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Measurement of gastric meal and secretion volumes using magnetic resonance imaging

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Abstract

MRI can assess multiple gastric functions without ionizing radiation. However, time consuming image acquisition and analysis of gastric volume data, plus confounding of gastric emptying measurements by gastric secretions mixed with the test meal have limited its use to research centres. This study presents an MRI acquisition protocol and analysis algorithm suitable for the clinical measurement of gastric volume and secretion volume. Reproducibility of gastric volume measurements was assessed using data from 10 healthy volunteers following a liquid test meal with rapid MRI acquisition within one breath-hold and semi-automated analysis. Dilution of the ingested meal with gastric secretion was estimated using a respiratory-triggered T1 mapping protocol. Accuracy of the secretion volume measurements was assessed using...
data from 24 healthy volunteers following a mixed (liquid/solid) test meal with MRI meal volumes compared to data acquired using gamma scintigraphy (GS) on the same subjects studied on a separate study day. The mean ± SD coefficient of variance between 3 observers for both total gastric contents (including meal, secretions and air) and just the gastric contents (meal and secretion only) was 3 ± 2% at large gastric volumes (>200ml). Mean ± SD secretion volumes post meal ingestion were 64 ± 51 ml and 110 ± 40 ml at 15 and 75 min, respectively. Comparison with GS meal volumes, showed that MRI meal only volume (after correction for secretion volume) were similar to GS, with a linear regression gradient ± std err of 1.06 ± 0.10 and intercept −11 ± 24 ml. In conclusion, (i) rapid volume acquisition and respiratory triggered \( T_1 \) mapping removed the requirement to image during prolonged breath-holds (ii) semi-automatic analysis greatly reduced the time required to derive measurements and (iii) correction for secretion volumes provided accurate assessment of gastric meal volumes and emptying. Together these features provide the scientific basis of a protocol which would be suitable in clinical practice.

Keywords: gastric volume, MRI, secretion

(Some figures may appear in colour only in the online journal)

1. Introduction

MRI is a very useful tool for studying both the structure and function of the stomach (Curcic et al 2010, Marciani 2011) as well as the mechanisms by which food is digested in and emptied from the stomach (Kwiatek et al 2009, Marciani et al 2007, 2012, 2013). A key advantage of MRI is that this imaging modality can acquire rapid measurements of multiple parameters of gastric function in a single scanning session without the use of ionizing radiation. Gastric emptying can be measured by the reduction in the volume of gastric contents over time on anatomical MRI scans (Schwizer et al 1992, Marciani et al 2000, 2001b, Steingoetter et al 2006, Fruehauf et al 2009), gastric motility can be assessed using cine imaging (Borovicka et al 1999, Marciani et al 2001c, 2005, Kwiatek et al 2006) and gastric secretion can be estimated by monitoring the dilution of the meal (Treier et al 2008, Goetze et al 2009, Sauter et al 2012). Based on these findings, gastric MRI has been proposed as a clinical investigation for the diagnosis of gastroesophageal reflux (Curcic et al 2014a), gastroparesis (Ajaj et al 2004) and functional dyspepsia (Tucker et al 2012), as well as to determine residual volumes in preoperative sedation ahead of surgery (Lobo et al 2009, Schmitz et al 2011, 2012). However, practical issues such as non-standardised meals, time consuming image acquisition and manual analysis protocols have restricted the use of gastric MRI studies outside specialist research centres. A key barrier to implementation in clinical practice is the time required to analyze the data, for instance the measurement of gastric volumes from the scans by manual or assisted-manual (Barrett and Mortensen 1996) drawing of the stomach content volume and air volume on every slice for every time point.

An additional complication is that gastric secretions within the stomach cannot be distinguished from the meal on standard MRI scans. Gastric secretion is highly variable between individuals and can complicate attempts to determine gastric meal volumes and, therefore,
gastric emptying and nutrient delivery to the small bowel (Kwiatek et al 2009). Gamma scintigraphy does not suffer from this problem as only the meal is labelled, but it cannot measure total gastric volume including intra-gastric air and secretions. This is important because total gastric volume has been closely linked to satiety scores of fullness in healthy individuals and patients with dyspepsia symptoms (Marciani et al 2001b, Treier et al 2008, Goetze et al 2009, Tucker et al 2012, Parker et al 2012c). A variety of MRI techniques can be used to assess gastric secretion independent of meal volume. Early work measured changes in the MR relaxation time ($T_2$) of locust bean gum solutions to estimate the amount of gastric secretions (Marciani et al 2001b). More recently estimates of gastric secretion volume in liquid meals have been made by adding Gadolinium contrast agent to the meal which substantially reduces the $T_1$ of the meal. This allows gastric secretion to be measured from the change in $T_1$ as the meal is diluted in the stomach (Treier et al 2008, Goetze et al 2009). However, published studies have used a dual flip angle sequence (and associated B1 map) to measure $T_1$ which required an excessively long breath hold not tolerated by many patients.

This study presents a rapid MRI acquisition protocol and analysis algorithm suitable for the practical measurement of gastric volume and secretion volume in clinical studies. Throughout we aimed to minimise acquisition and analysis time without sacrificing measurement accuracy. Acquisition time was kept to a minimum, with a single breath-hold scan for volume measurements and $T_1$ maps generated from a respiratory triggered acquisition that removed the need for prolonged breath holds, and subsequent recovery time. Semi-automatic volume analysis software was used to greatly reduced the time required to measure gastric volumes (Total Gastric Volume (TGV), gastric content volume (GCV) and intra-gastric air volume) and was applied to a study of the fate of a liquid meal swallowed by healthy volunteers. Reproducibility was assessed by the co-efficient of variance of the volumes measured by 3 observers. Measurement of gastric secretion and gastric meal volumes were validated by comparison of MRI measurements with gastric scintigraphy measurements that include only the labelled meal.

2. Materials and methods

An illustration and explanation of the terms used in this study and a flow diagram of the different meals and measurements taken are shown in figure 1.

2.1. Meals

2.1.1. Liquid test meal  This meal was used in the assessment of total gastric volumes, gastric content volumes and intra-gastric air volumes. The meal was made up of 200 ml vanilla fortisip (Nutricia Clinical) and 200 ml water (300 kcal, 11.6 g fat). To this 0.4 ml of paramagnetic contrast agent was added (0.5 mmol l$^{-1}$ Gd-DOTA; Dotarem®, Laboratorie Guerbet, Aulnay-sous-Bois, France) to increase the contrast between the meal and the surrounding tissues and to measure dilution (Parker et al 2012b).

2.1.2. Mixed test meal  This meal was used in the assessment of secretion and meal only volumes. This contained the 400 ml liquid test meal (above) with 12, 1% food grade agar beads (Marciani et al 2001a) (Agar-agar; Cuisine-innovation: Dijon, France), which adds no nutrient value to the meal. 7.0 g of barium sulphate per 100 ml of agar solution (E-Z Paque:
1370
Buckinghamshire, UK, Ph Eur 96% w/w) was added to the agar beads to increase the density and prevent floating within the liquid meal (Parker et al 2012a).

2.1.3. Meal ingestion After baseline images had been acquired, at time $T = -10 \text{ mins}$, volunteers drank 200 ml of the liquid test meal at a rate of 100 ml min$^{-1}$ and were then imaged ($T = -5 \text{ min}$). The remaining 200 ml of the test meal was consumed at a rate of 100 ml min$^{-1}$. If volunteers were consuming the mixed meal the agar beads were swallowed with the second 200 ml of liquid meal at a rate of 3 beads per 50 ml. Regular imaging commenced at $T = 0 \text{ mins}$.

2.2. Study population

Both liquid and mixed meal studies were approved by the local research ethics committee (10/H0408/52 liquid meal, 12/EM/0114 Mixed Meal). All volunteers gave written informed consent. Healthy volunteers had no history of gastro-intestinal diseases and were suitable for MRI scanning. All subjects abstained from alcohol and strenuous exercise for 24 h prior to each study day and arrived at the test centre after an overnight fast. 10 healthy volunteers (5 male, mean age 22, range 19–26 years) consumed the liquid test meal and underwent the gastric volume measurement protocol (given below). Data were used to assess the fast analysis algorithm for measuring gastric volume (TGV, GCV and intra-gastric air). A further

Figure 1. (a) A representative image of the stomach is presented with (b) an explanation of the terms used in this study and (c) a flow diagram of the different meals and measurements taken in the volume and secretion studies.
24 healthy volunteers (male, mean age 48, range 19–69) consumed the mixed test meal and underwent both the gastric volume and secretion protocols. Data were used for assessing gastric secretion.

2.3. MRI study protocol

Imaging was carried out using a Philips 1.5 T Achieva scanner with a 16 channel XL Torso Coil placed over the abdomen. Gastric volumes were determined from transverse balanced turbo field echo (bTFE) scans covering the stomach with 50 slices of 5 mm thickness, no slice gap, in-plane resolution 2.0 × 1.77 mm², FOV 400 × 320 mm², TE/TR 1.5/3.0 ms, SENSE 2.0, FA 80°, data acquired in a short 16 s breath hold. Thin slices were used to reduce partial volume effects and a high flip angle was used to give good contrast between the fluid contents of the stomach and surrounding walls. Secretion volumes were estimated from T₁ maps generated from a series of respiratory-triggered IR-EPI acquisitions (13 TIs 50–1000 ms) (Cox et al 2011) acquired with 5 slices of 8 mm thickness with slice gap of 5 mm, in plane resolution of 3 × 3 mm², matrix size 112 × 112, SENSE factor 2.0, half scan factor 0.625, TE = 31 ms, TR minimum = 3000 ms. This method of T₁ mapping allows data to be acquired relatively quickly without breath-holding. Triggering of the inversion pulse was altered with an additional variable delay to allow a range of TIs, whilst acquiring the images at the same time point within the late expiration phase of the respiratory cycle. Total acquisition time for this sequence ranged between 39–65 s depending on the length of the respiratory cycle of each volunteer.

Scans to measure gastric volume were carried out before feeding, and at T = −5, 0, 5, 10, 15, 30, 45, 60, 75, 90, 120 min after the whole meal had been consumed. Volume data at 60 min was only acquired after the mixed meal, and 3 subjects who consumed the liquid only meal did not have a scan at 45 min due to the constraints caused by interleaving of 2 subjects during acquisition. T₁ maps to measure gastric secretion were acquired at 15 and 75 min after the meal.

2.4. In vitro dilution calibration

To calibrate dilution against meal T₁, the T₁ of the 400 ml liquid test meal was measured during sequential dilution with simulated gastric secretions at 37 °C (Rayment et al 2009). This data was then fitted to the following equation to convert T₁ into relative concentrations of meal and secretion.

\[
\frac{1}{T_1} = \frac{1}{aC_{gd} + T_{1GS}} + (cC_{gd} + d)C_{gd}
\]

where \(C_{gd}\) is the concentration of Gd-DOTA in µM and \(T_{1GS}\) the modelled \(T_1\) of pure gastric secretion (Treier et al 2008), and \(a\), \(c\) and \(d\) are fitted parameters related to the baseline concentration of contrast agent, concentration dependence of the relaxivity of the contrast agent, and relaxivity of the undoped component of the meal.

3. Image analysis

3.1. Gastric volumes

Software was written using IDL® 6.4 (Research Systems Inc., Boulder, Colorado, USA) to allow fast processing of the data to determine the content and intra-gastric air volume of the
gastric lumen (Parker et al. 2012b). The algorithms used to define the contents and intra-gastric air (and hence total volume: intra-gastric air + contents) are described below.

3.1.1. Gastric contents volume (GCV). The observer started the analysis using data from a time point when the stomach was most full (e.g. between $T = 0$ and 30 min). The observer defined a ‘seed’ point in the bright signal corresponding to the liquid phase of the meal and then moved a slider to define a minimum signal threshold. All voxels above this threshold connected to the seed point in the slice were used to define a binary mask. A closing filter (dilate then erode) of $3 \times 3$ pixels was applied to this mask to fill any small holes within the region. The edge of the region was then displayed, allowing the observer to interactively refine the threshold to define the contents (see figures 2(a) and (b)). The final threshold level was saved and then seed points in the remaining slices including stomach contents were defined either manually (observer clicking) or automatically (new seed generated from centre of mass of previous slice’s mask, selected by the observer). After this had been repeated for all the slices showing stomach contents, three types of editing were applied as necessary.

(a) Refining spatial limits. If the mask of the stomach had ‘leaked out’ where the stomach wall was very thin then the observer could draw a limit line which stopped the mask from extending in that direction (figures 2(c) and (d)).

(b) Filling a hole. If there was a darker region within the stomach content which was not included within the mask (e.g. solid agar bead components or poorly mixed meal) then the observer could click within the hole and all the region inside the hole became part of the mask (figures 2(e) and (f)).

(c) Adding at the edge. If the threshold levels set did not include all the stomach contents at the edges, probably due to susceptibility artefacts, then the observer could draw on the correct edge and fill the corresponding hole created (figures 2(g) and (h)).

Additional editing was generally required only on a few slices per time point and this was not time-consuming since each hole was filled by a single ‘click’ and limits and edges were set with a ‘press-move-release’ action. After editing, the gastric contents were completely defined.

3.1.2. Volume of intra-gastric air. The volume of intra-gastric air was defined using a similar method as for the contents except that a maximum signal threshold rather than a minimum signal threshold was used to define the dark area corresponding to air.

Finally, to ensure that pixels at the air/content boundary were handled properly, any voxel lying between the 2 masked regions in the vertical direction was assigned to the appropriate mask based on its signal intensity. Mean intensity levels of each region (intra-gastric air and contents separately) were calculated and the voxel was then assigned to the region having the closest intensity. The total number of voxels in each region in all slices were then calculated and converted to a volume using information on the voxel resolution.

Total Gastric Volumes (TGV) were computed from the sum of the gastric content and air volumes.

For each volunteer the initial threshold levels were set according to the protocol above on the first data set analysed. These threshold levels (from the first data set analysed) were used for subsequent data sets of the same volunteer to increase the speed of analysis. However, occasionally, when the stomach was nearly empty, the signal intensity in the gastric contents became more heterogeneous and less bright, as the milk protein within the meal separated in the acidic environment and it was necessary to reduce the threshold intensity level for the accurate description of the contents to be completed.
Figure 2. Illustrations of the post-processing analysis algorithm for measurement of gastric volumes. (a) First seed point is positioned in the stomach (black plus sign). (b) Minimum threshold level correctly set so that edges (white line) of the mask defined are at the edges of the stomach contents. (c) Example of mask ‘leaking out’ of stomach (white arrows). (d) Example of observer defined correction for ‘leaking out’ by setting a limit on the slice (solid black line). (e) Example of a hole in the mask (long white arrow, black box). (f) Hole has been filled by clicking inside it (black cross). (g) Example of edge missing from mask defined (white arrow head). (h) Observer has corrected edge (solid black line) by adding the edge in the correct place.
3.1.3. Gastric secretions. Maps of the $T_1$ of the stomach content were generated on a voxel-by-voxel basis by a 3-parameter fit to the inversion recovery model

$$M_{TI} = M_0 \left( 1 - \alpha \exp^{-\frac{TI}{\tau}} \right)$$

using the Powell minimisation algorithm (figures 3(a) and (b)), where $M_{TI}$ is the signal at inversion time $TI$, $M_0$ the equilibrium magnetisation and $\alpha$ is a parameter that takes account of the degree of inversion of the magnetization. $R^2$ was also calculated for each voxel to determine the goodness of the fit. Images at long $TI$ were used to visualise the stomach, the edge of the stomach was then defined manually by the observer, on all the slices. A histogram of $T_1$ values within the region for voxels with an $R^2$ of >0.8 was calculated with a bin size of 20 ms (figure 3(c)). Using a look up table of the data from the *in-vitro* calibration experiment each histogram bin was assigned to a percentage of meal and secretions (100–meal %); e.g. 220–240 ms: 100% meal, 0% secretions; 460–480 ms: 50% meal, 50% secretion). The total volume of contents measured at that time point was then used to estimate the volume of meal ($V_{\text{meal}}$) and volume of secretions ($V_{\text{sec}}$) using the following equations

$$V_{\text{meal}} = \sum_{i=0}^{n} c_i \cdot \frac{V \cdot P_i}{100 \cdot N}$$

*Figure 3.* (a) Example of raw IR-EPI images at increasing $TI$ times for a single slice through the main body of the stomach. (b) $T_1$ map of the corresponding slice shown in (a), with stomach region defined (white line). (c) Histogram of stomach region showing distribution of $T_1$ data from all slices acquired. Note the inhomogeneous distribution of secretions in the stomach. There is a layer of secretion (i.e. highly dilute gastric contents) that is clearly visible above the Gd-labelled meal on (b) with the undiluted meal represented by a narrow peak in the distribution of $T_1$ data in the histogram (c) and the dilute region represented by the visible plateau beyond the peak.
\[
V_{sec} = \sum_{i=0}^{n} c_i \cdot \frac{V \cdot (100 - MP_i)}{100 \cdot N} 
\]

where \(c_i\) is the number of counts in histogram bin \(i\), \(V\) is the total volume measured at that time point from the bTFE images, \(n\) is the number of bins in the histogram, \(P_i\) is the percentage of meal corresponding to histogram bin \(i\) and \(N\) is the total number of counts in the histogram. For \(T = 15\) min, \(n = 100\) corresponding to a maximum \(T_1\) of 2000 ms and for \(T = 75\) min, \(n = 200\) corresponding to a maximum \(T_1\) of 4000 ms.

3.1.4. Inter-observer reproducibility. To determine the variability of the volumes measured by different observers, 3 observers analysed all volume data from 10 healthy volunteers who had consumed the liquid test meal. Coefficient of variance for each volume measured was calculated and plotted against the mean volume measured by all three observers. This was done for TGV, GCV and intra-gastric air separately. Generation of the \(T_1\) map and subsequent estimated meal and secretion volumes were compared between 2 observers using Bland-Altman plots (Bland and Altman 1999) for all data from the 24 healthy volunteers who had consumed the mixed meal.

3.1.5. Accuracy of secretion estimations. To determine whether the estimations of secretions were reasonable, the meal volumes were compared to Gamma Scintigraphy (GS) data obtained from the same subjects who ingested the same meal with a radionuclide tracer (on a separate occasion). The difference between gastric contents volume measured by MRI and GS provides an estimate of secretion volume because meal volumes calculated from MRI include secretion and those from GS do not. 0.5 MBq of the non-absorbable marker In-111-indium chloride was added to the liquid component of the meal and 5 MBq Tc-99m-MAA (Technescan® LyoMAA (DRN4378), Mallinckrodt Medical B.V.,The Netherlands) to the agar beads. For imaging, radioactive markers were fixed to the subject at the right costal margin, both anteriorly and posteriorly. Subjects stood in front of a Mediso Gamma Camera (Nucline X-Ring-R, Budapest, Hungary) and a 30 s acquisition of anterior and posterior images were acquired. The first 200 ml of the liquid test meal was given at 100 ml min\(^{-1}\) and the subject was imaged \((T = -5\) min scan). The remaining liquid meal was then given at 100 ml min\(^{-1}\) with 12 agar beads swallowed whole (3 beads per 50 ml). This two-stage technique allowed the In-111 overlap with the Tc-99 on the GS images to be corrected.

Meal volumes from GS at the same time points \((T = 15\) and \(T = 75\) min) were compared to the MRI volumes (meal + secretions) as well as the meal only estimated volumes. Linear regression using data from both time points was used to compare the MR estimates of the meal volumes with and without secretions, to the volumes obtained from GS.

4. Results

4.1. Gastric volumes

The semi-automatic algorithm for measuring stomach contents and air volumes required the observer to make far fewer mouse ‘clicks’ per image compared to manual, or assisted manual drawing. For a slice which required no editing, the air and contents could be defined by just 2 clicks compared to at least 7–10 per region for manual drawing. This reduced observer input, allowed faster processing, with typical times of 5–10 min to complete the volume analysis for the first volume (where threshold levels are set) for a stack of images covering the entire
stomach; the timing depends on the amount of additional editing needed. For subsequent volumes where no threshold levels need to be set a typical dataset (contents and air) could be defined in 2–3 min. This results in a total processing time of 30–40 min for the full 12 time points.

The percentage coefficient of variance of volume measurements made by 3 observers can be seen in figure 4. The larger volumes had smaller coefficient of variance, since the differences between observers was generally at the very edges of the masks which has the smallest relative effect for large volumes. For content and total gastric volumes over 200 ml the mean ± SD percentage coefficient of variance was 3 ± 2% for TGV and 3 ± 2% for GCV. Volumes under 100 ml showed much larger percentage coefficient of variance as indicated on figure 4.

4.2. Gastric secretions

Data from the in-vitro dilution experiments are shown in figure 5. The fitted parameters of equation (1) for the dilution of the meal with simulated gastric secretions are as follows:

\[ a = -1.4 \cdot 10^{-4} \text{ms} \cdot \mu \text{M}^{-1}, \quad c = 2.1 \cdot 10^{-8} \text{ms}^{-1} \cdot \mu \text{M}^{-2}, \]

\[ d = 1.7 \cdot 10^{-3} \text{ms}^{-1} \cdot \mu \text{M}^{-1}. \]

\( T_1 \) maps were successfully generated in all 24 subjects. Data were subsequently excluded in 5 data sets (N = 3 at T = 15 and N = 2 at T = 75 min) due to poor positioning of the slice stack in the antrum. A further 3 data sets from T = 75 mins were also excluded due to poor fitting of the data. The mean ± SD number of voxels included in the histograms at T = 15 mins was 2042 ± 640 and at T = 75 mins it was 1124 ± 461. The \( R^2 \) constraint used to exclude voxels reduced the number of voxels by 18 ± 9% for the T = 15 min data and 25 ± 14% for the T = 75 min data, predominately from the edges of the regions, where wall motion generated errors in the data. The mean ± SD secretion volume at 15 min was 64 ± 51 ml and increased to 110 ± 40 ml by 75 min. Figure 6(a) shows that the total amount of secretion increased between the 2 time points for all subjects except one and the average amount of increase in secretion volume was 52 ± 29 ml. The two individuals with high levels of secretion at the earlier time point maintained this high level of secretion later.
4.3. Accuracy of secretion estimations

The comparison of GCV and meal only volumes measured using MRI with gastric content volumes measured with GS is shown in figure 6(b) and mean values for the 2 time points are given in table 1. There was close agreement between MRI and GS meal volumes when the estimated volume of secretions was removed from the gastric contents volume data. The linear regression gradient ± std err was 1.06 ± 0.10 for the meal only and GS meal volumes compared to 0.85 ± 0.10 for the gastric contents.
4.4. Inter-observer variability

The inter-observer variability for gastric secretion measurements assessed using Bland-Altman plots, can be seen in figure 7. For the estimated secretion volumes there was a mean difference of −1 ml (CI −11 to 9 ml) between observers for $T = 15$ min and −1 ml (CI −16 to 14 ml) for the $T = 75$ min time points.

5. Discussion

This paper reports the development and validation of a method for the rapid and reliable assessment of gastric meal and secretion volumes from magnetic resonance imaging (MRI). The findings demonstrate rapid secretion after consumption of this nutrient liquid test meal that continues throughout the emptying process such that secretion contributes more than half the volume of the gastric contents 75 min after ingestion of this meal.

The algorithm proposed to measure gastric volumes significantly reduced the user input compared to manual drawing, even when manual drawing is assisted by a live-wire edge detection. Typical times for analysis for manually segmenting this data would be of the order of 3–4 h, compared to 30–40 min using the semi-automated method. This allows data to be processed much faster. Reproducibility of results were maintained with minimal differences between results between three observers for gastric volumes exceeding 200 ml, although the

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<th>Mean volume ± SD/ml</th>
<th>Linear regression and correlation with GS data</th>
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| GCV MRI             | 402 ± 58     | 202 ± 50     | 0.85 ± 0.10 | 121 ± 25 | 0.808b |
| Meal only           | 338 ± 50     | 92 ± 36      | 1.06 ± 0.10 | −11 ± 24 | 0.867b |
| volume MRI$^a$      |               |              |             |          |        |
| Meal volume GS      | 310 ± 32     | 112 ± 61     |             |          |        |

$^a$ This data has been corrected removing the estimated secretion volume from the GCV.

$^b$ Significant to $p < 0.001$, $N = 40$.

Figure 7. Graphs showing Bland Altman plots of estimated meal volumes ($a$) and secretions ($b$) for 2 observers. $T_{15}$ shown as solid diamonds, $T_{75}$ shown as open squares. Differences in gastric contents volume measured following ingestion of a 400 ml meal were usually less than 10 ml.
variation between observers increased as the gastric content volume reduced. This finding is due to the fact that the majority of inter-observer variation comes from setting the initial threshold level which primarily affects the edges of the region defined. This has a greater relative impact for smaller than larger gastric volumes, although the absolute volume differences were never large. Similar trends were previously reported by Fruehauf and colleagues (Fruehauf et al 2011) where inter-observer variation for an alternative semi-automatic approach of gastric volumes was \(\sim 8\% \) at 400 ml, compared to \(\sim 3\% \) found for the semi-automatic method described here. It should be noted that the relatively high percentage variation in measurements of gastric volumes at low volumes (<100 mL) may not be critical in practice, because key metrics used to describe gastric function (e.g. half emptying time) are acquired early during gastric emptying. Moreover these measurements are derived from models that fit the entire emptying curve that can exclude or down weight later data points (Elashoff et al 1982, Kwiatek et al 2009). Further validation work will be needed to develop the software into a commercial package suitable for use in clinical practice.

This algorithm can also be applied to other liquid meals which show good contrast between the contents and stomach wall (Murray et al 2013). It can also be used for meals containing solids such as the agar beads used in this study; however the thresholding method does not always work well if there is a large range of intensity levels across the stomach (e.g. after ingestion of a normal mixed meal) in which case further user input may be required to define the volumes accurately (Sweis et al 2013). Recently there have been other semi-automatic algorithm proposed to segment intra-gastric contents and air by Banerjee et al (Banerjee et al 2014) and Bharucha et al (Bharucha et al 2014). The algorithm proposed by Bharucha follows a similar methodology to the one described in this work, as it uses both thresholding and morphological filtering with manual edits to achieve segmentation of the stomach and air and the semi-automated analysis was completed in similar time scales (<5 min per time point). However differences in the imaging sequences used to acquire data meant that residual gastric fluids present at baseline can also be assessed using the proposed segmentation algorithm as this fluid remained bright on the images, where it was dark for the Bharucha study. The Banerjee algorithm requires the observer to define a set of starting points throughout the volume of the stomach data which are at the boundaries of the contents and air. These starting points can be generated in several ways including thresholding (as used here). Once defined these points are used to define the edges of the regions automatically by using a live-wire boundary to connect adjacent points within each slice. This algorithm is also dependent on good contrast between the intra-gastric contents, air and the gastric walls. As a consequence it experiences similar problems to the algorithm proposed here when contrast is poor. Further automation of the Banerjee algorithm allowing for consecutive time points to be processed assuming minimal changes to the shape and volume of the stomach and contents. The temporal resolution and possibility of large changes in position of the stomach within the imaging stack from repositioning of subjects may make the Banerjee algorithm less appropriate for clinical studies.

The original method used to quantify dilution via the \(T_1\) of gadolinium doped meals required prolonged breath-holds (Treier et al 2008) which may not be practical for use with all patients, especially when combined with MRI volume scans which also require subjects to hold their breath. The rapid EPI sequence used in this study allows subjects to breath freely during the measurement of \(T_1\) potentially increasing the patient compliance and improving data quality in patients unable to hold their breath. An alternative method to measure both gastric volumes and gastric content \(T_1\) during free breathing has recently been proposed (Curcic et al 2014b), using a golden angle radial sequence. In this study the meal and secretion volumes were highly reproducible between observers at both time points after the automatic removal of
poorly fitted data at the very edges of the stomach where the quality of the $T_1$ fit was reduced due to wall motion. The errors in measuring $T_1$ from the observers are likely to be small when compared to other errors such as measurement reproducibility; however, these are hard to quantify as repeating measurements within the dynamics of gastric emptying is not appropriate. Treier et al (Treier et al 2008) showed that the spread in $T_1$ increased with increasing dilution of a test meal confined within an intra-gastric balloon and suggested that the accuracy of quantification of secretions would decrease with increasing $T_1$, although this would depend on the details of the $T_1$ measurement protocol and to what extent it was optimal for measuring the short $T_1$ of the meal. These technical issues may explain the unexpected drop in secretion level, seen between the $T = 15$ min and $T = 75$ min time points, in one of the subjects, who had a very high secretion volume, although as the total volume of this subject dropped over 200 ml in the 60 min between scans this drop in secretion may be a real effect due to rapid emptying of the secretion layer independent of the meal. A unique strength of the current study is that the accuracy of measurement of gastric meal and secretion volumes by MRI was validated by comparison to independent estimates of meal volume by gamma scintigraphy in the same subjects. Correcting the gastric content volume for the secretion volumes resulted in closer agreement between measurements of meal volume by the two modalities, with a linear regression gradient close to 1 and intercept almost zero. The percentage difference in meal volume between the two measurement techniques was higher at 75 min because, as demonstrated by Fidler et al (Fidler et al 2009), small changes in emptying rates between the two experimental days will have larger effects on the absolute volumes by the later time point.

The $T_1$ mapping method does have some limitations. First, although EPI overcomes the problem of motion artefacts, the EPI readout, used in measuring secretions makes it difficult to obtain measurements in the antral area of the stomach due to artefacts from air distorting the images. As a result of this issue, 5 out of the total 48 data sets had to be excluded due to poor positioning of the slice stack in this region. A further 3 data sets at the 75 min time point also had to be excluded due to poor fitting of the $T_1$ data. Second, the poor spatial resolution of the EPI acquisition may result in errors in the $T_1$ map at the edges of the stomach due to partial volume effects. Third this methodology is meal dependent, it works best in meals that mix easily with the gastric secretions and would not necessarily be appropriate for other liquid meals that congeal, precipitate or phase separate in the acid environment. Fourth, small movements in the distal stomach wall also reduce the accuracy of volume measurements (a source of error for all imaging techniques) (Treier et al 2008). Another issue with the proposed methodology is that the area sampled does not cover the entire stomach, due to both wall movements and susceptibility artefacts in the distal stomach. This technique assumes that measurements acquired from the proximal stomach are applicable to the whole organ. However, in fact, at the early time point, there is layering of secretion in the stomach due to gravity which depends on body position. The proximal stomach will therefore contain relatively more meal and less secretion than the distal stomach and this is the probable reason for the small but significant systematic overestimation of meal volume/underestimation of secretion volume seen in figure 6(b) at the 15 min time point. It was also observed that the longest $T_1$ and hence most concentrated gastric secretions were found at the stomach walls and formed a ring surrounding the liquid meal. By the 75 min time point the secretions are well mixed with the meal and the data from the proximal stomach should be a good representation of total gastric contents. All these observations confirmed previous results that have shown that the mixing of secretion into the meal takes time and is not homogeneous. Finally, it is worth noting that all MR relaxation time measurements are field strength dependent and measurements at 3.0T would need separate in-vitro calibration data. The use of EPI acquisition at higher field strength
would increase the image distortions, however alternative imaging sequences based on the bTFE acquisition could be used.

Notwithstanding these limitations, the proposed method has been shown to provide accurate measurements of gastric secretion that are sensitive to individual variation in secretion volumes, able to identify subjects that secreted large amounts of acid compared to those who secreted small amounts (figure 6(a)). Moreover the histogram of the T1 maps provides visualisation of the mixing processes that occur during gastric emptying of a meal. More frequent scanning may be necessary if the kinetics of secretion and emptying are to be explored in more detail (Sauter et al 2012). This technology may well have a role in clinical studies of reflux disease, functional dyspepsia and related disorders. (Tucker et al 2012, Curcic et al 2014a). Further it may be of great value in pharmacological studies that aim to either suppress gastric secretion or suppress gastro-esophageal reflux (Sweis et al 2013).

In conclusion we present an MRI acquisition and analysis protocol that provides accurate measurements of gastric volumes and secretion volumes with excellent inter-observer reproducibility for both variables. Gastric meal volumes obtained by this system were very similar to those obtained independently by gastric scintigraphy on a separate day. The protocol is patient friendly with rapid acquisition reducing the requirement to obtain data during breath-holding scans. Post-processing is rapid with a greatly reduced time required to derive measurements. Together these features provide the scientific basis of a protocol which would be suitable in clinical practice.

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