Cine MRI assessment of motility in the unprepared small bowel in the fasting and fed state: Beyond the breath-hold

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\textbf{Abstract}

\textbf{Background:} The symptoms of functional bowel disorders are common in postprandial but investigations are generally undertaken in the fasted state using invasive procedures. MRI provides a noninvasive tool to study the gastrointestinal tract in an unperturbed, fed state. The aim of this study was to develop a technique to assess small bowel motility from cine MRI data in the unprepared bowel in fasting and fed states.

\textbf{Methods:} Fifteen healthy volunteers underwent a baseline MRI scan after which they consumed a 400 g soup. Subjects then underwent a postprandial scan followed by further scans at regular intervals. Small bowel motility was assessed using single-slice bFFE cine MRI. An optimized processing technique was used to generate motility data based on power spectrum analysis of voxel-signal changes with time. Interobserver variability (n = 15) and intra-observer (n = 6) variability were assessed. Changes in the motility index were compared between fasted and immediate postprandial state.

\textbf{Key Results:} Excellent agreement between observers was seen across the range of motility measurements acquired, with intraclass correlation coefficient (ICC) of 0.979 (P < 0.0001) and Bland-Altman limits of agreement 95% CI: –28.9 to 45.9 au. Intra-observer variability was low with ICC of 0.992 and 0.960 (2 observers, P < 0.0001). Changes from the fasted to immediately postprandial state showed an average increase of 122.4% ± 98.7% (n = 15).

\textbf{Conclusions & Inferences:} This optimized technique showed excellent inter and intra observer agreement. It was sensitive to changes in motility induced feeding. This technique will be useful to study contractile activity and regional patterns along the gastrointestinal tract under physiological conditions.

\textbf{KEYWORDS}
cine MRI, fasting, fed state, gastrointestinal motility, MRI

Abbreviations: AUC\textsubscript{power spectrum}, Area under the power spectrum; ICC, intra-class correlation coefficient; MRE, magnetic resonance enterography; ROI, region of interest; SD\textsubscript{Jac}, standard deviation of the Jacobian.
1 | INTRODUCTION

Conventional manometry of small bowel motility has provided valuable insights into motor function pathophysiology of the gastrointestinal tract.1,2 The technique has limitations, with naso-duodenal or oro-duodenal intubation being a difficult, uncomfortable, and invasive procedure for patients. Moreover, manometry techniques are not generally used in the lower sections of the small bowel due to difficulties with access and the invasiveness of the technique.3,4 Furthermore, misinterpretation of the manometry recordings can occur if nonocclusive contractions occur and large spacing between ports mean that motor patterns can be mis-defined.5 The tube may also interfere with normal feeding making it particularly difficult to study physiological changes from the fasting state and the effect of nutrient intake.

Over the last 10 years, MRI has proven to be a useful tool to probe the unprepared physiology of the gastrointestinal tract.6-9 It is particularly suitable for longitudinal or repeated studies, and its versatility allows for multiple physiological parameters to be monitored in a single scanning session. Magnetic resonance enterography (MRE) is used to evaluate the small bowel after the ingestion of an oral contrast agent. It involves distending the bowel artificially to produce detailed images of the bowel wall10 and induces bowel wall motion to move the large amount of oral contrast agent through the GI tract, which can then be studied using cine MRI.

Motility measurements following oral contrast preparation using a cine MRI acquisition have made significant advances in recent years.8,11-17 but to date quantification of wall motion has either involved looking for contractions across the lumen13,14,16 or using registration methods,15,17 which work well in the deliberately distended bowel where the walls are clearly visualized. However, bowel distension with a hypo-osmotic solution may not be truly physiological and so cannot study true fasting motility patterns and may not represent the full range of motility patterns in the postprandial state. The ability to study motility in the postprandial state, or the transition between the 2 states, has many potential advantages in furthering our understanding of physiology and the origin of symptoms which many patients experience after feeding. Moreover, it is particularly important for the pharmaceutical sciences because the rate and extent of drug dissolution and absorption from solid oral dosage forms is highly dependent upon gastrointestinal motility.18,19 Furthermore, the use of bowel distension limits the use of MRE in pediatric and elderly populations.

The unprepared small bowel can be imaged using the same high spatial and temporal resolution cine acquisitions as for the prepared bowel11,12,14. However, the postprocessing techniques used to parameterize the motility may need to be refined for images, which do not delineate the small bowel wall clearly and show different patterns of motility.

The aim of this study was to develop an analysis technique to assess motility from cine MRI data acquired in the fasted and fed, unprepared small bowel. Inter- and intra-observer variability, and the sensitivity to changes in motility caused by feeding were investigated.

2 | MATERIALS AND METHODS

This study was approved by the local Ethics Committee of the University of Nottingham (H19062014). This study is registered on www.clinicaltrials.gov with identifier NCT02717117. All subjects gave informed written consent. The study design, subjects, and data sets used have been reported previously.20

2.1 | Unprepared bowel data acquisition

Fifteen healthy volunteers (age 29 ± 10 years, BMI 24 ± 5 kg/m²) were recruited from the local campus population. Subjects with any disease or taking medication (eg loperamide, codeine, metoclopramide, hyoscine butylbromide, mebeverine, ondansetron) that affects gastric emptying or small bowel transit were excluded. Standard MRI exclusion criteria were applied.

This study was open label. Subjects were scanned using a 1.5T Philips Achieva MRI scanner (Philips Healthcare, Best, the Netherlands) using the 16 element torso (XL-TORSO, Philips Healthcare) coil at the Sir Peter Mansfield Imaging Centre, University of Nottingham. They underwent a baseline fasting scan defined as \( t = -20 \) minutes time point. They were asked to consume a soup meal (cream of chicken soup (400 g) (or mushroom for vegetarians) (Heinz, Wigan, UK) within 20 minutes then the subjects underwent a first immediate postprandial scan (defined as \( t = 0 \) minutes). MRI data collection was subsequently repeated every 15 minutes for the first 60 minutes where the subjects remained in the scanner and then every 30 minutes up to 270 minutes where subjects were allowed to leave the scanner between scans if they chose to.20 The subjects
were scanned using a range of sequences. At each timepoint, scans were acquired to assess small bowel motility using a single slice bTFE cine MRI acquisition (with reconstructed in-plane resolution $1.49 \times 1.7\, \text{mm}^2$, slice thickness 10 mm, echo time (TE) $= 1.52\, \text{ms}$, repetition time (TR) $= 3.0\, \text{ms}$, flip angle 80º, SENSE 2.0), of 1 minute free-breathing, temporal resolution of 1 s, this was repeated at 6 contiguous parallel coronal planes through the small bowel as previously described. The total scan time for motility was 6 minutes. The subjects were instructed to take shallow gentle breaths for the duration of the motility acquisition.

2.2 | Data analysis

2.2.1 | Motility assessment

Free breathing data were processed using GIQuant (Motilent, Ford, UK). The algorithm corrects respiratory motion before applying the nonlinear optic flow registration as described previously to correct local deformation caused by bowel wall motion and model intensity changes caused by luminal flow. The data output from the image registration were further analyzed using a customized graphical user interface written in MATLAB (MathWorks, Natick, MA).

On MRI, the unprepared small bowel has a very different appearance to the prepared bowel required for MRE (Figure 1). In the prepared bowel, there is clear definition of the bowel walls and obvious peristaltic motion is visible through the time series across most of the small bowel. In the unprepared bowel, the bowel wall is not always visible and bolus movement of the chyme between segments is common postprandially. Therefore, a different approach for quantifying the motility of the unprepared small bowel was developed, based on the registration parameter $C$ which represents the change in signal intensity between timepoints, within a defined region of interest (ROI) placed over the small bowel loops. This parameter is modeled simultaneously with the deformation during the registration process and is intended to capture any signal intensity changes not occurring from in-plane motion (ie through plane motion and flow).

To allow sensitivity to both oscillatory events such as mixing of contents during peristalsis and forward propulsion of boluses of chyme, the power spectrum analysis of the image registration parameter $C$ was developed similar to that proposed by van der Paardt et al and Sprengers et al. Initially, to remove zero frequency data, the mean of $C$ through time for each pixel was calculated and subtracted from each pixel value (Figure 2A). Then for each pixel in the data, the power spectrum (Fourier transform of time course of mean-subtracted $C$, smoothed to reduce noise and with the first timepoint removed to eliminate data not in the steady state, and then multiplied by its complex conjugate) was calculated (Figure 2B), generating data with frequency information up

![Image](https://via.placeholder.com/150)

**FIGURE 1**  Figure illustrating the difference between (A) the prepared bowel with clear definition of the bowel wall and bright luminal contents (data not from the current study, bowel preparation of 2% mannitol with 0.2% locust bean gum) and (B) the unprepared bowel with less visible bowel wall and brighter contents only when chyme moves into the segment

**FIGURE 2**  (A) Graph showing the variation in intensity of the $C$ parameter with time for 3 abdominal regions. Solid black line is a small ROI from the upper small bowel (jejunum), dashed black line is a small ROI from the lower small bowel (ileum), solid gray line is a small ROI from the ascending colon representing a known low motility region of the GI tract. (B) Corresponding power spectrum of the data in A.
to 0.48 Hz. The area under the power spectrum was calculated as a summary metric (AUC\_power\_spectrum), and maps created to visualize the regions of higher motility. This metric was intended to reflect both segmental oscillations and bolus movement of contents, typically seen postprandially.\(^{25}\) When regions of interest were defined, average data for the ROI was calculated from the pixel by pixel measurements within the ROI of the AUC\_power\_spectrum maps.

### 2.3 | 2.2.2. Observer variability

To determine the variability in the results due to observer definition of the region of the small bowel loops, the following analyses were carried out.

1. **Interobserver variability**: 2 observers, 1 experienced (CH over 10 years) in viewing small bowel MRI data and 1 inexperienced (AK <2 years) drew regions around all the visible small bowel segments across the 6 coronal slices acquired. This was repeated for all 15 subjects scanned at all timepoints pre- and post-test meal. The number of regions depended on the spatial separation of the different bowel loops. If there was a large amount of visceral fat separating the loops, more regions were drawn to encompass all the small bowel loops.

2. **Intra-observer variability**: The same 2 observers drew regions around the visible small bowel segments from only 6 subjects (chosen to have different body composition: 3 normal BMI (22.3, 22.9, 22.9 kg/m\(^2\)) and 3 high BMI (30.6, 27.1, 29.1 kg/m\(^2\)). The changing body composition resulted in images with very different contrast of the edges of the small bowel loops and therefore represented the maximum range of tissue contrast that would be seen across all subjects. Regions were defined by drawing the ROIs to encompass only the visible small bowel loops ignoring intra-abdominal visceral fat and other tissues; the regions were defined twice on the images with at least 1-month interval between observations.

### 2.3.1 | Changing motility in response to feeding using 2 motility metrics

We examined the strength of correlation between 2 motility analysis techniques using the total power (AUC\_power\_spectrum) and the standard deviation of the Jacobian (SD\_JAC, a previous published metric for motility\(^{3}\)). SD\_JAC looks at the geometric changes from image registration and is currently used for small bowel motility in the prepared bowel. In addition, we showed how both metrics change with feeding by looking at the mean change in the metric between the fasted (t = −20 min) and the immediately postprandial data (t = 0 min).

### 2.4 | Statistical analyses

All statistical analysis was carried out using Graph Pad Prism 7.0 (La Jolla, CA). All data were tested for normality using the D’Agostino and Pearson’s normality test. Interobserver variability was investigated using a Bland-Altman plot to determine the 95% confidence limits of agreement. Correlation between observers was measured using the Intra-class correlation coefficient (ICC) using a 2-way random effects model, with a single rater and absolute agreement.\(^{26}\) Intra-observer variability was also investigated with Intra-class correlation coefficients using a 2-way mixed effects model and single rater and absolute agreement. The 95% confidence limits of agreement were also calculated. Pearson’s correlation coefficient was used to measure the strength of correlation between the measurement of motility using AUC\_power\_spectrum and SD\_JAC for the fasting and immediately postprandial data sets.

### 3 | RESULTS

#### 3.1 | Visualization of high motility regions

Example maps of AUC\_power\_spectrum of fasting and postprandial data are shown in Figure 3 for 1 subject. Example maps of SD\_JAC are also shown for comparison. The AUC\_power\_spectrum maps show lower values in regions of known low motility (eg liver) compared to the SD\_JAC maps. There also appears to be a larger change (AUC\_power\_spectrum 122.4\% ± 98.7%, SD\_JAC 31.8\% ± 20.7%) (calculated by dividing the fed-fasted data by the fasted data across all the 15 subjects) across the small bowel between the fasted and immediately postprandial states which suggests the AUC\_power\_spectrum maps may have a better range to define the differences between sporadic movements occurring during fasting and large scale movements following ingestion of the meal.

#### 3.2 | Interobserver variability of small bowel motility

The variation in both AUC\_power\_spectrum and SD\_JAC across different timepoints, averaged over all healthy volunteers at each timepoint, and covering all regions of small bowel from the 6 slices, measured by each observer is shown in Figure 4A (error bars shown are SEM). This graph shows low measured motility at baseline in the fasting state followed by a significant increase postprandially which then persists for the majority of the imaging period. The degree of correlation between 2 observers was assessed using the ICC to be 0.979 and P < 0.0001, n = 195 (Figure 4B). Interobserver variability was assessed using the Bland Altman plot\(^{27}\) (Figure 4C) which showed a mean difference of 8.5 au between small bowel motility measurements, with a 95% confidence interval of −28.9 to 45.9 au as indicated by the upper and lower dotted lines (Figure 4C).

#### 3.3 | Intra-observer variability of small bowel motility

The correlation between the 2 analyses for AUC\_power\_spectrum performed by each observer was also assessed using the ICC and Bland-Altman limits of agreement (Table 1), showing good...
FIGURE 3  Example of motility maps generated by the software for a single volunteer across the 6 slices acquired, visualizing the areas of high motility. A and B illustrate the fasting and the fed state motility maps. C represents the different motility maps generated by the area under the power spectrum (AUC_{power spectrum}) and SD_{JAC} motility parameters. S: slice number. Regions of small bowel have been highlighted on the images.
agreement between the analyses (ICC > 0.9 for all data). Figure 5 plots \( \text{AUC}_{\text{power spectrum}} \) measurements against time for the 6 subjects with normal and high BMI, showing good agreement between analyses at most timepoints for both body compositions.

3.4 | Changing motility in response to feeding

The subjects showed an increase between their fasting and initial postprandial motility measurement for both analysis methods (\( \text{AUC}_{\text{power spectrum}} \) 122.4% ± 98.7%, \( \text{SD}_{\text{JAC}} \) 31.8% ± 20.7%), although there was considerable spread within the data across the subjects (Figure 6). The correlation between the techniques was significant \((r = 0.9, P < 0.0001, n = 30)\).

4 | DISCUSSION

This study has described and evaluated an optimized technique for the analysis of gut motility from MRI images, specifically addressing the differences in appearance and motility of the small bowel wall in unprepared bowel MR images, compared to images obtained following bowel preparation with oral luminal contrast agent. The images in fasting and fed conditions showed poor small bowel wall definition in places. Coupled with the different motility patterns of bolus propulsion as well as peristalsis meant previously published analysis techniques, which are based on geometric changes over the measurement period to generate the motility metric, were not as appropriate to use in fasting and fed studies. Maps of \( \text{AUC}_{\text{power spectrum}} \) indicated a better discrimination of the bowel tissue compared to the more widely published \( \text{SD}_{\text{JAC}} \)\(^{8,15,28}\) with lower noise in the regions where non/low-motility is expected, and a bigger range of motility indices. The proposed method utilizes changes in signal intensities that occur when the small bowel contents move between segments in regions showing bolus movement of contents (duodenum and jejunum) as well as those exhibiting more oscillatory motion (ileum), rather than looking for continuous motion throughout the time series.

The appearance of the unprepared bowel, which may contain multiple collapsed loops which are not always easily identified, means the definition of the ROIs is more subjective than for the prepared bowel with distended lumen. However, our interobserver variability data suggested that ROI definition had a small effect on the results compared to the changes in motility seen following eating the meal. Excellent correlation between observers was seen across the whole range of \( \text{AUC}_{\text{power spectrum}} \) measurement acquired and the Bland-Altman limits of agreement were low compared to the range of values measured. The postprandial changes over time show a rapid increase in motility following ingestion of the soup meal with levels returning towards baseline much later after the meal had been consumed.\(^{20}\)

Intra-observer variability was also low with a small range of Bland-Altman limits of agreements for both observers across 2 very different body shapes. The ICC was excellent with all data >0.9.

|         | ICC   | Bland-Altman       |
|---------|-------|--------------------|
|         |       | MD, mean difference |
| Observer 1 | 0.992 | **Table 1** Table summarizing the \( \text{AUC}_{\text{power spectrum}} \) intra-class correlation coefficient (ICC) and Bland-Altman from observer 1 and observer 2 |
|         |       | 95% CI             |
|         |       | −7.2 to 19.4 au    |
|         |       | Measurement range (27.2–254.5 au) |
|         |       | **n** = 78         |
| Observer 2 | 0.960 | **MD** 1.0 au      |
|         |       | 95% CI             |
|         |       | −41.2 to 43.19 au  |
|         |       | Measurement Range (28.4–281.0 au) |
|         |       | **n** = 78         |
This study presents an optimized analysis of MRI data to assess the motility of the unprepared fasting and fed small bowel. The MRI method proposed removes the need for intubating subjects, which can be a stressful procedure, often requiring fluoroscopy to place the catheter and rarely covering the entire length of the small bowel. Orocecal transit times can be measured using breath tests but these include changes due to gastric emptying and do not give information about the specific motor function of the small bowel but are an indirect measure of small bowel motility. Scintigraphy transit studies involving ionizing radiation do not provide information about the motor patterns seen in the small bowel. The proposed techniques (including further frequency analysis of the power spectra) will be useful to study the time scales of contractile activity and regional patterns along the gastrointestinal tract in health and
disease. Information on small bowel motility can be obtained in conjunction with mapping the bowel liquid pockets using MRI\(^2\) furthering understanding of the effects of motility on the fluid environment of the bowel. These combined insights could help with advancing in vitro/in vivo predictive dissolution studies of oral dosage forms under similar, undisturbed conditions.

The data from the normal and high BMI subjects would indicate that there is a potential for over estimating motility in the higher BMI subjects. These subjects all presented with high motility indices postprandially; however, only 1 of the 3 showed high baseline data. These larger motility indices may have been measured as regions of high signal intensity in the fat as it moves into and out of the imaging plane during respiratory motion. As this may not be fully corrected by the registration algorithm these movements will be interpreted as bowel motility. Further studies of higher BMI subjects is needed to understand the factors contributing to the larger motility metric measured and whether poor registration is a factor.

Other factors, which could also influence the signal intensity, are field inhomogeneities and metallic artifacts. To some extent, overall changes in image intensity across the image due to these factors are removed from the \(\text{AUC}_{\text{power spectrum}}\) analysis using the registration parameter C which models the signal changes and not the absolute values. An empty bowel has a different intensity to a filled bowel; however, movement of the contents either between loops or from 1 section to another show similar changes in intensity levels. Other meal contents should have similar motility patterns to the meal in this study, but would need investigating to determine whether the sensitivity is the same as the soup meal, particularly for a more solid meal, which may have a much lower signal intensity in the small bowel.

There were limitations to our study. Due to the time-consuming nature of drawing all the individual ROIs on the motility data the intra-observer repeated measurements were confined to just 6 subjects, not the full 15 available (used for the interobserver data). Drawing of ROIs for a single timepoint took around 5-10 minutes depending on the anatomy including the loading of each data set into the software. However, the intra-observer data were chosen from subjects who had very different small bowel anatomical appearances due to their differing BMIs, providing the observers with contrasting data for drawing the regions. Smoothing of the data before calculating the power spectrum reduces the effects of isolated poor mis-registration of the data. However, it will not eliminate the effects completely and these datasets will slightly overestimate the small bowel motility present.

In conclusion, this study describes an optimized analysis technique to evaluate small bowel motility in the physiological fasting and postprandial states using registered cine MRI datasets. This method showed excellent agreement between measurements of intra- and interobservers as well as showing the sensitivity of the technique to changes in motility induced by ingestion of a meal. Cine MRI scanning is available on most clinical scanners worldwide; therefore, future studies have real potential to translate and improve our knowledge of the small bowel environment in health and disease.

ACKNOWLEDGMENTS

This study was supported by the National Institute for Health Research Nottingham Biomedical Research Centre. The views expressed are those of the authors and not necessarily those of the National Health Service (NHS, UK), the National Institute for Health Research (NIHR, UK) or the Department of Health. This study was funded by a scholarship from the Kuwait University and from a research grant from the Nottingham University Hospitals’ Charity (PP-Gordon Moran-Nov14). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health. LM is supported in part by Award # HHSF223201510157C: “In Vivo Predictive Dissolution (IPD) to Advance Oral Product Bioequivalence (BE) Regulation” by the U.S. Food and Drug Administration (FDA); this manuscript represents the position of the authors and not necessarily that of the FDA.

CONFLICT OF INTEREST

The remaining authors have no competing interests. AM is the CEO of Motilent Limited, a medical imaging analysis company.

AUTHOR CONTRIBUTIONS

AK, CLH, LM, RCS, PAG and GWM designed the research. AK, AN recruited the participants. AK collected the data. AK, AM, and CLH, analyzed the data. AK, CLH, LM, and GWM wrote the manuscript draft. All authors revised the final manuscript.

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REFERENCES

1. Hansen MB. Small intestinal manometry. Physiol Res. 2002;51(6):541-556.
2. Malagelada JR, Stanghellini V. Manometric evaluation of functional upper gut symptoms. Gastroenterology. 1985;88(5):1223-1231.
3. Malagelada C, Malagelada JR. Small bowel motility. Curr Gastroenterol Rep. 2017;19(6):26.
4. Seidl H, Gundling F, Pfeiffer A, Pehl C, Schepp W, Schmidt T. Comparison of small-bowel motility of the human jejunum and ileum. Neurogastroenterol Motil. 2012;24(8):e373–e380.
5. Arkwright JW, Dickson A, Mauder SA, et al. The effect of luminal content and rate of occlusion on the interpretation of colonic manometry. Neurogastroenterol Motil. 2013;25(1):E52–E59.
6. Schwizer W, Steingoetter A, Fox M. Magnetic resonance imaging for the assessment of gastrointestinal function. *Scand J Gastroenterol*. 2006;41(11):1245-1260.

7. Fruehauf H, Menne D, Kwiatek MA, et al. Inter-observer reproducibility and analysis of gastric volume measurements and gastric emptying assessed with magnetic resonance imaging. *Neurogastroenterol Motil*. 2011;23(9):854-861.

8. Menys A, Taylor SA, Emmanuel A, et al. Global small bowel motility: assessment with dynamic MR imaging. *Radiology*. 2013;269(2):442-449.

9. Marciani L, Garsed KC, Hoad CL, et al. Stimulation of colonic motility by oral PEG electrolyte bowel preparation assessed by MRI: comparison of split vs single dose. *Neurogastroenterol Motil*. 2014;26(10):1426-1436.

10. Griffin N, Grant LA, Anderson S, Irving P, Sanderson J. Small bowel MR enterography: problem solving in Crohn’s disease. *Insights Imaging*. 2012;3(3):251-263.

11. Wakamiya M, Furukawa K, Kanasaki S, Murata K. Assessment of small bowel motility function with cine-MRI using balanced steady-state free precession sequence. *J Magn Reson Imaging*. 2011;33(5):1235-1240.

12. Ohkubo H, Kessoku T, Fuyuki A, et al. Assessment of small bowel motility in patients with chronic intestinal pseudo-obstruction using cine-MRI. *Am J Gastroenterol*. 2013;108(7):1130-1139.

13. Menys A, Atkinson D, Odille F, et al. Quantified terminal ileal motility during MR enterography as a potential biomarker of Crohn’s disease activity: a preliminary study. *Eur Radiol*. 2012;22(11):2494-2501.

14. Froehlich JM, Waldherr C, Stoupis C, Erturk SM, Patak MA. MR motility imaging in Crohn’s disease improves lesion detection compared with standard MR imaging. *Eur Radiol*. 2010;20(8):1945-1951.

15. Odille F, Menys A, Ahmed A, Punwani S, Taylor SA, Atkinson D. Quantitative assessment of small bowel motility by nonrigid registration of dynamic MR images. *Magn Reson Med*. 2012;68(3):783-793.

16. Bickelhaupt S, Froehlich JM, Cattin R, et al. Differentiation between active and chronic Crohn’s disease using MRI small-bowel motility examinations - initial experience. *Clin Radiol*. 2013;68(12):1247-1253.

17. Hahnemann ML, Nensa F, Kinner S, Gerken G, Lauenstein TC. Motility mapping as evaluation tool for bowel motility: initial results on the development of an automated color-coding algorithm in cine MRI. *J Magn Reson Imaging*. 2015;41(2):354-360.

18. Oberle RL, Amidon GL. The influence of variable gastric emptying and intestinal transit rates on the plasma level curve of cimetidine: an explanation for the double peak phenomenon. *J Pharmacokinet Biopharm*. 1987;15(5):529-544.

19. Oberle RL, Chen TS, Lloyd C, et al. The influence of the interdigestive migrating myoelectric complex on the gastric emptying of liquids. *Gastroenterology*. 1990;99(5):1275-1282.

20. Khalaf A, Hoad CL, Menys A, et al. MRI assessment of the postprandial gastrointestinal motility and peptide response in healthy humans. *Neurogastroenterol Motil*. 2017;31:e13182.

21. Moran GW, Leslie FC, McLaughlin JT. Crohn’s disease affecting the small bowel is associated with reduced appetite and elevated levels of circulating gut peptides. *Clin Nutr*. 2013;32(3):404-411.

22. Hamy V, Dikalos N, Punwani S, et al. Respiratory motion correction in dynamic MRI using robust data decomposition registration - application to DCE-MRI. *Med Image Anal*. 2014;18(2):301-313.

23. van der Paardt MP, Sprengers A, Zijta FM, Lamericis R, Nederveen AJ, Stoker J. Noninvasive automated motion assessment of intestinal motility by continuously tagged MR imaging. *J Magn Reson Imaging*. 2014;39(1):9-16.

24. Sprengers Andre MJ, van derPaardt MP, Zijta Frank M et al. Use of continuously MR tagged imaging for automated motion assessment in the abdomen: a feasibility study. *J Magn Reson Imaging*. 2012;36(2):492-497.

25. Silverthorn DU. *Human Physiology: an integrated Approach*. London: Pearson; 2012.

26. Koo TK, Li MY. A guideline of selecting and reporting intraclass correlation coefficients for reliability research. *J Chiropr Med*. 2016;15(2):155-163.

27. Bland JM, Altman DG. Statistical methods for assessing agreement between two methods of clinical measurement. *Lancet*. 1986;1(8476):307-310.

28. Plumb AA, Menys A, Russo E, et al. Magnetic resonance imaging: quantified small bowel motility is a sensitive marker of response to medical therapy in Crohn’s disease. *Aliment Pharmacol Ther*. 2015;42(3):343-355.

29. Geypens B, Bennink R, Peeters M, et al. Validation of the lactose-[C-13]uride breath test for determination of orocecal transit time by scintigraphy. *J Nucl Med*. 1999;40(9):1451-1455.

30. Maurer AH. Gastrointestinal motility, part 2: small-bowel and colon transit. *J Nucl Med Technol*. 2016;44(1):12-18.

31. Mudie DM, Murray K, Hoad CL, et al. Quantification of gastrointestinal liquid volumes and distribution following a 240 mL dose of water in the fasted state. *Mol Pharm*. 2014;11(9):3039-3047.

How to cite this article: Khalaf A, Nowak A, Menys A, et al. Cine MRI assessment of motility in the unprepared small bowel in the fasting and fed state: Beyond the breath-hold. *Neurogastroenterol Motil*. 2019;31:e13466. [https://doi.org/10.1111/nmo.13466]