Ultrasound Elastography in Inflammatory Bowel Diseases: A Systematic Review of Accuracy Compared with Histopathological Assessment

Arianna Dal Buono, a, Arianna Dal Buono, a, b Francesco Faita, b Laurent Peyrin-Biroulet, c Silvio Danese, d Mariangela Allocca d

a IBD Center, Department of Gastroenterology, Humanitas Clinical and Research Center - IRCCS, Rozzano, Milan, Italy
b Italian National Research Council Institute of Clinical Physiology, Pisa, Italy
c Department of Gastroenterology and Inserm NGERE U1256, University Hospital of Nancy, University of Lorraine, Vandœuvre-lès-Nancy, France
d Gastroenterology and Endoscopy, IRCCS Ospedale San Raffaele, and University Vita-Salute San Raffaele, Milan, Italy

Corresponding author: Mariangela Allocca, MD, PhD, IBD Center, Department of Gastroenterology, Ospedale Vita-Salute San Raffaele, IRCCS, Milan, Italy. Tel.: +39026432069; E-mail: mariangela.allocca@gmail.com

Abstract

Background and Aims: Ultrasound elastography [USE] is an innovative, non-invasive, promptly available, ancillary technique that has been proposed in the evaluation of intestinal fibrosis as a monitorable biomarker, in terms of stiffness. The non-invasive estimate of fibrosis by USE appears appealing for dedicated physicians, in order to optimise the treatments for inflammatory bowel disease [IBD] patients [surgical vs non-surgical]. We aimed to systematically review literature evidence on ultrasound elastography in IBD patients.

Methods: For this qualitative systematic review, we searched PubMed, EMBASE, and Scopus to identify all studies, published until October 2021, investigating the application of USE in IBD patients compared with histopathological assessment.

Results: Overall, 12 papers published between 2011 and 2019 were included. A total of 275 IBD patients were included: 272 Crohn's disease [CD] [98.9%] and three ulcerative colitis [UC] [1.1%]. Seven [58.3%] and four [41.6%] studies investigated strain elastography [SE] and shear wave elastography [SWE], respectively; in one study [0.1%] both techniques were addressed. The histological evaluation was largely conducted on surgical specimens and in two studies endoscopic biopsies were also included. The histological assessment was semi-quantitative in all the included studies, except for two where the fibrosis was evaluated only qualitatively. In 10/12 publications USE could accurately distinguish inflammation from fibrosis in the examined bowel tracts.

Conclusions: From the preliminary available data, an overall moderate-to-good accuracy of USE in detecting histological fibrosis [10/12 studies] was found. Point shear wave elastography has been shown to perform superiorly. Further studies are needed to confirm these evidences.

Key Words: Elastography; shear waves; bowel ultrasound; inflammation; inflammatory bowel disease

1. Introduction

Chronic inflammatory bowel diseases [IBDs] are relapsing-remitting and progressive conditions that lead to irreversible bowel damage. Especially the stricturing phenotype of Crohn's disease [CD] and late stages of ulcerative colitis [UC] are characterised by the development of fibrosis in the affected bowel tract. Fibrotic strictures have a multifactorial biological basis that involves the activation of mesenchymal cells that overproduce and deposit extracellular matrix. Soluble molecules such as cytokines and growth factors [i.e., transforming growth factors, tumour necrosis factor, interleukins] trigger this activation, with a subsequent remodelling of the tissue by matrix metalloproteinases and other fibrogenic enzymes.

The predominancy of fibrosis is believed to be less responsive to medical treatments and often requires a surgical intervention [i.e., resection, strictureplasty]. For this reason, distinguishing between IBD patients with a primarily inflammatory or a fibrotic disease has a relevant impact on clinical management.

In a recent systematic review, the sensitivity in detecting fibrosis of cross-sectional imaging techniques has been assessed at around 80% for both computed tomography [CT] and magnetic resonance imaging [MRI]. The main features that help the radiologists in distinguishing between inflammation and fibrosis are bowel wall thickness, mural contrast enhancement, mesenteric vascularity, and mesenteric fat stranding.

To date, neither a scoring system nor standardised criteria have been established for differentiating fibrosis at cross-sectional imaging, thus remaining an unsolved challenge for dedicated physicians.

Ultrasound elastography [USE] is an innovative, non-invasive, promptly available, ancillary technique that has been proposed in the evaluation of intestinal fibrosis as a monitorable imaging biomarker, in terms of stiffness.
As concerns technical aspects, USE assesses the elastic properties of soft tissues by acoustic or mechanical stimulation: the tissue response to the stress is processed and codified as an image with a scale of colours or as a quantitative measurement corresponding to the estimated stiffness value. The main types of USE are shear wave elastography [SWE] and strain elastography [SE]. The stimulus for the measured stress ranges from acoustic radiation force impulse imaging [ARFI] to mechanical or physiological palpation. In detail, point-SWE [pSWE] estimates a quantitative value of a specific point of the examined tissue, whereas two-dimensional SWE [2D-SWE] codifies a colour map that reflects the stiffness of a wider portion of the examined tissue. The application of USE has already been incorporated in the diagnostic algorithms of diseases of the liver, breast, pancreas, and thyroid, especially for neoplastic lesions.

Thanks to recent technological advancements, USE is implemented and usable in real time. However, there are no current international guidelines instructing on the applications of elastography in the field of IBD.

The role of USE in the management of IBD is currently under investigation, and its validation requires precise knowledge of the corresponding histological features. So far, data from literature on USE accuracy in the field of IBD are mostly derived from small cohorts and have never been comprehensively reviewed specifically and exclusively in comparison with histology as a reference standard. The purpose of our systematic review is to provide an exhaustive overview of the corresponding histological features. So far, data from literature on USE accuracy in the field of IBD are mostly derived from small cohorts and have never been comprehensively reviewed specifically and exclusively in comparison with histology as a reference standard. The purpose of our systematic review is to provide an exhaustive overview of the available data on USE in IBD patients.

2. Technique and principles of ultrasound elastography

Elastography evaluates the tissue elasticity, defined as the tendency of that tissue to resist deformations by an applied force, or to return to its original shape once the force is removed. Biologically, a stiff region displays less deformation compared with healthy surrounding tissue when the same stress stimulus is applied. The technologies currently used and commercially available in US machines are divided into two main types: strain [SE] and shear wave elastography [SWE] [Figure 1]. These types of elastography differ in the process used to measure tissue deformation in response to an applied force.

In detail, the applied force could be a mechanical internal [exploiting physiological periodic compression induced by circulatory and/or respiratory motion] or external [generated by hand through the US transducer that is gently pressed against explored tissues] pressure [Figure 1]. Alternatively, tissues can be stressed by imposing a low-frequency ARFI stimulus generated by the US device itself. In SE, the induced tissue displacement is traced between pairs of echo frames and then the strain is calculated from their gradient. Through the use of a colour map, the different strains are encoded within a two-dimensional image that can be instantly visualised together with the conventional B-mode US image. The SE is a semi-quantitative technique that cannot measure the elasticity of the examined tissue as an absolute value, since the absolute value of the applied stress is unknown.

In SWE, the dynamic stress induces shear waves that propagate perpendicular to the US beam. The speed of the generated shear waves is measured and returns quantitative estimates of the tissue elasticity. Technically, two SWE methods can be distinguished: the point-SWE [pSWE] and the 2-dimensional-SWE [2D-SWE]. In the p-SWE, the speed of the shear wave is measured in a single specific location [ROI]; the 2D-SWE produces a quantified colour map of the distribution of shear wave velocities in a wider region.

3. Methods

This work was conducted in accordance with the Cochrane Handbook and Preferred Reporting Items for Systematic Reviews and Meta-Analyses [PRISMA] recommendations for reporting systematic reviews.

3.1. Data sources and search strategy

We designed a comprehensive search strategy and searched PubMed/MEDLINE, Embase, and Scopus up to October 2021 to identify eligible studies. A hand-search of abstracts from the annual meetings of Digestive Disease Week, the American College of Gastroenterology, the European Crohn’s and Colitis Organisation, and the United European Gastroenterology Week, up to 2021, was also performed.

The search query employed both an exhaustive list of keywords and index terminology whenever possible. The following key words and corresponding Medical Subject

![Figure 1](Image)

**Figure 1.** Technique and principles of ultrasound elastography. A. The technologies currently used and commercially available in ultrasound [US] machines are divided into two main types: strain [SE] and shear wave elastography [SWE]. In SE, the induced tissue displacement is traced between pairs of echo frames, then the strain is calculated from their gradient. Technically, two SWE methods can be distinguished: the point-SWE [pSWE] and the 2-dimensional-SWE [2D-SWE]. In the p-SWE, the speed of shear wave is measured in a single specific location [ROI]; the 2D-SWE produces a quantified colour map of the distribution of shear wave velocities in a wider region. B. An example of SE applied to bowel wall is shown. SWE: shear wave elastography.
Heading/Entree terms were used: ‘elastography’, ‘ultrasound elastography’. The Medline search strategy was: [Ultrasound elastography] OR [elastography] OR [shear wave] OR [acoustic radiation force impulse imaging] OR [strain elastography] AND [inflammatory bowel disease] OR [Crohn's disease] OR [Colitis, Ulcerative': Mesh] OR [Crohn] AND [histology] OR [histopathological] OR [microscopic] OR [histopathology] OR [pathology] NOT animals. The full search strategy is available in the Supplementary material. No date or language filters were employed in the search. The literature search was performed and verified by two authors [FF, ADB].

3.2. Inclusion and exclusion criteria
The inclusion criteria were: a] studies investigating the application of USE in IBD patients; b] studies including a histopathological confirmation of the analysed bowel segment; c] studies on different USE modalities [i.e., SWE and SE]. No restriction on the type of study was applied. Full-text papers, conference abstracts, and case reports were included. Studies on paediatric populations [<16 years old] were excluded. All editorials, letters, or review articles were excluded. Animal studies were excluded as recommended in the Cochrane Handbook of Systematic Reviews of Interventions.14

3.3. Selection process, data extraction, and quality assessment
Two review authors [FF, ADB] independently screened the titles and abstracts yielded by the search. Full reports were obtained for all titles that appeared to meet the inclusion criteria or where there was any uncertainty. Disagreements were resolved through collegial discussion. The reasons for excluding trials were recorded. When there were multiple articles for a single study, the latest publication was used. The studies were reviewed for patients’ selection and features, technical aspects, USE, and histological assessment. When the USE assessment was done through classes based on the analysis of qualitative colour maps, it was considered semi-quantitative; when the USE measurements were reported as absolute values, it was considered a quantitative assessment. Finally, when the USE

Figure 2. Preferred Reporting Items for Systematic Reviews flow diagram.
measurements were not ordered into classes of severity, it was considered as a qualitative assessment.

The quality of the included studies was assessed with the Quality Assessment of Diagnostic Accuracy Studies [QUADAS-2] checklist.16 This tool includes four domains: patient selection, index test, reference standard, and flow and timing. The risk of bias is evaluated across all four domains, and the first three domains are also assessed in terms of concerns regarding applicability. The QUADAS-2 allows expression of an overall judgment as ‘low risk of bias’ or ‘low concern regarding applicability’ in case of assignment of ‘low’ to most/all domains relating to bias or applicability. If a study is judged ‘high’ or ‘unclear’ regarding one or more domains, then it may be judged ‘at risk of bias’ or as having ‘concerns regarding applicability’.

4. Results
The literature search revealed 177 publications: after excluding any duplicates, 158 were screened. After reviewing

| Reference, first author | Year | Study design | N | UC | CD | US elastography technique | US device | Results on accuracy of USE |
|-------------------------|------|--------------|---|----|----|----------------------------|-----------|---------------------------|
| Stidham RW17            | 2011 | Case series  | 7 | 7  | SE | Zonare Medical Systems [Z-1], Mountain View, CA | Fibrostenotic bowel was stiffer than the normal tissue resection margins in all subjects \[p = .0009\] |
| Havre RF18              | 2014 | Prospective  | 9 | 9  | SE | HV900 [Hitachi Medical Corporation, Tokyo, Japan] | Mean SR of 3.33 [SD, 5.21] in CD patients with increased tissue fibrosis [score ≥1] |
| Dillman JR19            | 2014 | Prospective  | 11| 3  | 8  | pSWE/2D-SWE | Acuson S3000 [Siemens Medical Solutions USA, Inc., Mountain View, CA] | High fibrosis vs low fibrosis score segments showed significantly greater mean shear wave speed \[p = 0.049\] |
| Baumgart DC20           | 2015 | Prospective  | 10| 10 | SE | Hitachi [Lübecke, Germany] | A higher collagen content was associated with RTE-assessed strain \[p <0.001\] and tensiometry-assessed strain \[p <0.001\] |
| Fraquelli M21           | 2015 | Prospective  | 23| 23 | SE | Philips iU22 [Philips Healthcare, Bothell, WA] | Area under the receiver operating characteristic curve: 0.917 [95% CI, 0.788 to 1.000] |
| Giannetti A27          | 2016 | Case report  | 1 | 1  | SE | Mylab Twice, Esaote, Italy | The SE pattern of reduced elasticity corresponded to histological fibrosis |
| Lu C22                 | 2017 | Prospective  | 105| 105 | pSWE | Philips Epiq 5, Philips IU-22 [Philips Healthcare, Bothell, WA] or Acuson S3000, [Siemens Medical Solutions, Malvern, PA] | Moderate correlation between SWE and muscular hypertrophy \[r = 0.59, p = 0.02\] |
| Serra C23              | 2017 | Prospective  | 26| 26 | SE | iU22 Philips [Philips, Bothell, WA, USA] | No significant correlation was found between mean SR and fibrosis score \[p = 0.877\], \[p = 0.002\] |
| Chen YJ24              | 2018 | Prospective, cross-sectional | 35| 35 | pSWE/2D-SWE | SuperSonic Imagine S.A., Aix-en-Provence, France | Sensitivity of 69.6% and specificity of 91.7% with AUC of 0.822 [95% CI, 0.685 to 0.960] \[p = 0.002\] |
| Quaia E25              | 2018 | Prospective  | 20| 20 | SE | iU22 Philips [Philips, Bothell, WA, USA] | Overall accuracy of SE alone of 30% to 35% in distinguishing fibrotic bowel; increased accuracy when combined with CEUS, US \[p <0.5\] for p-SWE [cutoff value >2.73 m/s] |
| Ding SS26              | 2019 | Retrospective | 25| 25 | SE, pSWE/2D-SWE | S2000 [Siemens Medical Solutions, Mountain View, CA] | Sensitivity of 75% and specificity of 100%, accuracy of 96% \[p <0.5\] for p-SWE [cutoff value >2.73 m/s] |
| Thimm MA28             | 2019 | Case report  | 3 | 3  | pSWE | EPIC scanner [Philips Healthcare] | Increased level of stiffness [1.58 m/s] consistent with histological fibrosis |

IBD, inflammatory bowel disease; UC, ulcerative colitis; CD, Crohn’s disease; US, ultrasound; USE, ultrasound elastography; SE, strain elastography; pSWE, point shear wave elastography; 2D-SWE, two-dimensional shear wave elastography; NS, not specified; SR, strain ratio; SD, standard deviation; AUC, area under the curve; CI, confidence interval; CEUS, contrast enhanced US; RTE, real-time elastography.

\(^{a}\)Exclusively IBD patients.
the title and abstract and, if necessary, the full publication, 146 records were rejected. After the reviewing process, a total of 12 publications met the inclusion criteria, all full-text [100%]. Figure 2 illustrates the screening and the selection process. All studies were single-centre experiences. The study design of the included papers was as follows: eight prospective studies [66.7%], one retrospective study [8.3%], and three case reports/series [25%]. In five studies there was a control group or a comparison with healthy tissue. None of the studies was excluded because of quality concerns.

Overall, a total of 275 IBD patients were included: 272 CD [98.9%] and three UC [1.1%]. Approximately half of the affected patients were male at 138 [50.2%], 121 were female [44%], and in 16 cases the gender was not specified [5.8%]. The age of the included patients ranged from 16 to 70 years.

4.1. Ultrasound elastography technique

Seven studies [58.3%] investigated the strain elastography [SE] as elastographic technique, and shear wave elastography [SWE] was adopted in four studies [41.6%]. In one study only [0.1%], authors explored the use of both techniques [SE, SWE]. The ultrasound elastography techniques and the devices used are presented in Table 1. As concerns the timing and the modality of the ultrasonographic evaluation, in most of the studies [10/12, 83.3%] the elastography assessment was pursued trans-abdominally and pre-operatively, and in two studies [16.7%] the assessment was ex vivo on the resected intestine. The evaluated segments were small and large bowel in the majority of the papers [66.7%] and small bowel only in four studies [33.3%]. Table 3 summarises further technical details of the included studies.

| Reference, first author | Risk of bias | Flow and timing | Applicability concerns | Overall risk | Author’s note |
|-------------------------|--------------|-----------------|------------------------|--------------|--------------|
| Stidham RW<sup>27</sup> | ☹ ☺ ☺ ☺ ☹ ☹ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ ☺ ☺ ☺ ☺ ☺ | Low/ moderate | Small sample size, intra-individual comparison only, concerns about validation and reproducibility |
| Havre RF<sup>21</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ ☺ ☺ | Low | Small sample, the primary observer was not blinded, ex vivo setup |
| Dillman JR<sup>19</sup> | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ ☺ ☺ | Low | Small sample size, ex vivo setup, single operator, concerns about validation and reproducibility |
| Baumgart DC<sup>20</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ ☺ | Low | Small sample size, inter- and intra-observer variability not assessed |
| Fraquelli M<sup>21</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ ☺ | Low | Small sample size, ultra-selected patients, concerns about validation and reproducibility |
| Giannetti A<sup>27</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Moderate/ high | Case report, concerns about validation and reproducibility |
| Lu C<sup>22</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Low | Possible patients’ selection bias, two different US devices, concerns about validation |
| Serra C<sup>23</sup> | ☺ ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Low | Small sample size, no control group, concerns about validation and reproducibility |
| Chen YJ<sup>24</sup> | ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Low | Small sample size, intra-individual comparison only, concerns about validation |
| Quaia E<sup>25</sup> | ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Low | Small sample size, concerns about the reference standard used, validation and reproducibility |
| Ding SS<sup>26</sup> | ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Moderate | Small sample size, retrospective design, concerns about validation and reproducibility |
| Thimm MA<sup>28</sup> | ☺ ☺ ☺ | ☺ ☺ ☺ | ☺ ☺ ☺ ☺ | Moderate/ High | Case report, concerns about validation and reproducibility |

The QUADAS tool<sup>16</sup> consists of four domains that assess: patient selection [was a consecutive or random sample of patients enrolled? was a case-control design avoided? did the study avoid inappropriate exclusions? could the selection of patients have introduced bias?]; index test [were the index test results interpreted without knowledge of the results of the reference standard? if a threshold was used, was it prespecified? could the conduct or interpretation of the index test have introduced bias?]; reference standard [is the reference standard likely to correctly classify the target condition? were the reference standard results interpreted without knowledge of the results of the index test?]; and flow of patients throughout the study design and timing of the index tests and reference standard [was there an appropriate interval between index tests and reference standard? did all patients receive a reference standard? did all patients receive the same reference standard? were all patients included in the analysis?].
The USE assessment was mainly semi-quantitative [i.e., with a colour map] in four studies,17,18,20,23 purely quantitative and qualitative in five,19,22–24,28 and one paper,27 respectively, and both semi-quantitative and quantitative in the works by Fraquelli et al. and by Ding et al.21,26 In most studies, the USE was performed by a single operator,17–19,21–24,26–28 and in five studies there was more than one sonographer [two to five].20–23,25 The inter-observer agreement was estimated as moderate and excellent by Quaia et al. and Fraquelli et al., respectively.21,25

4.2. Histological evaluation

The histological definition of fibrosis was ‘the evidence of increased submucosal collagen deposition’ in all the included studies. The use of Masson trichrome staining was specified in six works,17,19–22,24,28 and one paper,27 respectively, and both semi-quantitative and quantitative in the works by Fraquelli et al. and by Ding et al.21,26 In most studies, the USE was performed by a single operator,17–19,21–24,26–28 and in five studies there was more than one sonographer [two to five].20–23,25 The inter-observer agreement was estimated as moderate and excellent by Quaia et al. and Fraquelli et al., respectively.21,25

Table 3. Technical details of the included studies.

| Reference, first author | Histological specimen | Examined intestine | Ex vivo/ pre-operatively | Control group/comparison with healthy tissue | USE assessment | Histologic assessment |
|-------------------------|-----------------------|--------------------|--------------------------|-------------------------------------------|----------------|----------------------|
| Stidham RW17            | Surgical resection    | Small bowel        | Trans-abdominal, pre-operatively | Yes                                       | Semi-quantitative | Semi-quantitative     |
| Havre RE18              | Surgical resection    | Small- and large-bowel | Ex vivo                  | Yes                                       | Semi-quantitative | Semi-quantitative     |
| Dillman JR19            | Surgical resection    | Small- and large-bowel | Pre-operatively and ex vivo | No                                        | Quantitative     | Semi-quantitative     |
| Baumgart DC20           | Surgical resection    | Small- and large-bowel | Trans-abdominal, pre-operatively | Yes                                       | Semi-quantitative | Semi-quantitative     |
| Fraquelli M21           | Surgical resection    | Small- and large-bowel | Semi-quantitative and quantitative | Qualitative                            |                 | Qualitative           |
| Giannetti A27           | Surgical resection    | Small bowel        | Trans-abdominal, pre-operatively | No                                        | Quantitative     | Semi-quantitative     |
| Lu C22                  | Surgical resection    | Small- and large-bowel | Trans-abdominal, pre-operatively | No                                        | Quantitative     | Semi-quantitative     |
| Serra C23               | Surgical resection    | Small- and large-bowel | Trans-abdominal, pre-operatively | No                                        | Quantitative     | Semi-quantitative     |
| Chen YJ24               | Surgical resection    | Small- and large-bowel | Trans-abdominal, pre-operatively | No                                        | Quantitative     | Semi-quantitative     |
| Quaia E21               | Endoscopic biopsies or surgical resection | Small bowel | Trans-abdominal, pre-operatively | No                                        | Semi-quantitative | Semi-quantitative     |
| Ding SS26               | Endoscopic biopsies or surgical resection | Small- and large-bowel | Trans-abdominal, pre-operatively | No                                        | Semi-quantitative | Semi-quantitative     |
| Thimm MA28              | Surgical resection    | Small bowel        | Trans-abdominal, pre-operatively | No                                        | Quantitative     | Semi-quantitative     |

USE, ultrasound elastography.

Overall, an accurate differentiation of inflammatory from prevalently fibrotic intestinal tracts, compared with histology, was found in all the included papers,17,19–22,24,28 except for the ones by Havre et al. and Serra et al.18,23 These authors concluded in their studies that USE could not accurately distinguish the grade of inflammation from fibrosis.18,23

4.3. Accuracy of elastography in detecting fibrosis

In addition to USE, the histological fibrosis was compared with further ultrasound features such as conventional US and/or contrast enhanced US [CEUS] in seven publications.21–23,27,28
| Reference, first author | Inflammation score | Fibrosis score | Main achievements |
|-------------------------|------------------|----------------|-------------------|
| Dillman JR, 19 | 0 = 0 inflammation | 0 = 0 architectural distortion, no abnormal Masson trichrome staining | Statistically significant difference of SWE throughout the classes of fibrosis [low vs high fibrosis AUC max = 0.91] |
| | 1 = low level of inflammation with scattered infiltrating mononuclear cells | 1 = o architectural distortion, mild abnormal Masson trichrome staining in mucosa/ submucosa |  |
| | 2 = moderate inflammation with multiple foci | 2 = substantial abnormal mucosal/submucosal Masson trichrome staining with modest distortion of architecture but without obscuration of the mucosal/submucosal border |  |
| | 3 = high level of inflammation with increased vascular density and marked wall thickening | 3 = transmural fibrosis with abnormal Masson trichrome staining in all histological layers, transmural architectural distortion |  |
| | 4 = maximal severity of inflammation with transmural leukocyte infiltration and loss of goblet cells | 4 = maximal severity of inflammation with transmural leukocyte infiltration and loss of goblet cells |  |
| Baumgart DC, 20 | 0 = 0 architectural distortion, no abnormal Masson trichrome staining | 0 = 0 increased collagen deposition | Higher collagen content in affected versus unaffected segments associated with RTE-assessed strain [p <0.001] |
| | 1 = low level of inflammation with scattered infiltrating mononuclear cells | 1 = increased collagen deposition in submucosa |  |
| | 2 = moderate inflammation with multiple foci | 2 = increased collagen deposition in submucosa and mucosa |  |
| Fraquelli M, 21 | 0 = 0 architectural distortion, no abnormal Masson trichrome staining | 0 = 0 increased collagen deposition | Strain ratio was significantly correlated with the bowel fibrosis [AUC for severe fibrosis = 0.917]. Bowel fibrosis was the only independent determinant of the strain ratio at multivariate analysis |
| | 1 = low level of inflammation with scattered infiltrating mononuclear cells | 1 = increased collagen deposition in submucosa |  |
| | 2 = moderate inflammation with multiple foci | 2 = increased collagen deposition in submucosa and mucosa |  |
| | 3 = high level of inflammation with increased vascular density and marked wall thickening | 3 = increased collagen deposition in muscularis mucosa, submucosa, and mucosa, as well as thickening and disorganization of the muscularis mucosa |  |
| | 4 = maximal severity of inflammation with transmural leukocyte infiltration and loss of goblet cells | 4 = increased collagen deposition in muscularis propria, muscularis mucosa, submucosa, and mucosa |  |
| | 5 = transmural fibrosis with abnormal Masson trichrome staining in all histological layers, transmural architectural distortion | 5 = increased collagen throughout all layers, including serosa |  |
| Lu C, 22 | 0 = 0 architecture alteration | 0 = 0 fibrosis | No correlation between SWE and fibrosis score |
| | 1 = minimal fibrosis limited to submucosa | 1 = minimal fibrosis limited to submucosa |  |
| | 2 = submucosal muscular layer fibrosis | 2 = submucosal muscular layer fibrosis |  |
| | 3 = submucosal and muscular layer fibrosis between 30% and 60%, with preserved layers | 3 = submucosal and muscular layer fibrosis between 30% and 60%, with preserved layers |  |
| | 4 = Massive transmural fibrosis >60%, effacement of normal layers | 4 = Massive transmural fibrosis >60%, effacement of normal layers |  |
| Serra C, 23 | 0 = 0 polymorphonuclear or mononuclear leukocytes infiltrates | 0 = 0 polymorphonuclear or mononuclear leukocytes infiltrates | No correlation between SE and fibrosis score |
| | 1 = Mild cryptitis, leukocytes infiltrates limited to mucosa | 1 = mild fibrosis limited to submucosa |  |
| | 2 = Moderate cryptitis, crypt abscess, and leukocytes infiltrates until the submucosa | 2 = Submucosal and muscular layer fibrosis <30% |  |
| | 3 = Severe transmural inflammation with leukocytes infiltrates in all the layers | 3 = submucosal and muscular layer fibrosis between 30% and 60%, with preserved layers |  |
| | 4 = Massive transmural fibrosis >60%, effacement of normal layers | 4 = Massive transmural fibrosis >60%, effacement of normal layers |  |
| Chen YJ, 24 | 0 = 0 inflammation | 0 = 0 inflammation | SWE was significantly different throughout the classes of fibrosis [mild/moderate vs severe fibrosis AUC = 0.822; sensitivity = 69.6%; specificity = 91.7%]. No difference in SWE observed with respect to inflammation classes. Combined SWE + US showed a moderate agreement with the classes of strictures |
| | 1 = slight inflammation only | 1 = slight inflammation only |  |
| | 2 = submucosal foci of inflammation and/or foci of transmural inflammation | 2 = increased submucosal fibrosis, septa into muscularis propria and/or septa through muscularis propria, increase in sub-serosal collagen |  |
| | 3 = significant, dissecting, confluent transmural inflammation | 3 = significant transmural scar, marked sub-serosal collagen |  |
Table 4. Continued

| Reference, first author | Inflammation score | Fibrosis score | Main achievements |
|------------------------|--------------------|----------------|------------------|
| Quaia E $^{21}$        | 1. mucosal ulceration [grade 0–3] | Sections were scored as positive for fibrosis if at least moderate fibrosis was observed which involved the submucosa or deeper layers $^{9}$ | SE was able to differentiate between fibrosis and inflammation with a maximal AUC of 0.885 |
|                        | 2. edema [grade 0–3]         |                |                  |
|                        | 3. quantity [grade 0–3] of neutrophilic infiltration |                |                  |
|                        | 4. depth [grade 0–4] of neutrophilic infiltration $^{a}$ |                |                  |

RTE; real-time elastography; SE, strain elastography; AUC, area under the curve; SWE, shear wave elastography.

$^{a}$According to Likert-like scales.

$^{b}$According to Bataille F, Klebl F, Rümmele P, et al. Histopathological parameters as predictors for the course of Crohn’s disease. *Virchows Arch* 2003;443:501–7.

$^{c}$According to Chiorean MV, Sandrasegaran K, Saxena R, et al. Correlation of CT enteroclysis with surgical pathology in Crohn’s disease. *Am J Gastroenterol* 2007;102:2541–50 and/or Borley NR, Mortensen NJ, Jewell DP, et al. The relationship between inflammatory and serosal connective tissue changes in ileal Crohn’s disease: evidence for a possible causative link. *J Pathol* 2000;190:196–202.

$^{d}$According to Gupta RB, Harpaz N, Itzkowitz S, et al. Histologic inflammation is a risk factor for progression to colorectal neoplasia in ulcerative colitis: a cohort study. *Gastroenterology* 2007;133:1099–105.

$^{e}$According to Theiss AL, Fuller CR, Simmons JG, Liu B, Sartor RB, Lund PK. Growth hormone reduces the severity of fibrosis associated with chronic intestinal inflammation. *Gastroenterology* 2005;129:204–19.

5. Discussion

This systematic review illustrates the present understanding of the capability of USE in detecting and quantifying, whenever possible, the degree of fibrosis within the bowel wall of IBD patients.

Collectively, the analysis of the published literature testifies an overall shared moderate-to-good accuracy of USE in detecting histological fibrosis [10/12 studies]. $^{17,18,22,26}$ In detail, the accuracy of USE varied from 35% to 91% in all the included studies. $^{17}$ However, important concerns are raised regarding the heterogeneity of the USE modalities investigated [SE, SWE], in terms of both the input application/stimulus and the biomarkers analysed [i.e., strain ratio, pSWE, etc.] that do not allow formulation of unequivocal accuracy data. In particular, since SE only allows semi-quantitative assessments of stiffness, these are difficult to compare longitudinally. With respect to technical aspects, all the studies included had a limited cohort of patients [from 1 to 105 patients] and the US devices used were of different manufacturers, the Philips iU22 being the most used $^{21,23,25}$ [Table 1]. These methodological gaps might be only overcome by multicentre studies adopting a common USE equipment.

A further explanation of the variation of USE accuracy observed in our systematic review is the heterogeneity of the bowel segments analysed in the included studies. Indeed, the studies investigating exclusively the ileum in CD patients, where the pathological processes involve the whole bowel wall, reported higher rates of accuracy and better correlation between USE measurements and histology. $^{17,21,22}$

Possible selection bias must be additionally addressed: the investigation of USE in advanced stenosis candidates for surgical resection might have returned higher rates of tissue fibrosis, possibly enhancing the accuracy of USE. The incorporation of a control group and the inclusion of different stages of disease in the study design are necessary to reduce this kind of bias.

The main strength of our systematic review was to address the accuracy of USE in detecting fibrosis exclusively in comparison with histological assessment. To our knowledge, this inclusion criterion was never adopted by previous systematic reviews on the topic of USE in IBD. $^{18}$ Of note, the definition of fibrosis was univocally adopted by all the studies, but the histological assessment varied from semi-quantitative $^{17–26}$ to merely qualitative, $^{27,28}$ thus limiting the uniformity of the data and a direct comparison between the studies. A further main finding emerges from our analysis, which is the lack of a standardised histological score to quantify fibrosis. Indeed, a strong discrepancy with respect to the reference standard adopted [i.e., also endoscopic biopsies, which limits the proper assessment of the submucosal fibrotic changes] and the histological quantification of fibrosis was found between the included studies.

It appears clear that USE has been more extensively investigated in CD than in UC [only one study included, three UC patients], but with the growing adoption of US also in the monitoring of UC this trend will reasonably change in the near future.

Interestingly, several publications suggested the integration of USE with conventional B-mode US and CEUS in order to gain greater accuracy. $^{21–25,27,28}$ Indeed, this concept has been broadly explored and is well known by experienced bowel sonographers who are used to combining different qualitative and quantitative features within activity scores. $^{33,34}$

The matter of operator dependency remains for USE, as for all ultrasonographic methods. When estimated, there was a moderate-to-high inter-reader agreement in SE measurements, $^{21,25}$ and we can speculate that inter-reader agreement might be superior in the case of SWE. A consensus on specific skills and training for USE operators is yet to be specifically established.

Another relevant issue regards the quality assessment of the included studies [Table 2]. $^{18}$ Indeed, despite addressing the main methodological features of scientific studies, many limitations could not be addressed [i.e., the exiguous sample size, the lack of validation, and reproducibility] [Table 2].

The main limitation of this work is that no meta-analysis was performed due to the lack of standardisation between the results of the included studies; further limitations derive from
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the discussed heterogeneity of the included studies [different techniques of USE, different reference standards]. In our view, the so far gathered data on USE deserve endorsement to be incorporated into the management algorithms of IBD, whereas USE does not appear to add any specific information to guide clinical decisions. Indeed, current European Federation of Societies for Ultrasound in Medicine and Biologys guidelines instruct on USE with a relatively low level of evidence and suggest using it to characterise bowel wall lesions exclusively in CD.\[35\]

The appeal of USE lies in its non-invasiveness and repeatability. Indeed, in the treat-to-target era a new physiological surrogate endpoint, such as the quantification of the intestinal fibrosis, would be warmly welcome by dedicated physicians.

Our systematic review endorses that elastography cannot replace the tissue specimen yet, at least in the field of bowel ultrasound and IBD. The applicability of this technique to the bowel wall, compared with parenchymal organs might be limited and challenged by the unique features of the intestine [i.e., peristalsis, the peritoneum, the structure in layers].

In conclusion, despite the data gathered so far, the role of USE in the detection and quantification of fibrosis in IBD patients requires additional research with properly designed randomised clinical trials. Moreover, long-term data on patients followed up with USE longitudinally over time are warranted as well.

The data underlying this article are available in its online Supplementary material and upon request to the corresponding author.

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Conflict of Interest
SD has served as a speaker, consultant, and advisory board member for Schering-Plough, AbbVie, MSD, UCB Pharma, Ferring, Cellerix, Millenium Takeda, Nycomed, Pharmacosmos, Actelion, Alphawassermer, Genentech, Grunenthal, Pfizer, Astra Zeneca, Novo Nordisk, Cosmo Pharmaceuticals, Vifor and Johnson & Johnson, Nikkiso Europe GMBH, Theravance. LP-B has served as a speaker, consultant, and advisory board member for Merck, Abbvie, Janssen, Genentech, Mitsubishi, Ferring, Norgine, Tillots, Vifor, Hospira/Pfizer, Celltrion, Takeda, Biogaran, Boehringer-Ingelheim, Lilly, HAC-Pharma, Index Pharmaceuticals, Amgen, Sandoz, Forward Pharma GmbH, Celgene, Biogen, Lycera, Samsung Bioepis, Theravance. MA received consulting fees from Nikkiso Europe, Mundipharma, Janssen, Abbvie, and Pfizer.

Author Contributions
FF and ADB performed the research. ADB and FF wrote the manuscript. MA, LPB, and SD critically reviewed the content of the paper. MA and FF conceived the subject of the paper, contributed to the critical interpretation, and supervised the project. All authors approved the final version of the manuscript.

Supplementary Data
Supplementary data are available at ECCO-JCC online.

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