, a definite proof that therapy escalation for patients in clinical remission not achieving mucosal healing will be beneficial is still lacking. Clinical trials are under way to answer this question. At the moment it appears to be helpful to summarize the current evidence available on mucosal healing and deep remission to support the everyday decision making for gastroenterologists.

98 words

**Introduction**

Assessing the activity of inflammatory bowel disease (IBD) is important for our daily practice treating patients with these chronic inflammatory diseases. The assessment of disease activity will guide our therapeutic decision and our choice of medication. Furthermore it is most important for clinical investigations of new treatment options and new drugs. The reduction of disease activity remains the most important endpoint in clinical trials.
However, the discussion on which parameters are most useful for this purpose is still ongoing and unresolved.

Assessment of activity of inflammatory bowel disease (IBD) can be performed on different levels such as clinical activity, biochemical activity (eg. by measuring CRP or fecal calprotectin), endoscopy, and histology. Clinical remission in a given IBD patient does not necessarily imply biochemical, endoscopic, or histologic remission. To evaluate biochemical, endoscopic, and histologic activity, an increasing degree of invasive measures (blood sample, endoscopy, biopsies) is required. Assessing activity in IBD has thereby analogies to the iceberg phenomenon where the clinical assessment on the surface may show clinical remission, but inflammatory activity may still be present on biochemical, endoscopic, and histologic level (Figure 1).
Histological remission as initial definition of mucosal healing

One of the first scientists and clinicians that used the term “healing” or “mucosal healing” within the field of IBD was Burton I. Korelitz, past chief of the Division of Gastroenterology at Lenox Hill Hospital in New York [1]. However, he used this term exclusively with respect to histological changes of the mucosa [1]. So when the term “mucosal healing” was introduced into IBD clinic it meant the absence of histological alterations of the mucosa. Korelitz was well aware that healing of IBD is not regarded to be possible as both Crohn’s disease (CD) and ulcerative colitis (UC) are regarded to be chronic diseases without spontaneous healing [2]. There may be an absence of symptoms and flares over years but mucosal inflammation may re-occur after remission for years or even decades (Figure 1).

Histological healing is difficult to determine especially in Crohn’s disease as the inflammation may be patchy and a biopsy could miss an inflammatory infiltrate only a few millimeters away [3]. Similarly, in UC the histological evaluation of a biopsy may be misleading [4]. Histological alterations may be absent from the rectum and sigmoid due to effective topical therapy despite the presence of inflammation further proximal in the colon that may not be obvious to the endoscopist [4, 5]. Histological healing would mean that we have to be sure that there had been an inflammatory infiltrate at a specific localization that completely disappeared upon therapy (or spontaneously). As is obvious this is hard or even impossible to prove as this would require frequent endoscopies with many biopsy samples and a labeling of former
biopsy locations. Due to the impracticability of this approach the overall acceptance of the concept of “histological healing” was very limited. [5]. Of note, newer techniques such as endomicroscopy suffer from the same shortcomings.

**Endoscopic remission as a new concept for mucosal healing.**

In contrast to the initial concept of “mucosal healing” as a “disappearance of inflammatory infiltrate” [2] recent original manuscripts and reviews on the topic have used the term under different meanings. The “newer” meanings of “mucosal healing” have been summarized again by Korelitz in a critical review [2]. One of the “newer meanings” of mucosal healing would be the absence of inflammation (“healed mucosa”) to the eye of the endoscopist, a definition that now has been applied in many clinical trials [6-16].

There is an obvious problem with this definition. One must assume the location of endoscopically normal mucosa has previously been inflamed [2]. Certainly this is easier to assess with endoscopy rather than histology as the area of evaluation is larger and small local differences and a patchy pattern would play a less important role. Nevertheless it requires that two endoscopical examinations are compared.

The definition also ignores that in endoscopically normal appearing mucosa there still may be histological inflammation. Another problem of this definition of course is that the inter-observer reproducibility of endoscopical IBD scores usually is very poor [17] and depends on the experience of the endoscopist [18] regardless of the technique used [19, 20] (it may be discussed whether a kappa between 0.7 and 0.8 is satisfying). Usually endoscopic
findings are assessed on fixed point scales or described by dichotomous variables (present/absent) \cite{18, 21}. However, as outlined by de Lange and colleagues “endoscopic features of mucosal inflammation are continuous variables” for which dichotomous decisions are artificial and always require individual decisions \cite{18}. The question arises how to interpret endoscopical findings indicating a clearly improved appearance of the mucosa in endoscopy with some or few remaining scattered erosions. A further important question arises with respect to endoscopical findings that cannot be interpreted as present inflammation but as residuals of former inflammation and a lack of complete normalization of the mucosa. Such findings would be pseudopolyps in an otherwise normal-appearing colon.

**Biochemical (fecal markers) remission as mucosal healing**

Fecal markers such as calprotectin or lactoferrin correlate very well with the degree and extent of infiltration of the mucosa by leukocytes. A good correlation between fecal calprotectin and the Crohn’s Disease Endoscopic Index of Severity (CDEIS) was reported in several studies \cite{22, 23}. There is also a good correlation of fecal calprotectin with the Simple Endoscopic Score for Crohn’s disease (SES-CD) which itself has a strong correlation with the CDEIS (correlation coefficient $r = 0.920$) and an excellent inter-observer reliability ($\kappa$ coefficients $0.791 - 1.000$) \cite{24}. In ulcerative colitis calprotectin correlates well with disease activity as determined by histology and endoscopy \cite{25, 26}. It is a familiar experience to endoscopists that the mucosa may appear completely normal (healed) in patients that still have a markedly elevated
fecal calprotectin. This would be an endoscopic remission but not biochemical remission, most likely reflecting a lack of histological remission with neutrophils still being present in the mucosal wall. It has been well established that calprotectin better correlates with histological findings (at least in UC) as compared to serum parameters or endoscopy [27-29].

**Mucosal healing and deep remission: The confused clinician**

Surprisingly, some recent trials have reported a higher relative amount of patients with mucosal healing compared to the percentage of patients with clinical remission, especially in UC [30]. In those trials usually the endoscopist defined whether mucosal healing was present. How can this be explained? One reason could be that those patients had concomitant irritable bowel syndrome that was responsible for their complaints but no relevant remaining inflammation (“IBS superimposed on IBD”). The argument is straightforward and logical but it probably does not explain all cases. Firstly, little or no information is available on the histological remission in those patients. Histological remission – if evaluated by biopsies – again may be patchy and the evaluated biopsies may not be representative. Damage to deeper layers of the mucosa may have occurred that are not visible to the endoscopist’s eye. Therefore is has to be challenged whether healed mucosa to the eye of the endoscopist is indeed the “most satisfying objective confirmation to support the clinical response” as outlined by Korelitz [2]. As he states the endoscopic healing “might be satisfactory for comparison in time for response to therapy in an individual case, but not for
mucosal healing as an entity and certainly not to be used as an index of response to therapy in trials.” [2]

To minimize the subjective component many clinical trials now apply the principle of a “central reader”. Not only does this make trials more complicated, more expensive and more time consuming. It substitutes the problem of a bias introduced by many subjective evaluations of the mucosal response to a bias introduced by one subjective interpretation of findings. The intra-observer agreement for many endoscopic scores is not satisfactory. It may well be argued that the subjective criteria used by a central reader may not be accepted by others and that there could be a reduction of bias by a “multi-subjective” view (as we assume is the case for multicenter trials as compared to monocentric studies). Of note, in a recent randomized-controlled trial in patients with UC the conclusion was significantly changed after blinded central review of endoscopic images, suggesting that central reading of endoscopy may be necessary for regulatory purposes [31]. However, the question about the best method of objective endoscopic assessment is far from being answered.

Korelitz suggested that histological healing should be the “minimal criterion for mucosal healing and preferably this information should be derived from multiple biopsy sites of previous inflammation” [2]. However, this would implicate that the evaluation of inflammation by a pathologist is objective. There have been studies on the inter-observer and intra-observer agreement of pathology findings [32]. Those results are not very encouraging. When a number of established criteria were used (excess of histiocytes in combination with a villous or irregular aspect of the mucosal surface and
granulomas) experienced pathologists could correctly classify 70% of CD patients and 75% of UC patients [32]. Especially in mild disease, there is still dispute as to whether the presence of a “physiological (minor) inflammation“ should be regarded as manifestation of IBD or not. Clinically unaffected siblings of IBD patients may show mild histological inflammation and increased cellular activation markers [33]. Cell counting will not solve the problem. The request for a “central pathology reader” also is not helpful as the same dilemma as for the central endoscopy reader will occur. Moreover, different pathologists have suggested different criteria to evaluate the presence or absence of “un-normal” inflammation (for an overview see [3, 34-37]. There is no agreement on that. Geboes for example suggested that the presence of neutrophils in the intestinal epithelium is an important discriminator for the presence or absence of inflammation. He therefore suggested that a combination of endoscopy and histology should be used to evaluate the presence of inflammation in IBD patients to finally judge whether mucosal healing has been achieved (see above).

**Mucosal healing and deep remission: The confused scientist**

CD and UC are regarded to be chronic diseases that never disappear. The concept of a healing of a part of the body affected by such a disease subsequently is surprising for scientists working on the elucidation of the pathophysiology of IBD.

However, there is another aspect that is disturbing. There have been reports that even in macroscopically and microscopically normal appearing mucosa
specific changes can be found that are characteristic for inflammation or at least changes that could be associated with the pathophysiology [38-45]. Changes of the microbiota in the lumen of the gut have been described in IBD patients despite the absence of detectable inflammation [46-51]. Could a “complete deep remission” be possible without normalization of the intestinal microbiome? The mucus layer of the mucosa may be changed also in normal appearing mucosa in endoscopy [52-56]. The normal fixation procedure of biopsies and the subsequent H&E staining does not allow evaluation of the mucus layer as it is destroyed during this procedure. A reduced thickness of the mucus layer in UC in remission has been described [54, 56, 57] as well as a reduced secretion of mucin [52, 53, 58-60] or defensins [61-64]. The question arises whether the mucosa can be termed as “normal” or “healed” if those changes are still present.

Epithelial cells may have an impaired barrier function despite a lack of inflammatory signs. Cytokine expression and cytokine secretion by immune cells may still be significantly increased despite a normal appearing histology. A normalization of those changes has been termed biochemical healing [65-68]. There are no data available with respect to the predictive value of “biochemical healing” and whether this would correlate to a more favorable disease outcome.

The confused scientist, however, is able to imagine a further level of “healing”. In macroscopically normal appearing mucosa with microscopically normal appearing cells that display normal cytokine expression and secretion levels, epigenetic changes may still be present that may trigger pathological responses upon minor stimuli [69-76]. Can a persistence of
epigenetic changes in otherwise normal mucosa be termed “mucosal healing”? Or do we have to achieve “epigenetic healing” to finally achieve the best outcome possible for our patients? These questions will have to be answered in the future. Currently we are just at the start of investigations into these aspects with the first interesting pieces of the puzzle being put together.

**Mucosal healing and deep remission: The confused “trialist”**

As mentioned above the terms “mucosal healing” and “deep remission” have been used in a number of trials with quite different meanings and definitions. The key confounder is the lack of unequivocal definition(s). Therefore, results and data from those trials with respect to mucosal healing cannot easily be compared. Nevertheless, this is done frequently. In most cases endoscopical investigation is used for the evaluation of “mucosal healing”. One crucial point is whether “mucosal healing” was defined simply as the absence of ulcers when ulcers had been seen previously or whether the absence of ulcerations and ulcers was investigated exactly at a place where those alterations had been found before.

The above is reflected in the way different trials have been reported. In the ACCENT 1 endoscopic sub-study the CDEIS was used for scoring and the complete absence of mucosal ulcerations that were observed at baseline was evaluated [77]. In the SONIC study in contrast no clearly defined score was used. Mucosal healing was defined as “complete absence of mucosal ulceration in the colon and terminal ileum” [78]. In the “Top-down versus step
up” study by Gert D’Haens and coworkers SES-CD was used for the evaluation of mucosal healing which was a secondary endpoint \[79, 80\]. Mucosal healing was defined as “absence of ulcers”. In the MUSIC trials again the CDEIS was applied. The definition of mucosal healing was “absence of ulcers and endoscopic remission defined as CDEIS < 6”. In the EXTEND study applying again SES-CD mucosal healing was seen as “absence of mucosal ulceration” \[81\]. As is obvious from those definitions, the question arises whether a few remaining aphthous lesions in a patient with severe and deep ulcers at the beginning of therapy also may be termed mucosal healing.

For UC the IOIBD attempted a consensus for mucosal healing in 2007: “absence of friability, blood, erosions and ulcers in all visualized segments of the gut mucosa”. According to the IOIBD experts the presence of an abnormal vascular pattern is still compatible with mucosal healing or “normal mucosa”. However, also in UC the definitions applied varied widely: In the ACT1 study mucosal healing was a secondary endpoint \[82, 83\]. The Mayo endoscopic subscore was used and mucosal healing was defined as “absolute subscore for endoscopy of 0 or 1” \[82, 83\]. The same definition was used for ULTRA 2 \[84\].

In studies on the outcome of therapy with 5-aminosalicylic acid the definition of mucosal healing largely defined the number of patients achieving this endpoint (Table 1). As an example, Vecchi et al. compared mesalazine 4g orally vs. 2+2g orally and enema in 2001 in patients with a clinical activity index (CAI) of 4-12 and used an endoscopic Rachmilewitz index <4 as definition of mucosal healing leading to 58% vs. 71% of patients
achieving this endpoint [85]. In 2002 Malchow compared Mesalazine 4g enema vs. 1g foam preparation in patients with a CAI>4 for 4 weeks and applied an endoscopic Rachmilewitz index < 2 as definition of mucosal healing leading to rates of 38% vs. 37% [86]. As one would expect, the different definitions used cause huge variation in defined endoscopic mucosal healing rates in patients with UC, which makes the comparison of efficacy of different drugs or formulations extremely difficult.

Table 1:

Association between the definitions of remission and mucosal healing and actual healing rates in patients with UC treated with mesalazine (MH). Abbreviations: Mc = multicenter, db = double-blind, RCT = randomized controlled trial.

<table>
<thead>
<tr>
<th>Author</th>
<th>Design</th>
<th>Study</th>
<th>Timing of endoscopy</th>
<th>Endoscopic index</th>
<th>Def. of MH</th>
<th>No of pat. Achieving MH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vecchi (2001)</td>
<td>Mc, RCT</td>
<td>Mesalazine 4g orally vs. 2+2g orally and enema</td>
<td>6 weeks</td>
<td>Rachmilewitz</td>
<td>Rachmilewitz &lt;4</td>
<td>58% vs. 71%</td>
</tr>
<tr>
<td>Malchow (2002)</td>
<td>Mc, db, RCT</td>
<td>Mesalazine 4g enema vs. 1g foam</td>
<td>4 weeks</td>
<td>Rachmilewitz</td>
<td>Rachmilewitz &lt; 2</td>
<td>38% vs. 37%</td>
</tr>
<tr>
<td>Mansfield (2002)</td>
<td>Mc, db, RCT</td>
<td>Balsalazide 6.75g vs. sulfasal. 3g</td>
<td>8 weeks</td>
<td>4 point scale</td>
<td>Score of 0= normal mucosa</td>
<td>27% vs. 25%</td>
</tr>
<tr>
<td>Hanauer (2007) Ascend</td>
<td>Mc, db, RCT</td>
<td>Asacol 4.8g vs. 2.4g</td>
<td>6 weeks</td>
<td>Descriptive, no score</td>
<td>Normal endoscopic finding</td>
<td>25% vs. 20%</td>
</tr>
<tr>
<td>Kamm (2007) MMX</td>
<td>Mc, db, RCT</td>
<td>MMX mes. 4.8g vs. 2.4g vs. placebo</td>
<td>8 weeks</td>
<td>Mod. Sutherland index</td>
<td>Mod Sutherland index &lt; 1</td>
<td>77% vs. 69% vs. 46%</td>
</tr>
</tbody>
</table>
One of the problems in endoscopic UC scores is the application of varying criteria (see table 2)

<table>
<thead>
<tr>
<th>Erythema</th>
<th>Baro</th>
<th>Powell-T</th>
<th>Levine</th>
<th>Rachmilewitz</th>
<th>Modified</th>
<th>Mayo</th>
<th>Sutherland</th>
</tr>
</thead>
<tbody>
<tr>
<td>Erythema</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Edema</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Granularity</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vascular pattern</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Friability</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Erosions</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ulceration</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exudate</td>
<td></td>
<td></td>
<td>+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Remission</td>
<td>0</td>
<td>0-1</td>
<td>0-2</td>
<td>0-1</td>
<td>0-1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

The reasons for such different definitions and endpoints may only be speculated.

Unfortunately we lack an unequivocal definition; all of the scoring systems published so far have certain limitations, which have led to the introduction of several additional scoring systems. From a patient’s and physician’s perspective, however, the use of one single scoring system would be most desirable to enable valid comparisons among study outcomes.
What is the additive value of deep remission as compared to mucosal healing?

“Deep remission” is another term that has been discussed as a treatment target in recent years. The definition, however, is unfortunately not clearer than the one of mucosal healing. In the EXTEND study “deep remission” was defined as clinical remission (CDAI <150) and complete mucosal healing as defined according to CDEIS \cite{13}. It is worthwhile to look a bit closer at this definition. If a patient with CD achieves mucosal healing but still has increased CDAI (no clinical remission) this may be due to superimposed IBS symptoms or the fact that without the presence of inflammation there is some bowel damage such as a fibrotic stricture or an internal fistula which might contribute to increased bowel frequency. Subsequently the lack of clinical remission is important for the patient and his/her clinical management (e.g. surgery of the stricture) but not for the medical (anti-inflammatory) management of the disease. Thus, the term “deep remission” in the definition outlined above is not useful and does not provide more information than mucosal healing. In fact – it contributes to confusion of scientists, clinicians and “trialists”.

How can we improve?

There should be standards on the definition of mucosal healing for clinical studies. It needs to be discussed – and finally decided – whether endoscopic mucosal healing, histologic mucosal healing or a combination of both can be standardized. Once agreement on definitions has been achieved, a given patient could be assessed by a –hopefully- simple binary coded tool that is
oriented according to the TNM classification of oncology. A proposal for such a tool is illustrated in Table 3. The number “1” stands for “active”, “0” for “remission” and “x” for “not assessed”. Of note CD activity assessment would require, in contrast to UC, not only measuring clinical activity, biochemical, endoscopic and histologic activity, but also imaging modalities (presence of fistulas, strictures). This simple approach has the potential to reduce the amount of potentially confusing new definitions to describe different combinations of activities in IBD.

Other definitions of mucosal healing (such as “biological mucosal healing”, “epigenetic mucosal healing”, “mucus layer healing” or “microbiota mucosal healing”) require further studies and prospective trials. At this point they are purely investigational and should not be used in clinical trials.

What would happen if such an agreement cannot be achieved? Then it would not make sense to discuss mucosal healing as a treatment target for IBD any further as this would be a treatment target that lacks a definition and subsequently is blurry, vague and indistinct.

**Table 3**: proposal of the CBEHI classification to assess CD activity. Example: a CD patient with C0B0(CRP)E1H1I0 would have clinical and biochemical remission, but endoscopic and histologic activity.

<table>
<thead>
<tr>
<th>Activity level</th>
<th>Definition</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clinical activity</strong></td>
<td>Remission: CDAI &lt;150</td>
<td>C0</td>
</tr>
<tr>
<td></td>
<td>Active: CDAI ≥150</td>
<td>C1</td>
</tr>
<tr>
<td><strong>Biochemical activity</strong></td>
<td>CRP normal</td>
<td>B0(CRP)</td>
</tr>
<tr>
<td></td>
<td>Elevated CRP</td>
<td>B1(CRP)</td>
</tr>
<tr>
<td></td>
<td>Calprotectin &lt;200µg/g</td>
<td>Calprotectin ≥200µg/g</td>
</tr>
<tr>
<td>------------------------------</td>
<td>-----------------------</td>
<td>-----------------------</td>
</tr>
<tr>
<td><strong>Endoscopic activity</strong></td>
<td>Remission: SES-CD &lt;4</td>
<td>Active: SES-CD ≥4</td>
</tr>
<tr>
<td><strong>Histologic activity</strong></td>
<td>Inactive</td>
<td>Active</td>
</tr>
<tr>
<td><strong>Imaging</strong></td>
<td>Inactive: no fistulas, no stenoses</td>
<td>Active: presence of either fistula and/or stenosis</td>
</tr>
</tbody>
</table>
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