Inflammation and fibrosis in Crohn’s disease: location-matched histological correlation of small bowel ultrasound features

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Received: 18 January 2020 / Revised: 20 May 2020 / Accepted: 26 May 2020 © The Author(s) 2020

Abstract

Purpose To evaluate the utility of mural and extramural sonographic features of Crohn’s Disease as potential imaging biomarkers of inflammation and fibrosis against whole-mount histological sections.

Methods Twelve Crohn’s disease patients (Mean age 35(25–69), 7 males) underwent small bowel ultrasound prior to small bowel resection. Two radiologists in consensus graded multiple parameters including mural, mucosal and submucosal thickness, submucosal/mesenteric echogenicity and clarity and mural Doppler signal in 50 selected bowel cross-sections. Matching with histological sampling sites was facilitated via scanning of the resected specimen. A histopathologist scored acute and chronic inflammation, and fibrosis (using histological scoring systems) following analysis of whole mount block sections. The association between sonographic observations and histopathological scores was examined via univariable and multivariable analysis.

Results In univariate analyses, bowel wall thickness (regression co-efficient and 95% CI 0.8 (0.3, 1.3) p = 0.001), mesenteric fat echogenicity (8.7(3.0, 14.5) p = 0.005), submucosal layer thickness (7.4(1.2, 13.5) p = 0.02), submucosal layer clarity (4.4(0.6, 8.2) p = 0.02) and mucosal layer thickness (4.6(1.8, 7.4) p = 0.001) were all significantly associated with acute inflammation. Mesenteric fat echogenicity (674(8.67, 52404) p = 0.009), submucosal layer thickness (79.9(2.16, 2951) p = 0.02) and mucosal layer thickness (13.6(1.54, 121) p = 0.02) were significantly associated with chronic inflammation. Submucosal layer echogenicity (p = 0.03), clarity (25.0(1.76, 356) p = 0.02) and mucosal layer thickness (53.8(3.19, 908) p = 0.006) were significantly associated with fibrosis. In multivariable analyses, wall and mucosal thickness remained significantly associated with acute inflammation (p = 0.02), mesenteric fat echogenicity with chronic inflammation (p = 0.009) and mucosal thickness (p = 0.006) with fibrosis.

Electronic supplementary material The online version of this article (https://doi.org/10.1007/s00261-020-02603-6) contains supplementary material, which is available to authorized users.

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Published online: 20 June 2020
Conclusion  Multiple sonographic parameters are associated with histological phenotypes in Crohn’s disease although there is overlap between ultrasonic stigmata of acute inflammation, chronic inflammation and fibrosis.

Graphic Abstract

Keywords  Crohn’s disease · Ultrasound · Inflammation · Fibrosis

Abbreviations
CD  Crohn’s disease
SBUS  Small bowel ultrasound
MRE  Magnetic resonance enterography
CDAI  Crohn’s Disease Activity Index
AIS  Histological acute inflammation score
CIS  Histological chronic inflammation score
MM  Muscularis mucosa

Introduction

Cross-sectional imaging assessment of disease activity is fundamental to management of Crohn’s disease (CD) [1, 2]. Both small bowel ultrasound (SBUS) and magnetic resonance enterography (MRE) are accurate for detecting active disease [3–5] although SBUS, however, has several advantages, including patient acceptability [6], relative simplicity and general availability.

Studies supporting SBUS’s ability to assess disease activity have employed a variety of clinical, serological and endoscopic comparators [7] as a single standard of reference is lacking. The effectiveness of clinical indices such as the Crohn’s Disease Activity Index (CDAI) is increasingly questioned [8, 9], while colonoscopy and mucosal biopsies only evaluate superficial disease [10, 11] in at best, a limited length of terminal ileum. Transmural inflammation characterises CD [12], so histological analysis of resected small bowel specimens arguably serves as the most complete reference standard to validate imaging biomarkers: it looks beyond the mucosa, and allows comprehensive assessment of all intestinal layers, including quantification of mural fibrosis. Although resection specimens have been employed in several studies investigating CT and MRE [13–17], relatively few SBUS studies have used such a standard of reference [18–22]. These have mainly focused on a limited number of mural observations, such as Doppler signal or tested pre-defined definitions of activity. None has systematically evaluated the full range mural and extra mural observations apparent on SBUS in CD. The purpose of our study was to evaluate the utility of multiple mural and extramural sonographic features as potential imaging biomarkers of inflammation and fibrosis against a whole-mount histological section reference standard.

Materials and methods

The study had research ethics committee approval. All patients gave written informed consent. Patients with a known diagnosis of CD over the age of 16 undergoing small bowel surgical resection between the dates of 01/01/2014 and 01/02/2015 were eligible. Members of the local research team identified suitable patients from the operating lists of two colorectal surgeons with a special interest in Crohn’s disease. Patients were excluded if they were pregnant (n = 0). In total, 16 patients were eligible, of whom four declined participation. The patient’s sex, age, Montreal classification, and reason for surgery were recorded.

SBUS

Consenting patients underwent SBUS within 2 weeks of their planned operation date. Patients in whom the surgery
was subsequently delayed beyond 2 weeks remained eligible providing they did not undergo any treatment changes, and the surgery was performed within 8 weeks of SBUS. Patients fasted for 4 h prior to SBUS and did not receive any oral or intravenous contrast. Two radiologists (20 years and 3 year’s experience in small bowel imaging) performed all ultrasounds in consensus. Prior imaging and clinical notes were reviewed before scanning to ensure the bowel segment scheduled for resection was interrogated. SBUS was undertaken on a standard platform (ELogiq E9, GE Medical Systems Ltd, Buckinghamshire, UK) using both curvilinear (C2-9) and high-resolution (9L and ML6-15) probes, including colour Doppler imaging (typical flow 6–9 m/s). Detailed observations (described below, see Online Appendix 1) were made for between 4 and 8 cross-sections through the abnormal small bowel in each patient. Sections were chosen to, as far as possible, encompass all the different US observations present in the diseased bowel segment. For example, if the bowel contained two different submucosal echogenicity patterns, both were included in the selected sections. Additionally, sonographically normal bowel sections were selected towards the periphery of diseased segment for comparison with diseased bowel.

Each SBUS section was systematically scored for multiple sonographic variables (for example wall thickness, mesenteric fat echogenicity, anti-mesenteric border clarity, submucosal layer echogenicity and mural Doppler vascular pattern) as detailed in Online Appendix 1 (Fig. 1). The observations were based on those already described in the literature, and the researchers own clinical experience. A pictorial and descriptive key was available to the radiologists for reference during their evaluation (Online Appendix 2).

**Co-localisation between US and histological sampling**

The location of each chosen cross-section of the bowel was recorded by the interrogating radiologists with reference to distances from fixed structures such as ileo-caecal valve or fistula (if present), using the method described by Punwani et al. [14]. To further facilitate co-localisation between bowel sections scored on SBUS and subsequent histological sectioning, the fixed resection specimen (and associated mesentery) was scanned by the same two radiologists who had performed the in vivo scan. Guided by the recorded locations, and considering any identifying sonographic characteristics of the bowel and/or surrounding tissues, the axial sections scored on the in vivo US were located on the surgical specimen and marked using blunt needles for subsequent histological sampling [14].

![Fig. 1 B-Mode US demonstrating multiple variables evaluated. Additionally, the presence of ulceration, increased Doppler vascular pattern and stenosis causing functional obstruction were recorded.](image-url)
Histological analysis

Histological assessment was performed by a specialist GI histopathologist (15 years of experience), blinded to the SBUS scoring and clinical information pertaining to underlying disease activity. Whole mount block sections taken through each identified cross-section were analysed after conventional H&E staining. For all histological analysis, abnormal bowel was compared to macroscopically “normal bowel” at the resection margins, which acted as an internal control. A previously described scoring system was used to classify the disease within the specimen. The inflammation was reported in each slice according to the method of Borley et al. as detailed in Online Appendix 3 [23]. Specifically, a histological acute inflammation score (AIS) was created by adding the scores for eosinophils, neutrophils, cryptitis, crypt destruction and ulceration. A histological chronic inflammatory score (CIS) was created by adding the scores for structural architectural changes, chronic inflammatory infiltrate, presence of granulomas and eosinophils in the lamina propria. These have been previously described in studies relating to Ulcerative colitis but applied to histological correlation studies in CD also [14, 24, 25].

Histological grading of mural fibrosis was undertaken on a three point scale, as previously described [13, 14] and detailed in Online Appendix 4. The specialist GI histopathologist further classified threshold values of the AIS (> 6), CIS (> 3) and fibrosis (> 0) for descriptive purposes only.

Of note, previous work has confirmed that mural thickness in formalin-fixed bowel specimens does not appreciably differ from that of fresh unfixed specimens, allowing direct comparisons of measured mural thickness between ultrasound and histology [14].

Statistical analysis

All analyses examined the association between the SBUS parameters and the histological outcomes. A feature of the data was that there were multiple measurements from each patient. This violated an assumption of independence, assumed by many statistical methods. To allow for these repeat measurements, the analysis was performed using multilevel regression methods.

AIS was considered a continuous outcome and analysed using multilevel linear regression. For this variable the results are presented as a regression co-efficient with 95% confidence intervals (Coefficient (95% CI)). The remaining outcomes (CIS and fibrosis score) were ordinal in nature and were analysed using multilevel ordered logistic regression (using the meologit command in the software program Stata 14, StataCorp LLC, Texas, USA). For these variables the results are presented as Odds Ratio with 95% confidence intervals.

The analyses were performed in two stages. First, the association between each separate SBUS parameter and each histological outcome was examined in a series of univariable analyses. Subsequently, the joint association of the parameters was examined in a multivariable analysis. Due to the relatively small sample size, the multivariable analysis was restricted only to those parameters showing some association with the outcome from the univariable analyses ($p \leq 0.1$). A backwards selection procedure was used to retain only the statistically significant variables in the final model.

One patient scored category 4 (uniform hypoechoic mesenteric fat) for mesenteric fat echogenicity which was omitted from the results as the finding occurred only once and this was felt too infrequent to be reliably considered as a discrete category in itself. Agreement between histological and ultrasonic bowel wall thickness measurements was tested using Bland-Altman limits of agreement.

Results

Patient characteristics

Twelve patients (7 male, mean age 35) were included. Patients had a range of phenotypes according to the Montreal classification (Table 1). The mean time interval between the ultrasound and surgery was 9 days. All patients had been clinically stable in the pre-operative period with no alteration in medication and were not on corticosteroids.

Specimen characteristics

In total, 50 small bowel cross-sections were scored from the 12 patients. Three to six sections were taken from individual resected segments with a median of 4 sections per segment (Online Appendix 5). The majority of sections demonstrated a combination of acute and chronic inflammation (AIS mean 7.6, range 0-14, CIS Mean 3.8, Range 0-6) and fibrosis (fibrosis score mean 0.6, range 0–2) (Online Appendix 5). Thirteen of the 50 sections demonstrated above threshold values for acute inflammation (> 6), chronic inflammation (> 3) and fibrosis (> 0), whilst a further 14 sections demonstrated above threshold values for both acute and chronic inflammation, but not fibrosis. Four sections demonstrated above threshold values of acute inflammation only, three sections demonstrated above threshold values of chronic inflammation only and six sections demonstrated above threshold values of fibrosis only.
Mural thickness

Overall, in vivo ultrasound estimates of wall thickness ranged from 3 to 12 mm (mean, 6.2 mm, median 5.5 mm) and histologically measured specimen wall thicknesses which ranged from 1.7 mm to 15 mm (mean 5.3 mm, median 4.7 mm). A scatter plot of ultrasonic against histological bowel wall thickness measurement is given in Fig. 2. The mean difference in measurements was 1 mm (95% Bland–Altman limits of agreement − 5 to 7 mm).

Sonographic parameter correlation with histological acute inflammation

Univariate analysis

Bowel wall thickness, higher bowel wall thickness category, mesenteric fat echogenicity, submucosal layer thickness, submucosal layer clarity and mucosal layer thickness were all individually significantly associated with the AIS (Table 2).

Greater bowel wall thickness (and higher categories of bowel wall thickness categories) were associated with higher acute inflammatory scores. A 1 mm increase in wall thickness was associated with a 0.8 unit increase in the AIS.

Table 1  Patient characteristics, interval between ultrasound and surgery and reason for surgery

| Patient | Age (years) | Sex | Montreal classification | Ultrasound scan to surgery (days) | Reason for surgery |
|---------|-------------|-----|-------------------------|----------------------------------|-------------------|
| 1       | 27          | M   | A1L1B2                  | 5                                | Obstructive stricture |
| 2       | 37          | F   | A2L1B2                  | 7                                | Obstructive stricture |
| 3       | 35          | M   | A2L1B3                  | 1                                | Abscess            |
| 4       | 24          | M   | A1L3B3p                 | 1                                | Obstructive stricture |
| 5       | 25          | M   | A1L1B2                  | 5                                | Obstructive stricture |
| 6       | 26          | M   | A1L3B3p                 | 0                                | Failed medical management |
| 7       | 45          | F   | A2L3B3                  | 5                                | Obstructive stricture |
| 8       | 69          | M   | A2L3B3                  | 6                                | Obstructive stricture |
| 9       | 34          | F   | A2L2B1                  | 11                               | Failed medical management |
| 10      | 23          | F   | A1L3B3                  | 44                               | Fistula            |
| 11      | 38          | M   | A1L3B1                  | 21                               | Failed medical management |
| 12      | 36          | F   | A1L1B3                  | 0                                | Obstructive stricture/adhesions |
| Mean    | 35          |     |                         |                                  |                   |

Fig. 2  A scatter plot of ultrasonic against histological bowel wall thickness measurement
Mesenteric fat echogenicity pattern one (focal hyper-echoic without fat wrap), two (focal hyperechoic with fat wrap) and three (stratified) were associated with higher AIS than category zero (normal). Of these, pattern two had the strongest association with AIS which was almost 9 units higher than sections with normal mesenteric fat (pattern zero).

A thickened submucosa (submucosal layer thickness pattern one), an ill-defined submucosa (submucosal layer clarity pattern one) and a thickened mucosa (mucosal layer thickness patterns one or two) were associated with higher AIS. Specimens with a thickened submucosa (submucosal layer thickness pattern one) had AIS that were over seven units higher than specimens with a normal submucosal thickness (submucosal layer thickness pattern zero).

Images with matched sonographic and histological descriptors are provided as examples of normal (Fig. 3) and abnormal segments (Fig. 4, 5).

### Multivariate analysis

The final multivariate model is summarised in Table 3.

Both bowel wall thickness and mucosal layer thickness were found to be independently associated with the acute histological score. After adjusting for these two variables, no further variables were found to be statistically significant.
As in the univariable analyses, a greater bowel wall thickness and a thickened mucosa with or without submucosal thickness (mucosal layer thickness patterns one or two) were associated with higher acute histological scores.

**Sonographic parameter correlation with histological chronic inflammation**

**Univariate analysis**

Mesenteric fat echogenicity, submucosal layer thickness and mucosal layer thickness were statistically associated with the CIS (Table 4).

All mesenteric fat patterns were associated with CIS, notably hyperechoic mesentery with fat wrapping (mesenteric fat echogenicity pattern two).

A thickened submucosa (submucosal layer thickness pattern one) and thickened mucosa with or without submucosal thickness (mucosal layer thickness patterns one or two) were associated with a higher CIS.

**Multivariate analysis**

Only mesenteric fat echogenicity was included in final model. No further variables were significant after factoring in this variable. As this was the only variable in the final model, the results are equivalent to those from the univariable analysis.

**Sonographic parameter correlation with histological fibrosis**

**Univariate analysis**

Submucosal layer echogenicity, submucosal layer clarity and mucosal layer thickness were significantly associated with fibrosis (Table 5).

Submucosal layer echogenicity could not be analysed using regression methods as all sections with a normal submucosal echogenicity (submucosal layer echogenicity pattern zero) demonstrated no fibrosis. It was thus not possible to take account of the repeat measurements per patient and caution should be exercised when interpreting the significance of this association. The highest fibrosis scores were for sections with a reduced submucosal echogenicity (submucosal layer echogenicity pattern one) and for specimens with an increased submucosal echogenicity containing low echogenicity bands (submucosal layer echogenicity pattern three).

Sections with an ill-defined submucosa (submucosal layer clarity pattern one) had higher fibrosis scores than sections with a normal, well defined submucosa (submucosal layer clarity pattern zero). Additionally, specimens with a thickened mucosa with or without submucosal thickness (mucosal layer thickness pattern one or two) had higher fibrosis scores than specimens with a normal mucosal thickness (mucosal
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Layer thickness zero). A mucosal layer thickness pattern one or two was associated with 50 times greater odds of being in the next highest fibrosis category.

Multivariate analysis

Only mucosal layer thickness was included in final model. There was no additional effect of any other factors after adjusting for this variable. As this was the only variable in the final model, the results were equivalent to those from the univariable analysis.

Discussion

We demonstrate that several sonographic features correlate with histologically graded acute inflammation, chronic inflammation and fibrosis. There is relatively little comparative data using full thickness pathological specimens in the literature. Previous comparisons have investigated Doppler ultrasound abnormalities only [18, 19], or intravenous contrast enhanced ultrasound (US) perfusion and bowel wall thickness [17]. Both Maconi et al. and Hata et al. pre-categorised sonographic findings of mural thickening and mural stratification patterns [20, 21], while Nylund et al. performed SBUS on the resected specimen only and reported their findings according to changes in the different mural layers [22].

We found that mucosal layer thickness was the only sonographic feature associated with all three histological categories. The mucosa is the primary layer involved

Table 3 Multilevel regression model demonstrating the statistically significant sonographic variables associated with the histological acute inflammatory score (AIS)

| Variable             | Category | Coefficient (95% CI) | p-value |
|----------------------|----------|----------------------|---------|
| Wall thickness       | –        | 0.6 (0.1, 1.1)       | 0.02    |
| Mucosal layer thickness | 0         | 0                    | 0.02    |
|                      | 1 and 2  | 3.4 (0.6, 6.3)       |         |

Fig. 5 B-mode US images (TOP) with matched histological sections demonstrating the importance of fat wrapping and mucosal thickening in evaluating active inflammation. a A histologically normal segment (AIS 0, CIS 1, fibrosis 0) presents with hypechoic mesenteric fat without fat wrapping (white star for US and black star for histology), a prominent, well defined, homogenous submucosa (white arrow for US and black arrow for histology) and a normal mucosa. b Histologically demonstrated fat wrapping (black stars showing extent), mild submucosal oedema (black arrow) and mild active mucosal inflammation (dashed black arrow) (AIS 8, CIS 4, fibrosis 1). Sonographically, there is hypechoic mesenteric fat with fat wrapping (white stars showing extent), a thickened, well defined, homogenous submucosa (white arrow) and mucosal/muscularis mucosa thickening (dashed white arrow). Both images are acquired from the same patient.
in inflammation. Histologically there are a range of processes ranging from architectural distortion of the crypts at the mucosal surface, infiltration by inflammatory cells and mucin depletions [10, 12]. Mucosal healing is the goal of successful treatment [26], and our findings demonstrate the importance of evaluating the mucosa during sonography; the presence of a thickened mucosa suggests ongoing inflammation. Furthermore, studies have demonstrated hyperplasia of the muscularis mucosa in fibrosis which may in part explain why the mucosal layer is thickened in sections demonstrating fibrosis, with and without active inflammation [27]. Nylund et al. reported that it was possible to identify the muscularis mucosa (MM) on US in 48 of 69 sections where it was histologically thickened, and 52 of 54 sections where it was histologically normal. Their results suggested that submucosal fibrosis was more likely to be present with a non-visible (normal) MM than in those with a visible (thickened) MM [22].

| Variable                        | Category | Odds ratio (95% CI) | p value |
|---------------------------------|----------|---------------------|---------|
| Wall thickness                  | –        | 1.37 (0.94, 1.99)   | 0.10    |
| Wall thickening                 | 0        | 1                   | 0.09    |
|                                | 1 or 2   | 9.45 (1.08, 82.9)   |         |
|                                | 3        | 17.6 (1.10, 281)    |         |
| Mesenteric fat echogenicity*    | 0        | 1                   | 0.009   |
|                                | 1        | 26.1 (0.40, 1688)   |         |
|                                | 2        | 674 (8.67, 52,404)  |         |
|                                | 3        | 131 (0.32, 54,209)  |         |
| Anti-mesentric border definition| 0        | 1                   | 0.24    |
|                                | 1        | 2.94 (0.50, 17.4)   |         |
| Mesentric border definition     | 0        | 1                   | 0.82    |
|                                | 1 or 2   | 1.24 (0.20, 7.74)   |         |
| Submucosal thickness            | 0        | 1                   | 0.02    |
|                                | 1        | 79.9 (2.16, 2951)   |         |
| Submucosal echogenicity         | 0        | 1                   | 0.20    |
|                                | 1        | 32.1 (0.62, 1674)   |         |
|                                | 2        | 199 (0.76, 51,659)  |         |
|                                | 3        | 89.6 (1.13, 7116)   |         |
| Submucosal clarity              | 0        | 1                   | 0.08    |
|                                | 1        | 14.5 (0.72, 295)    |         |
| Mucosal layer thickness         | 0        | 1                   | 0.02    |
|                                | 1 and 2  | 13.6 (1.54, 121)    |         |
|                                | 1 and 2  | 2.87 (0.34, 24)     |         |
| Mural Doppler pattern           | 0        | 1                   | 0.33    |
|                                | 1 and 2  | 2.87 (0.34, 24.2)   |         |

Table 4 The association of sonographic variables with the chronic inflammatory score (CIS) examined through univariable analyses

| Variable                        | Category | Odds ratio (95% CI) | p value |
|---------------------------------|----------|---------------------|---------|
| Wall thickness                  | –        | 1.14 (0.83, 1.56)   | 0.44    |
| Wall thickening                 | 0        | 1                   | 0.15    |
|                                | 1 or 2   | 25.0 (0.81, 770)    |         |
|                                | 3        | 28.5 (0.57, 1419)   |         |
| Mesenteric fat echogenicity*    | 0        | 1                   | 0.06    |
|                                | 1        |                     |         |
|                                | 2        |                     |         |
|                                | 3        |                     |         |
| Anti-mesentric border definition| 0        | 1                   | 0.49    |
|                                | 1        | 0.56 (0.11, 2.91)   |         |
| Mesentric border definition     | 0        | 1                   | 0.47    |
|                                | 1 or 2   | 0.44 (0.05, 4.13)   |         |
| Submucosal thickness            | 0        | 1                   | 0.23    |
|                                | 1        | 5.93 (0.32, 110)    |         |
| Submucosal echogenicity         | 0        | †                   |         |
|                                | 1        |                     |         |
|                                | 2        |                     |         |
|                                | 3        |                     |         |
| Submucosal clarity              | 0        | 1                   | 0.03    |
|                                | 1        | 25.0 (1.76, 356)    |         |
| Mucosal layer thickness         | 0        | 1                   | 0.006   |
|                                | 1 and 2  | 53.8 (3.19, 908)    |         |
| Mural Doppler pattern           | 0        | 1                   | 0.50    |
|                                | 1 and 2  | 0.49 (0.06, 3.98)   |         |

Table 5 The association of sonographic variables with the fibrosis score examined through univariable analyses

The results are summarised in the form of odds ratios (and corresponding confidence intervals) due to the ordinal nature of the fibrosis histological score. The odds ratios represent the relative change in the odds of the next highest outcome score in one situation relative to another. For the one continuous variable (BWT) the coefficients indicate the change in the odds of being in the next category for one-unit increase in that measure. For the categorical variables, the odds ratios indicate the difference of being in the next outcome category for each ultrasound category relative to a baseline category. p-values indicating the significance of the results are also presented

*Omitting one patient with category 4
†Unable to calculate odds ratios as all results in one category having same outcome. Analysis using Kruskal–Wallis test
Bowel wall thickness was significantly correlated with histological scores of acute inflammation, but interestingly not with chronic inflammation or fibrosis. Bowel wall thickness is the most frequently investigated parameter to assess disease presence and activity within the published literature [7]. Wilkens et al. have demonstrated that US is highly accurate in measuring bowel wall thickness, reporting that US measured mural thickness is on average 0.4 mm thicker than histological measurements, whilst MRE measurements are 1.4 mm thicker [17]. We also found reasonable agreement between ultrasound and histological bowel wall thickness measurement, albeit with some outliers. Previous studies across a range of imaging modalities have demonstrated bowel wall thickness is more strongly associated with active inflammation than fibrosis [13, 14]. For example, Moreno et al. compared sonographic findings with endoscopy and endoscopic biopsies in 178 segments (across 30 patients). Bowel wall thickness was the strongest US predictor of endoscopic remission with a normal bowel wall thickness (<3 mm) predicting endoscopic remission with a PPV of 97.1% (77.1–99.7%) [28]. These findings are unsurprising as acute inflammation is associated with histological findings of oedema and inflammatory infiltrate which subsequently results in mural thickening [23]. However, studies using endoscopic biopsies cannot assess for any relationship between wall thickness and fibrosis, which requires full thickness specimen histology. Indeed, using histological specimens, Wilkens et al reported a significant correlation of 0.4 between ultrasonic bowel wall thickness measurements and histological fibrosis, although they found no such relationship using MRI measured wall thickness. Other MRE studies using full thickness specimen histology have however demonstrated a correlation between bowel wall thickness and fibrosis [15, 29], which is at odds with our findings. The dominant process in fibrosis is smooth muscle hyperplasia (in addition to proliferation of fibroblasts and myofibroblasts) which may not lead to the magnitude of mural thickening seen with acute inflammation [27]. However, it is reasonably established that fibrosis thickens the bowel wall so our negative findings are most likely due in part to the relatively small number of sections with fibrosis and the relatively limited histological scoring system used.

Abnormal patterns of mesenteric fat were associated with acute and chronic inflammation but not fibrosis. Our categorisation of mesenteric fat echogenicity incorporated various parameters including hypertrophy and fat wrapping. Mesenteric fat changes have infrequently been investigated in other histological correlation studies. Maconi et al. reported that patients with mesenteric fat hypertrophy exhibited higher clinical (CDAI) and serological (CRP and ESR) markers of inflammation than those without mesenteric fat hypertrophy [30]. This would support our findings that mesenteric fat abnormalities are linked to mural inflammation rather than mural fibrosis. Pallotta demonstrated that mesenteric fat hypertrophy was associated with the presence of fistulae at histology but not with the presence of strictures [31]. Seven of our cohort had obstructive strictures which histologically often exhibited a mixture of acute and chronic inflammation, and only 1 had a fistulae. Our data therefore suggests stricturing disease is associated with mesenteric fat changes. Indeed, histologically, both fat wrapping and mesenteric thickening are known to be associated with acute and chronic inflammation [23]. It has been proposed that these processes may be a combination of at least two events: perivascular inflammation with mesenteric fibrosis and muscularis propria contraction (noting that mural fibrosis and mesenteric fibrosis are different entities). As such alterations in mesenteric fat echogenicity, represent transmural extension of more advanced Crohn’s disease [32]. Whilst many of the patients in our cohort presented with obstructive strictures all but one of the resected segments demonstrated features of acute or chronic inflammation or both.

We found that many sonographic features of the submucosal layer are also associated with inflammation and fibrosis in the univariate analyses. We tested both submucosal layer thickness, echogenicity and clarity as individual features. This arguably adds to the complexity of scoring system. Clarity specifically looks at the interface between the submucosa and the other bowel wall layers (as opposed to echogenicity) and may reflect a different pathological process, for example transmural cellular infiltration. Indeed, the fact that submucosal echogenicity and clarity had differing associations with the various histological findings does suggest they are giving, in part, different information. A thickened or an ill-defined submucosal was associated with active inflammation, while a thickened submucosal was also associated with chronic inflammation. Fibrosis was associated with reduced submucosal echogenicity, increased submucosal echogenicity with hypoechoic bands and an ill-defined submucosa. Hata et al. combined many of the individual sonographic criteria we describe into discrete patterns, and reported that mural thickening without stratification was associated with severe inflammation whilst mural thickening with stratification was associated with moderate inflammation [21]. Maconi et al concluded that stenosis with stratified or mixed echo pattern shows a significantly greater mucosal and submucosal fibrosis in comparison to a hypoechoic echo pattern [20]. Nylund et al. performed in vitro US on 14 resected small bowel segments and classified the submucosa into three categories (Echo-rich, Sporadic echo poor elements and diffuse echo poor elements) [22]. This parameter is comparable to our variable for submucosal layer echogenicity. They demonstrated that increasing echo poor elements in the submucosa were linked to fibrosis. These findings are confirmed by our study; we demonstrated that highest fibrosis scores were for specimens with a reduced
submucosal echogenicity (pattern one) and for specimens with an increased submucosal echogenicity containing low echogenicity bands (pattern three).

Interestingly, we found no correlation between Doppler US findings and inflammation, and in this regard is at odds which much of the current literature. For example, in a study of 10 patients, Sasaki et al., reported significantly greater inflammatory cell infiltration \( p = 0.043 \) and mural vascularity \( p = 0.047 \) in specimens with increased mural Doppler flow \[18\]. None of the cohort were taking steroids which could have potentially help explain our findings by reducing acute inflammation. However, by definition, a patient cohort destined for surgery is likely to have received long term anti-inflammatory medication and have less active disease than a cohort prior to initiation of immunosuppressive therapy for example. The more chronic nature of our cohort may in part explain the lack of action between histological activity and doppler signal.

This study had limitations. The number of patients we evaluated were relatively low but at par with other studies reporting imaging and histological correlation in CD. We had numerous data points from the same patients for which we adjusted our statistical methodology, although this was not always possible. There was some minor variation in the number of histological sections (3–6, median 4) taken from individual resected segments, in order to maximise the variety of sonographic features being assessed. Whilst there is potential for individual patients to dominate the population, these seems unlikely given the relatively even spread of sections across the cohort. One finding, mesenteric fat hypertrophy was scored in a solitary category, 4 on a single occasion only. This was omitted from the assessment of this variable as it could not be reliably handled in the statistical analysis and as such this is a limitation. This was a single centre-study undertaken by the same two sonographers and the same histopathologist. Whilst this has the advantage of standardising interpretation it impacts the generisability of our findings especially given that US is anecdotally more operator dependent than other imaging modalities. We used a consensus process for the ultrasonic observation and did not test inter-observer variation. However, detailed analysis of inter-observer agreement between multiple reader for the various observations across the cohort. Finding, mesenteric fat hypertrophy was scored in a solitary category, 4 on a single occasion only. This was omitted from the assessment of this variable as it could not be reliably handled in the statistical analysis and as such this is a limitation. This was a single centre-study undertaken by the same two sonographers and the same histopathologist.

In conclusion, multiple sonographic parameters are associated with acute inflammation, chronic inflammation and fibrosis. As expected there is overlap in US findings, akin to histology. Segments that demonstrate mucosal, submucosal and mural thickening (> 3 mm) together with abnormal mesentry and loss of definition of the submucosal layer likely have underlying acute inflammation. Fibrosis is more likely to be present in segments with mucosal thickening, and if the submucosa is hypoechoic (either diffuse or focally) or ill-defined.

Author contribution GB: Study design, patient recruitment, data collection, data analysis, data interpretation, drafting manuscript. MRJ: Study design, patient recruitment, data collection, data analysis, data interpretation, drafting manuscript. AH: Study design, revising manuscript. PB: data analysis, data interpretation. AW: Patient recruitment, revising manuscript. RC: Patient recruitment, revising manuscript. SH: Study design, revising manuscript. SAT: Study design, patient recruitment, data collection, data analysis, data interpretation, drafting and revising manuscript.

Funding This study was undertaken at the Comprehensive Biomedical Research Centre, University College Hospital, London, which received a proportion of the funding from the Crohn’s and colitis UK, Royal College of Radiologists. SAT is a NIHR senior investigator.

Compliance with ethical standards

Conflict of interest SAT is a research consultant to Robarts. All other authors declare that they have no conflict of interest.

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