



# Feasibility of *in vivo* magnetic resonance elastography of mesenteric adipose tissue in Crohn's disease

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**Background:** Although there is growing evidence that functional involvement and structural changes of mesenteric adipose tissue (MAT) influence the course of Crohn's disease (CD), its viscoelastic properties remain elusive. Therefore, we aimed to investigate the viscoelastic properties of MAT in CD using magnetic resonance elastography (MRE), providing reference values for CD diagnosis.

**Methods:** In this prospective proof-of-concept study, 31 subjects (CD: n=11; healthy controls: n=20) were consecutively enrolled in a specialized care center for inflammatory bowel diseases (tertiary/quaternary care). Inclusion criteria for the CD patients were a clinically and endoscopically established diagnosis of CD based on the clinical record, absence of other concurrent bowel diseases, scheduled surgery for the following day, and age of at least 18 years. Diagnoses were confirmed by histological analysis of the resected bowel the day after MRE. Subjects were investigated using MRE at 1.5-T with frequencies of 40–70 Hz. To retrieve shear wave speed (SWS), volumes of interest (VOIs) in MAT were drawn adjacent to CD lesions (MAT<sub>CD</sub>) and on the opposite side without adjacent bowel lesions in patients (MAT<sub>CD\_Opp</sub>) and controls (MAT<sub>CTRL</sub>). The presented study is not registered in the clinical trial platform.

**Results:** A statistically significant decrease in mean SWS of 7% was found for MAT<sub>CD\_Opp</sub> vs. MAT<sub>CTRL</sub> (0.76±0.05 vs. 0.82±0.04 m/s, P=0.012), whereas there was a nonsignificant trend with an 8% increase for

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MAT<sub>CD</sub> vs. MAT<sub>CD,Opp</sub> (0.82±0.07 vs. 0.76±0.05 m/s, P=0.098) and no difference for MAT<sub>CD</sub> vs. MAT<sub>CTRL</sub>. Preliminary area under the receiver operating characteristic curve (AUC) analysis showed diagnostic accuracy in detecting CD to be excellent for SWS of MAT<sub>CD,Opp</sub> [AUC =0.82; 95% confidence interval (CI): 0.64–0.96] but poor for SWS of MAT<sub>CD</sub> (AUC =0.52; 95% CI: 0.34–0.73).

**Conclusions:** This study demonstrates the feasibility of MRE of MAT and presents preliminary reference values for CD patients and healthy controls. Our results motivate further studies for the biophysical characterization of MAT in inflammatory bowel disease.

**Keywords:** Inflammatory bowel disease; Crohn's disease (CD); magnetic resonance imaging (MRI); magnetic resonance elastography (MRE); mesenteric adipose tissue (MAT)

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## Introduction

Crohn's disease (CD) is a severe condition with a growing incidence worldwide, with an incidence of over 6 per 100,000 person-years in broad parts of Western Europe and North America and an annual rise of up to 11% in newly industrialized countries (1). It is characterized by chronic recurrent inflammation, frequently followed by complications such as strictures, abscesses, and fistulas (1-3). The exact underlying pathology remains unclear.

In 1932, Crohn *et al.* first described hyperplasia of mesenteric adipose tissue (MAT) adjacent to intestinal inflammation (4). Corresponding indirect signs were described by Sellink *et al.* in 1974, who observed an increased inter-enteric spacing in abdominal radiography with oral contrast agent administration (5). The hyperplastic fat wrapping—also known as creeping fat—around the intestine is pathognomonic of CD and is thus not found in other inflammatory bowel diseases such as ulcerative colitis.

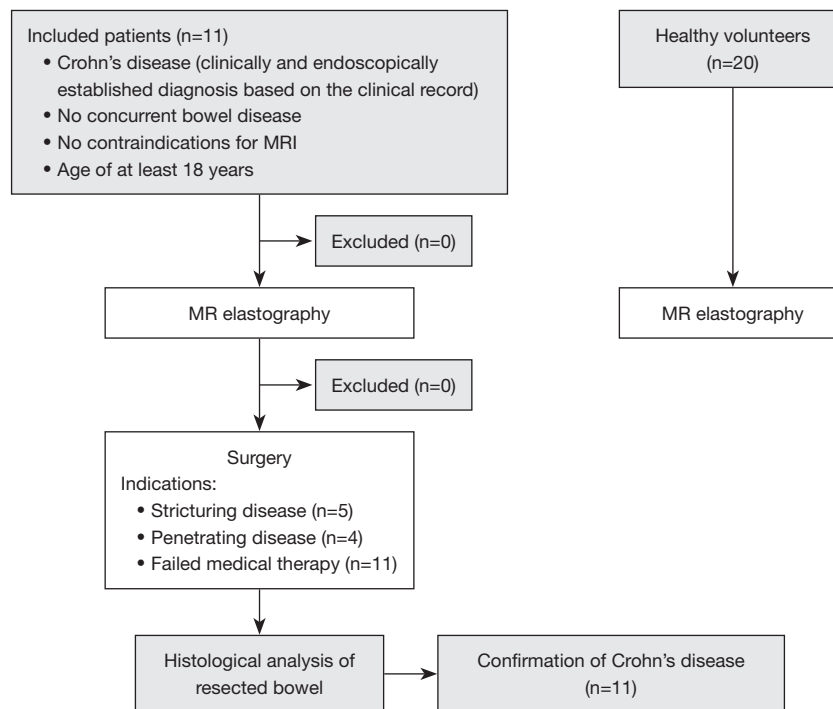
More recent studies indicate that beyond the morphological changes, structural and functional aspects of MAT influence progression, prognosis, and outcome in CD (6-9). Increased permeability of the bowel wall with bacterial translocation to the MAT is presumed to have an active role in the course of the disease (10-12). In contrast, MAT is also considered a potential secondary barrier around the inflamed intestine with localized anti-inflammatory function and a protective effect against a systemic inflammatory response. It can be distinguished histopathologically from healthy MAT by its altered extracellular matrix, distinctively small adipocytes, and a specific microenvironment defined by higher levels of adipokines and immune cell infiltration (8,13). All these new insights have led to the assumption of MAT as a distinct organ and a possible target for future

therapies (14,15). Different adipocytokines secreted by visceral fat were considered potential therapeutic targets (15). For example, inhibitors of the adipocytokine visfatin, which is known to be elevated in CD, were investigated in phase I trials (15-19). Nevertheless, no approved therapeutic agent has yet been found to target MAT in CD (15).

Quantitative imaging techniques, among them magnetic resonance elastography (MRE), have recently been extended to the assessment of intestinal involvement in inflammatory bowel disease (20-22). MRE has emerged as a method for the quantitative mapping of biomechanical properties of different biological tissues (23). Several links between viscoelasticity and clinical endpoints have been identified. In a feasibility study on patients with inflammatory bowel disease compared to healthy volunteers, shear wave speed (SWS) representing stiffness of the bowel wall was significantly increased in patients with CD and ulcerative colitis (20). Avila *et al.* correlated the stiffness value to intestinal fibrosis in patients with CD, allowing predictions on the further clinical course based on the fibrosis score (21).

Imaging methods for direct visualization of what surgeons and pathologists describe as creeping fat are evolving, but the influence of its biomechanical signature remains elusive (24-26). We hypothesize that the development of creeping fat as a secondary barrier to CD-related inflammation leads to altered viscoelastic properties, which can be noninvasively measured by MRE.

Therefore, we conducted a study investigating the viscoelastic properties of MAT in CD patients and healthy subjects using *in vivo* MRE. We present this article in accordance with the STARD reporting checklist (27) (available at <https://qims.amegroups.com/article/view/10.21037/qims-23-41/rc>).



**Figure 1** Flow diagram of the included patients. MR, magnetic resonance.

## Methods

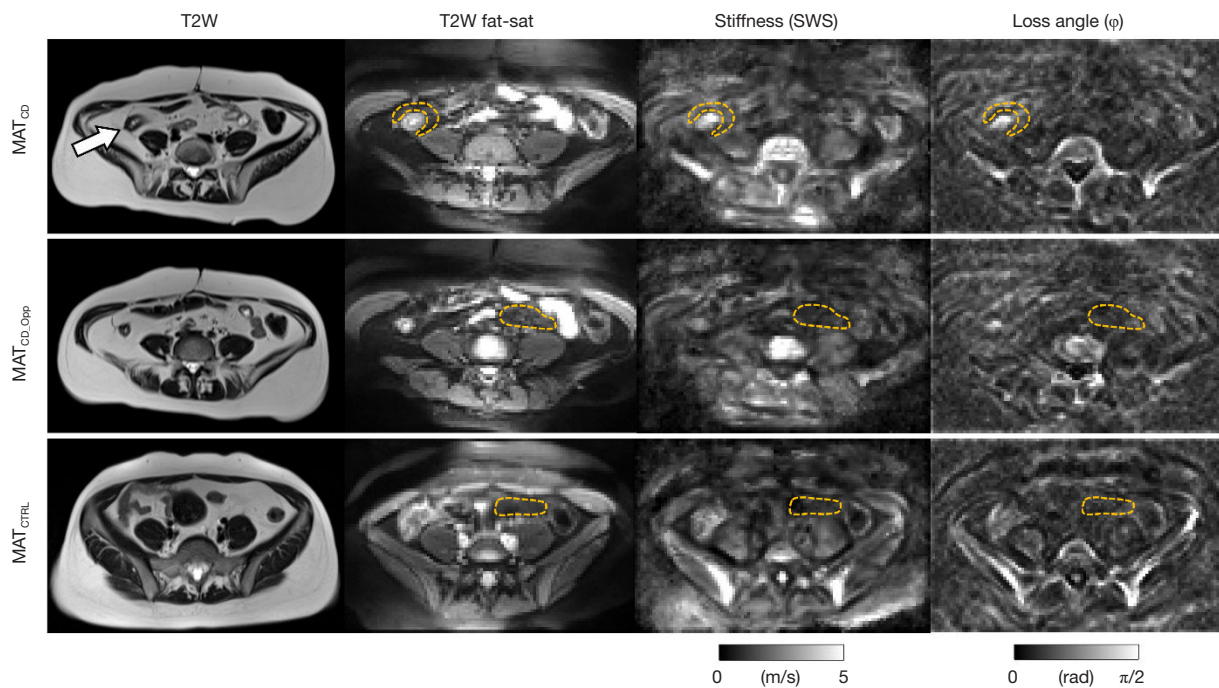
### Subjects

The study was approved by the Institutional Review Board of Charité – Universitätsmedizin Berlin [protocol code (A1/075/17) and date of approval (5-11-2017)] and was conducted in accordance with the Declaration of Helsinki (as revised in 2013). Written informed consent was obtained from patients. Our institution is a specialized care center for inflammatory bowel diseases (tertiary/quaternary care). All elastography examinations were prospectively performed at a single institution from July 2019 through December 2020. Thirty-one subjects were consecutively included: CD patients (n=11) and healthy controls (n=20). Only patients with CD and no other inflammatory bowel diseases were included to focus on the elastography of creeping fat. For reasons of availability, more healthy individuals were examined. Inclusion criteria for patients were a clinically and endoscopically established diagnosis of CD based on the clinical record with clinically indicated surgery, absence of other concurrent bowel diseases, absence of general contraindications to MRI, and age of at least 18 years. Exclusion criteria were defined as increased peristalsis and fecal impaction. All patients met the eligibility criteria, and

none were excluded. CD-related characteristics of patients are reported according to the Montreal classification, and diagnoses were confirmed by histological analysis of the resected bowel the day after MRE, which was used as standard of reference in all patients (28). The indication for surgery was stricture in five patients, penetrating disease in four patients, and failed medical therapy in all patients. *Figure 1* shows a flow diagram of the included patients. Of note, MRE data on the bowel wall have been reported before (20).

### Tomoelelastography

All examinations were performed on a 1.5-T MRI scanner (Magnetom Aera, Siemens Healthineers, Erlangen, Germany) using the spine-array coil in combination with an 18-channel body phased-array coil positioned on the patient's abdomen. Multifrequency MRE at 40, 50, 60, and 70 Hz with tomoelelastography postprocessing was performed using 4 compressed-air drivers (20) and a single-shot spin-echo echo planar imaging (EPI) sequence (29,30). Further MRE scan parameters were: slices in axial orientation, 25; matrix, 100×66; resolution, 3×3×5 mm<sup>3</sup>; TR, 3,070 ms, and TE, 50 ms; and total MRE acquisition



**Figure 2** Cases. The yellow dotted lines show delineated VOIs in the MAT. The upper row shows a 41-year-old woman with CD with a SWS of  $0.80 \pm 0.10$  m/s and  $\phi$  of  $0.47 \pm 0.05$  rad in the  $\text{MAT}_{\text{CD}}$ . The inflamed terminal ileum is indicated by an arrow. In the same patient, the middle row shows  $\text{MAT}_{\text{CD\_Opp}}$  with a decreased SWS of  $0.74 \pm 0.06$  m/s and  $\phi$  of  $0.44 \pm 0.11$  rad. The lower row depicts a 31-year-old healthy woman with an SWS of  $0.84 \pm 0.15$  m/s and  $\phi$  of  $0.58 \pm 0.21$  rad ( $\text{MAT}_{\text{CTRL}}$ ). T2W, T2-weighted; SWS, shear wave speed;  $\phi$ , loss angle;  $\text{MAT}_{\text{CD}}$ , MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $\text{MAT}_{\text{CD\_Opp}}$ , MAT without adjacent bowel lesion;  $\text{MAT}_{\text{CTRL}}$ , MAT of healthy controls; VOIs, volumes of interest.

time, 5:17 min. Subjects were breathing freely throughout MRE acquisition. No oral or intravenous contrast media or spasmolytic agents were administered. Quantitative maps of SWS (m/s) and loss angle of the complex modulus ( $\phi$ , rad) representing stiffness and viscosity-related dispersion of stiffness over frequency, respectively, were generated using the publicly available tomoelastography processing pipeline (<https://bioqic-apps.charite.de>) (31-35). This pipeline offers a combination of two viscoelastic parameters that have been previously proposed for the investigation of many organs such as intestine (20), liver (36,37), kidney (38), parotid glands (39), and prostate (40). Reconstruction of SWS is based on the  $k$ -multifrequency dual elasto-visco ( $k$ -MDEV) inversion which represents a phase-gradient-based inversion with a Laplacian unwrapping method (35). Reconstruction of  $\phi$  is based on the MDEV inversion which represents a direct inversion using a magnitude formulation and a gradient-based unwrapping method (41). Of note,  $\phi$  ranges from 0 rad for pure solids to  $\pi/2$  rad for pure fluids.

### Volume of interest (VOI) analysis

VOIs were drawn on axial SWS maps using Matlab R2021b (MathWorks, Natick, MA, USA). Manually matched fat-saturated T2-weighted magnetic resonance images were used to locate MAT in healthy controls ( $\text{MAT}_{\text{CTRL}}$ ) and bowel lesions in patients to place VOIs in adjacent MAT ( $\text{MAT}_{\text{CD}}$ ). Additionally, VOIs in CD patients were also placed in MAT on the opposite side without adjacent bowel lesions ( $\text{MAT}_{\text{CD\_Opp}}$ ). VOIs were drawn independently by two readers (reader 1, LJJ: senior radiology resident with more than 5 years of experience; reader 2, RR: board-certified radiologist with over 10 years of experience in abdominal MRE). For the two readers, it was apparent on MRI scans whether participants were patients with CD or healthy subjects. Further clinical data were not available to the readers at the time of radiological assessment. In consensus, bowel lesions were defined as the most severely affected pathologically wall-thickened ( $>3$  mm) intestinal loops. Moreover, VOIs were drawn on gluteal subcutaneous adipose tissue for inherent comparison. VOI placement is illustrated in *Figure 2*.

**Table 1** Demographic baseline data of patients and healthy controls

| Parameters                                  | Patients (n=11) | Healthy controls (n=20) | P value |
|---|-----------------|-------------------------|---------|
| Gender (women/men), n (%)                   | 6/5 (54.5/45.5) | 10/10 (50.0/50.0)       | –       |
| Age (years), median [interquartile spacing] | 38 [25]         | 31 [10]                 | 0.47    |
| BMI (kg/m <sup>2</sup> ), mean ± SD         | 21.8±3.4        | 22.0±2.6                | 0.84    |

BMI, body mass index; SD, standard deviation.

**Table 2** Clinical details of CD patients according to the Montreal classification (29)

| Montreal classification            | Patients (n=11) |
|------------------------------------|-----------------|
| Age group at initial diagnosis     |                 |
| A1, ≤16 years at diagnosis         | 1               |
| A2, 17–40 years at diagnosis       | 8               |
| A3, >40 years at diagnosis         | 2               |
| Disease extent                     |                 |
| L1, terminal ileum                 | 7               |
| L2, colon                          | 2               |
| L3, ileocolon                      | 2               |
| L4, upper gastrointestinal tract   | 0               |
| Disease behavior                   |                 |
| B1, nonstricturing, nonpenetrating | 2               |
| B2, stricturing                    | 5               |
| B3, penetrating                    | 4               |
| B3p, perianal penetrating          | 0               |
| Perianal disease modifier          |                 |
| No perianal disease                | 9               |
| Perianal disease present           | 2               |
| Medication history                 |                 |
| Corticosteroids                    | 3               |
| Infliximab                         | 2               |
| Adalimumab                         | 2               |
| Azathioprine                       | 4               |

CD, Crohn's disease.

### Statistical analysis

Continuous variables were described as mean ± standard deviation for data with normal distribution and as median with interquartile range (IQR) for data without normal distribution. Categorical variables were described as

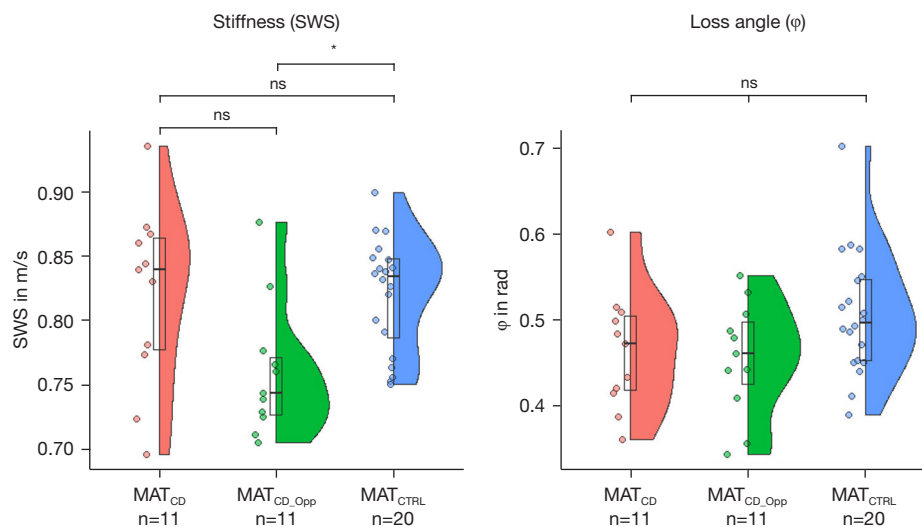
percentages. Group values of patients (MAT<sub>CD</sub> and MAT<sub>CD\_Opp</sub>) and healthy controls (MAT<sub>CTRL</sub>) were tested for differences using one-way analysis of variance (ANOVA) with Bonferroni-adjusted post-hoc analysis. Group values of gluteal subcutaneous adipose tissue between patients and healthy controls were compared with Welch's two sample *t*-test. Preliminary estimates of diagnostic accuracy in predicting CD were obtained by area under the receiver operating characteristic curve (AUC) analysis with a 95% confidence interval (CI). The Pearson correlation coefficient (*r*) was determined to assess the relationship between MRE parameters, age, body mass index (BMI), duration of disease, and C-reactive protein. All cases were re-evaluated to assess reproducibility by calculating intraclass correlation coefficients (ICCs) for interobserver agreement with 95% CI. Interobserver reliability was classified from poor to excellent (poor, <0.50; moderate, 0.50–0.75; good, 0.75–0.90; excellent, >0.90) (42). A P value <0.05 was considered to indicate statistical significance. R Studio (version 1.3.1093; R-Foundation, Vienna, Austria) was used for statistical analysis.

### Results

MRE examinations were feasible in all 11 patients (median age: 38 years; IQR: 25 years; mean BMI: 21.8±3.4 kg/m<sup>2</sup>; 6 women) and 20 healthy controls (median age: 31 years, IQR: 10 years; mean BMI: 22.0±2.6 kg/m<sup>2</sup>; 10 women). An overview of demographic baseline data is shown in *Table 1*. Patients had a mean duration of CD of 150.0±111.4 months and a median C-reactive protein of 36.25 mg/L (IQR: 88.6 mg/L). Further CD-related clinical details of patients are listed in *Table 2*. *Figure 2* shows representative cases of a CD patient and healthy control. Median VOI size was 7.6 cm<sup>3</sup> (IQR: 6.9 cm<sup>3</sup>).

### Group values

As shown in *Figure 3*, a statistically significant decrease



**Figure 3** Box plots (displaying median, upper, and lower quartiles, and whiskers) combined with half-violin plots and experimental data points of stiffness and  $\phi$  comparing  $MAT_{CD}$ , on the  $MAT_{CD\_Opp}$ , and in  $MAT_{CTRL}$ . Statistically significant differences are indicated by an asterisk (\*) for  $P < 0.05$ ; n.s., not significant. SWS, shear wave speed;  $\phi$ , loss angle;  $MAT_{CD}$ , MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $MAT_{CD\_Opp}$ , MAT without adjacent bowel lesion;  $MAT_{CTRL}$ , MAT of healthy controls.

in mean SWS of 7% was found for  $MAT_{CD\_Opp}$  vs.  $MAT_{CTRL}$  ( $0.76 \pm 0.05$  vs.  $0.82 \pm 0.04$  m/s,  $P = 0.012$ ), whereas a nonsignificant trend with an 8% increase in mean SWS was found for  $MAT_{CD}$  vs.  $MAT_{CD\_Opp}$  ( $0.82 \pm 0.07$  vs.  $0.76 \pm 0.05$  m/s,  $P > 0.99$ ) and no difference for  $MAT_{CD}$  vs.  $MAT_{CTRL}$  ( $0.82 \pm 0.07$  vs.  $0.82 \pm 0.04$  m/s,  $P > 0.99$ ). For mean  $\phi$ , no significant difference was found for any of these pairwise comparisons with  $P = 0.179$ , 1.000, 0.342; respectively ( $MAT_{CD}$ :  $0.46 \pm 0.07$  rad;  $MAT_{CD\_Opp}$ :  $0.46 \pm 0.07$  rad;  $MAT_{CTRL}$ :  $0.82 \pm 0.04$  rad). Moreover, no significant difference was found for subcutaneous adipose tissue between CD patients and healthy controls (SWS:  $1.00 \pm 0.22$  vs.  $1.08 \pm 0.24$  m/s,  $P = 0.38$ ;  $\phi$ :  $0.52 \pm 0.04$  vs.  $0.55 \pm 0.05$  rad,  $P = 0.09$ ).

### Correlation analysis

Pairwise Pearson correlation analysis of MRE parameters, age, BMI, duration of disease, and C-reactive protein did not reveal any significant correlations, except for a strong correlation between age and duration of disease ( $r = 0.83$ ,  $P = 0.01$ ) and a strong negative correlation between  $\phi$  of  $MAT_{CD\_Opp}$  and BMI ( $r = -0.79$ ,  $P = 0.004$ ). Table 3 provides an overview of all parameters of the correlation analysis. Of note, there was one data point missing for the C-reactive protein value and three data points for the duration of the disease because these patients were referred to our

department of surgery from another hospital.

### Diagnostic performance

Preliminary AUC analysis showed an excellent diagnostic accuracy of SWS of  $MAT_{CD\_Opp}$  in detecting CD with an AUC = 0.82 (95% CI: 0.64–0.96), whereas accuracy of SWS of  $MAT_{CD}$ ,  $\phi$  of  $MAT_{CD}$  and  $\phi$  of  $MAT_{CD\_Opp}$  was poor to moderate with an AUC = 0.52, 0.67, 0.69 (95% CI: 0.34–0.73, 0.48–0.84, 0.51–0.85), respectively. Moreover, sensitivities, specificities, positive predictive values, and negative predictive values with optimal cut-offs using the Youden index are listed in Table 4. A cross-tabulation of SWS and  $\phi$  results against the reference standard according to the AUC analysis is shown in Table 5.

### Reproducibility

Interobserver agreement was good for  $MAT_{CD}$  with ICC = 0.83 (95% CI: 0.39–0.95) for SWS and ICC = 0.90 (95% CI: 0.63–0.97) for  $\phi$ , good to excellent for  $MAT_{CD\_Opp}$  with ICC = 0.83 (95% CI: 0.37–0.95) for SWS and ICC = 0.92 (95% CI: 0.70–0.98) for  $\phi$ , and good for  $MAT_{CTRL}$  with ICC = 0.84 (95% CI: 0.56–0.94) for SWS and ICC = 0.90 (95% CI: 0.75–0.96) for  $\phi$ . Moreover, agreement between the two readers for SWS and  $\phi$  is shown in Bland-Altman plots in Figure 4.

**Table 3** Correlation analysis with Pearson correlation coefficients  $r$  (upper triangle array) and corresponding  $P$  values (lower triangle array) for all demographic and imaging parameters of patients ( $n=11$ )

| Categories                         | SWS of MAT <sub>CD</sub> | $\varphi$ of MAT <sub>CD</sub> | SWS of MAT <sub>CD_Opp</sub> | $\varphi$ of MAT <sub>CD_Opp</sub> | BMI                        | Age                      | Duration of disease      | CRP                      |
|------------------------------------|--------------------------|--------------------------------|------------------------------|------------------------------------|----------------------------|--------------------------|--------------------------|--------------------------|
| SWS of MAT <sub>CD</sub>           | –                        | 0.38<br>(–0.28 to 0.80)        | 0.43<br>(–0.23 to 0.82)      | 0.21<br>(–0.45 to 0.72)            | –0.03<br>(–0.62 to 0.58)   | –0.22<br>(–0.72 to 0.44) | 0.36<br>(–0.31 to 0.79)  | –0.13<br>(–0.68 to 0.51) |
| $\varphi$ of MAT <sub>CD</sub>     | 0.25                     | –                              | 0.43<br>(–0.23 to 0.82)      | 0.27<br>(–0.39 to 0.75)            | –0.05<br>(–0.63 to 0.57)   | 0.15<br>(–0.49 to 0.69)  | 0.16<br>(–0.49 to 0.69)  | 0.27<br>(–0.39 to 0.75)  |
| SWS of MAT <sub>CD_Opp</sub>       | 0.19                     | 0.19                           | –                            | 0.49<br>(–0.16 to 0.84)            | –0.28<br>(–0.75 to 0.38)   | –0.43<br>(–0.82 to 0.23) | –0.26<br>(–0.74 to 0.40) | 0.37<br>(–0.30 to 0.79)  |
| $\varphi$ of MAT <sub>CD_Opp</sub> | 0.54                     | 0.42                           | 0.13                         | –                                  | –0.79*<br>(–0.94 to –0.36) | –0.44<br>(–0.82 to 0.22) | –0.57<br>(–0.87 to 0.05) | 0.21<br>(–0.45 to 0.72)  |
| BMI                                | 0.93                     | 0.89                           | 0.41                         | 0.004*                             | –                          | 0.35<br>(–0.32 to 0.79)  | 0.35<br>(–0.32 to 0.79)  | –0.42<br>(–0.81 to 0.24) |
| Age                                | 0.52                     | 0.67                           | 0.19                         | 0.18                               | 0.30                       | –                        | 0.83*<br>(0.46 to 0.95)  | 0.14<br>(–0.50 to 0.68)  |
| Duration                           | 0.38                     | 0.71                           | 0.53                         | 0.14                               | 0.40                       | 0.01*                    | –                        | 0.09<br>(–0.54 to 0.65)  |
| CRP                                | 0.72                     | 0.44                           | 0.29                         | 0.56                               | 0.22                       | 0.70                     | 0.86                     | –                        |

Values in parentheses indicate 95% CI of  $r$ . \*, significant correlation ( $P < 0.05$ ). SWS, shear wave speed; MAT<sub>CD</sub>, MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $\varphi$ , loss angle; MAT<sub>CD\_Opp</sub>, MAT without adjacent bowel lesion; BMI, body mass index; CRP, C-reactive protein; CI, confidence interval.

**Table 4** Diagnostic performance

| Categories                         | Sn               | Sp               | NPV              | PPV              | AUC              | Cut-off  |
|------------------------------------|------------------|------------------|------------------|------------------|------------------|----------|
| SWS of MAT <sub>CD</sub>           | 0.36 (0.13–0.63) | 0.80 (0.65–0.95) | 0.70 (0.54–0.85) | 0.50 (0.20–0.80) | 0.52 (0.34–0.73) | 0.85 m/s |
| SWS of MAT <sub>CD_Opp</sub>       | 0.75 (0.59–0.91) | 0.82 (0.59–0.99) | 0.64 (0.43–0.86) | 0.88 (0.73–0.99) | 0.82 (0.64–0.96) | 0.78 m/s |
| $\varphi$ of MAT <sub>CD</sub>     | 0.90 (0.78–0.99) | 0.46 (0.20–0.71) | 0.71 (0.40–0.99) | 0.75 (0.59–0.88) | 0.67 (0.48–0.84) | 0.43 rad |
| $\varphi$ of MAT <sub>CD_Opp</sub> | 0.85 (0.71–0.99) | 0.45 (0.20–0.71) | 0.63 (0.33–0.99) | 0.74 (0.57–0.89) | 0.69 (0.51–0.85) | 0.44 rad |

Values in parentheses indicate 95% CI. Sn, sensitivity; Sp, specificity; NPV, negative predictive value; PPV, positive predictive value; AUC, area under the receiver operating characteristic curve (with an optimal cut-off using the Youden index); SWS, shear wave speed; MAT<sub>CD</sub>, MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $\varphi$ , loss angle; MAT<sub>CD\_Opp</sub>, MAT without adjacent bowel lesion; CI, confidence interval.

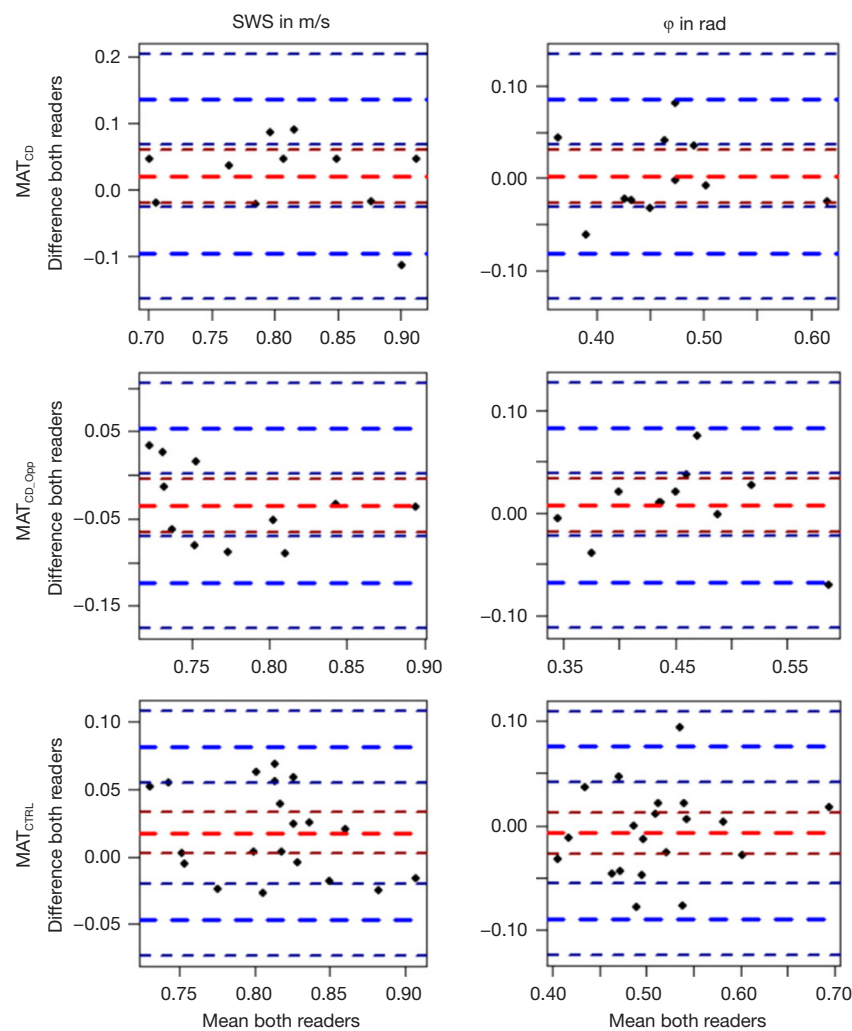
**Table 5** Comparison of the cut-off-based MRE diagnosis between the number of CD patients and healthy volunteers

| Categories                         | MRE diagnosis, n (%) |           | Total, n |
|------------------------------------|----------------------|-----------|----------|
|                                    | Wrong                | Right     |          |
| SWS of MAT <sub>CD</sub>           | 11 (35.5)            | 20 (64.5) | 31       |
| SWS of MAT <sub>CD_Opp</sub>       | 7 (22.6)             | 24 (77.4) | 31       |
| $\varphi$ of MAT <sub>CD</sub>     | 9 (29.0)             | 22 (71.0) | 31       |
| $\varphi$ of MAT <sub>CD_Opp</sub> | 10 (32.3)            | 23 (74.2) | 31       |

CD, Crohn's disease; MRE, magnetic resonance elastography; SWS, shear wave speed; MAT<sub>CD</sub>, MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $\varphi$ , loss angle; MAT<sub>CD\_Opp</sub>, MAT on the opposite side without adjacent bowel lesions.

## Discussion

In this proof-of-concept study, we demonstrate the feasibility of *in vivo* MRE of MAT and present preliminary reference values for CD patients and healthy controls. AUC results show diagnostic accuracy in detecting CD to be excellent for SWS of MAT<sub>CD\_Opp</sub> (AUC = 0.82, 95% CI: 0.64–0.96) but poor for SWS of MAT<sub>CD</sub> (AUC = 0.52, 95% CI: 0.34–0.73). We found a statistically significant decrease in mean stiffness of 7% for MAT<sub>CD\_Opp</sub> compared to MAT<sub>CTRL</sub>. This observation may suggest a general softening of MAT in CD with a trend toward 8% higher stiffness



**Figure 4** Bland-Altman plots show agreement between the two readers for SWS and  $\phi$ . Single red bold dashed line = mean difference; paired blue bold dashed lines = 95% limits of agreement (paired dark red and blue thin dashed lines = lower and upper limits of 95% CI). SWS, shear wave speed;  $\phi$ , loss angle;  $\text{MAT}_{\text{CD}}$ , MAT with adjacent bowel lesion; MAT, mesenteric adipose tissue;  $\text{MAT}_{\text{CD\_Opp}}$ , MAT without adjacent bowel lesion;  $\text{MAT}_{\text{CTRL}}$ , MAT of healthy controls; CI, confidence interval.

in areas of focal inflammatory bowel lesions ( $\text{MAT}_{\text{CD}}$ ). This result could be attributed to the increased density of network elements by characteristically small adipocytes in CD, and enhanced leukocyte infiltration, which both contribute to the secondary barrier function of creeping fat. This biophysical signature may prevent bacterial translocation from the bowel walls with CD-related increased permeability, reducing the inflammatory response from systemic to localized. Moreover, the statistically significant negative correlation between  $\phi$  of  $\text{MAT}_{\text{CD\_Opp}}$  and BMI may be related to an increase in fluid-like properties of MAT with decreasing BMI in CD. Nevertheless, further

studies with histopathological reference are needed to corroborate this preliminary hypothesis.

As the composition of extracellular matrix components of MAT is constantly changing during inflammation (13), a noninvasive imaging modality allowing the characterization of these changes would be a valuable new diagnostic tool. Despite the growing body of imaging studies aiming to quantify creeping fat in CD, little is known about its viscoelastic properties. For instance, van Schelt *et al.* examined seven patients with active CD and seven healthy volunteers using tomoelastography from 30–60 Hz and histopathological assessment in a subset of patients (43).



Preliminary results show no significant difference for mean SWS (CD:  $0.80 \pm 0.21$  vs. healthy:  $0.67 \pm 0.07$  m/s,  $P=0.18$ ), whereas a significant increase was found for mean  $\phi$  (CD:  $0.58 \pm 0.15$  vs. healthy:  $0.45 \pm 0.08$  rad,  $P=0.02$ ). While our results are of the same order of magnitude, we found no significant group differences for  $\phi$ . Possibly, this could be related to the use of mannitol solution as an oral contrast agent in patients but not in healthy controls and to the increased inflammation in active CD compared with our study with a rather fibrotic CD. We assumed a predominance of fibrotic over acute inflammatory changes in our CD patient collective as the indication for surgery was stricture in five patients and failed medical therapy in all patients, indicating a long-standing disease course. Desreumaux *et al.* observed a shift in the ratio of intra-abdominal fat to total body fat on MRI in CD patients compared to controls, indicating MAT hyperplasia in CD patients (26). As another approach to noninvasively quantify the presence of creeping fat, Li *et al.* developed a CT-based “mesenteric creeping fat index”, which indirectly assesses creeping fat by using vascular findings as a surrogate (25). The proposed index allowed the authors to predict the extent of creeping fat in surgical specimens in a prospective study population with an AUC of 0.76 (25). Moreover, in a study using CT-enterography, Sakurai *et al.* found that mesenteric findings such as hypervascularity (comb sign) and enlarged mesenteric lymph nodes, rather than mural findings, were highly correlated with the endoscopically determined severity of mucosal ulcerations in CD (44).

Many MRI studies (12,45–49) and recently also a few MRE studies (20,21,30) assessed the bowel wall and not MAT. For instance, a recent MRE study conducted in a patient population overlapping with the present study showed significantly increased stiffness and loss angle of the bowel wall in inflammatory bowel disease (20). Compared to the present study, there was a strong correlation of  $\phi$  between the bowel wall and adjacent MAT in CD patients ( $r=0.84$ ,  $P=0.001$ ), whereas no correlation was found for SWS ( $r=0.16$ ,  $P=0.64$ ). In another study, Avila *et al.* also reported increased stiffness in patients with CD as well as a link between advanced fibrosis on MRI and clinical events, e.g., surgical resection (21). Ultrasound-based elastography techniques have been used as diagnostic tools for the assessment of the bowel wall for more than two decades, however, without quantitative analysis of MAT (50).

Our study has limitations. First, we only investigated a small number of patients in this proof-of-concept study.

Second, all CD patients and healthy controls have been reported previously (20). However, there is no overlap of MRE data as the earlier study focused on the bowel wall in inflammatory bowel disease. Third, intestinal adhesions and pathologically altered movements in CD might have influenced the comparison with healthy controls. Furthermore, no spasmolytic agents were administered to enable comparison with healthy controls in this explorative setting. Fourth, we have not performed a test-retest reproducibility assessment. However, we have found a good to excellent interreader agreement and another study investigating the prostate using the same tomoelastography setup demonstrated a good test-retest reproducibility (51). Fifth, the BMI distribution in our cohort reflects normal adult weight, which prevents an in-depth analysis of the interaction between BMI and MRE parameters. Nevertheless, in this exploratory proof-of-concept study, it is beneficial to avoid high variability in BMI as a confounding factor. Sixth, a biometric power analysis could not be performed in this exploratory proof-of-concept study as this is the first study to investigate the MRE-based viscoelastic properties of MAT in CD and no prior values were available. However, the results of this study provide a basis for planning future prospective studies. Finally, although all patients underwent surgery the day after the MRE scan and CD was histopathologically confirmed, there was no uniform or quantified account available for the extent of creeping fat or for the extent of intestinal inflammation and fibrosis. Nevertheless, we were able to measure MAT directly adjacent to pathological intestinal lesions. Still, other factors, such as mesenteric inflammation and fibrosis, might have had an influence. Future studies should take this into consideration to further improve the correlation of quantitative parameters. Using intraoperative ultrasound elastography or tabletop MRE of surgical specimens would allow a direct comparison of the mechanical properties of creeping fat with histopathology.

## Conclusions

In this proof-of-concept study, we demonstrate the feasibility of *in vivo* MRE of MAT and present preliminary reference values for CD patients and healthy controls. Our results provide a basis for future study planning and motivate further studies for the biophysical characterization of MAT in CD in a bigger patient collective with histopathological scoring. Threshold values

of creeping fat would be desirable, possibly enabling predictions of early therapy response and the course of CD in clinical routine.

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## Footnote

*Reporting Checklist:* The authors have completed the STARD reporting checklist. Available at <https://qims.amegroups.com/article/view/10.21037/qims-23-41/rc>

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INC Research, Innate Pharma, InSightec Ltd., Inspiremd, InVentiv Health Clinical UK Ltd., Inventivhealth, IO Biotech ApS Copenhagen, IOMEDICO, IONIS, IPSEN Pharma, IQVIA ISA Therapeutics, Isis Pharmaceuticals Inc., ITM Solucin GmbH, Jansen-Cilag GmbH, Kantar Health GmbH (CRO), Kartos Therapeutics, Inc., Karyopharm Therapeutics, Inc., Kendle/MorphoSys AG, Kite Pharma, Kli Fo Berlin Mitte, Kura Oncology, Labcorb, La Roche, Land Berlin, Lilly GmbH, Lion Biotechnology, Lombard Medical, Loxo Oncology, Inc., LSK BioPartners, USA, Lundbeck GmbH, LUX Biosciences, LYSARC, MacroGenics, MagForce, MedImmune Inc., MedImmune Limited, Medpace, Medpace Germany GmbH (CRO), MedPass (CRO), Medtronic, Medtraveo GmbH, Merck, Merrimack Pharmaceuticals Inc., MeVis Medical Solutions AG, Millenium Pharmaceuticals Inc., Miltenyi Biomedicine GmbH, Bergisch Gladbach, miRagen Boukider, Mologen, Monika Kutzner Stiftung, MophoSys AG, MSD Sharp, Nektar Therapeutics, NeoVacs SA, Netzwerkverbund Radiologie, Neurocrine Biosciences Inc., US, Newlink Genetics Corporation, Nexus Oncology, NIH, NOGGO Berlin, Nord-Ostdeutsche Gesellschaft e.V., Novartis, Novocure, Nuvisan, Ockham oncology, Odonate Therapeutics San Diego, OHIRC Kanada, Oppilan Pharma Ltd., London, Orion Corporation Orion Pharma, OSE Immunotherapeutics, Parexel CRO Service, Pentixal Pharma GmbH Perceptive, Pfizer GmbH, PharmaCept GmbH, Pharma Mar, Pharmaceutical Research Associates GmbH (PRA), Pharmacyclics Inc., Philipps, Philogen s.p.a. Siena, Pliant Therapeutics San Francisco, PIQUR Therapeutics Ltd., Pluristem, PneuRX.Inc., Portola Pharmaceuticals, PPD (CRO), PRaint, Precision GmbH, Premier-Research, Priovant Therapeutics USA, Provectus Biopharmaceuticals, Inc., Psi-Cro, Pulmonx International Sarl, Quintiles GmbH, Radiobotics ApS, Regeneraon Pharmaceuticals Inc., Replimune, Respicardia, Rhythm Pharmaceuticals, Inc. Boston USA, Roche, Salix Pharmaceuticals Inc., Samsung, Sanofi, Sanofis-Aventis S.A., Sarepta Therapeutics, Cambridge, US, Saving Patient's Lives Medical B.V., Schumacher GmbH, Seagen, Seattle Genetics, Servier (CRO), SGS Life Science Sercives (CRO), Shape Memorial Midical Inc., USA, Shire Human Genetic Therapies, Siemens, Silena Therapeutics, SIRTEX Medical Europe GmbH, SOTIO Biotech, Boston, Spectranetics GmbH, Spectrum Pharmaceuticals, Stiftung Charite/BIH, St. Jude Medical, Stiftung Wolfgang Schulze, Syneos Health UK, Ltd., Symphogen, Taiho Oncology, Inc., Taiho Pharmaceutical Co., Target Pharma Solutions Inc., TauRx

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*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The presented study is not registered in the clinical trial platform. The study was approved by the Institutional Review Board of Charité – Universitätsmedizin Berlin [protocol code (A1/075/17) and date of approval (5-11-2017)] and was conducted in accordance with the Declaration of Helsinki (as revised in 2013). Written informed consent was obtained from patients.

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