Small bowel transit of a bran meal residue in humans: sieving of solids from liquids and response to feeding

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Abstract

Background—Ileal motor patterns are adapted to the propulsion of viscous meal residue, such as bran, which accumulates in the distal ileum postprandially.

Aims—To examine the effects of a second liquid/solid meal on ileal emptying.

Subjects and methods—Eleven healthy fasting subjects consumed a 1.47 MJ pancake containing 15 g bran and 5 MBq Technetium-99m labelled amberlite resin (meal A). Gastric emptying and transit through the left upper quadrant (proximal) and right lower quadrant (distal) small bowel regions and colon were assessed scintigraphically. Transit was compared with and without a second Indium-111 liquid/solid DTPA labelled 2.28 MJ meal (B) given three hours after the first meal.

Results—Gastric emptying of meal A was slower than meal B (the time for 50% of the activity to leave the stomach (T50) being 113 (11) minutes versus 48 (3) minutes respectively, p<0.01, n=11). Both meals passed rapidly through the proximal small bowel (T50 meal A = 57 (14) minutes versus T50 meal B = 42 (11) minutes). Transit of meal A through the distal small bowel was much slower (T50 more than 390 minutes versus 176 (29) minutes for meal B, p<0.01), resulting in meal B overtaking meal A and entering the colon earlier. Ingestion of the second meal (B) resulted in significantly less meal A marker entering the colon (5 (3)% at 11 hours than when meal A was taken alone (18 (4)% (p<0.05, n=8).

Conclusions—The distal small bowel selectively retains bran, allowing liquid phase markers through to the colon. Consuming a second liquid/solid meal does not stimulate ileal transit of bran which seems to be propelled quicker by fasting motor patterns.

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Absorption of fluids and nutrients in the upper small intestine is usually rapid,1,2 such that the terminal ileum receives relatively small volumes of fluid containing indigestible, often viscous meal residues. The flow of this residue increases two to four hours postprandially,3 and the pharmacology and specific motor patterns and reflexes of the ileum4,5 discussed below, can be seen as an adaptation to its unique environment, lying between the jejunum and colon.

During fasting, aboral flow through the terminal ileum is greatest immediately before the passage of a phase III of the migrating motor complex,1 but also occurs with discrete clustered contractions and infrequent prolonged propulsive pressure waves which migrate distally in both man and dogs.5–7 Unlike those seen in dogs,8,9 human phase IIIIs do not regularly pass through the terminal ileum5 and emptying of the terminal ileum is more erratic, occurring for the most part without obvious relation to readily recognisable motor patterns.10

The discrete clustered contractions and the isolated prolonged propagated pressure waves, in dogs at least, are powerful, propulsive motor patterns which may be regarded as an adaptation to allow propulsion of the highly viscous material which may accumulate in the distal ileum.

Bran contains a substantial amount of indigestible, viscous material which will accumulate in the distal ileum. When compacted and dehydrated, it is one of the most viscous materials which the ileum is required to transfer, and we considered this to be a good test of ileal propulsive ability. Furthermore in subjects with functional bowel disorders, bran is poorly tolerated11 and associated with increased sensation of bloating.12 Previously it has been suggested that patients with irritable bowel syndrome in whom bloating is the main complaint show a defect in ileocolonic transfer of bran.13

The aim of the present study was therefore to explore further the handling of bran by the normal ileum prior to studying this phenomenon in the irritable bowel syndrome. In addition, we were particularly interested to determine whether the transit of bran through this region would be stimulated as previously described13 by feeding a second meal, either via a reflex, the “gastroileal response”, or as a direct effect of the increased flow through the terminal ileum which follows two to four hours postprandially.3

Materials and methods

SUBJECTS

Twelve healthy volunteers (four females, eight males; aged 19–47 years) were recruited into a two part study, each part separated by at least two weeks. All were free from any organic or functional gastrointestinal disorder, and taking no medication known to alter gastrointestinal
motility. Women were required to have a negative pregnancy test within the 24 hour period prior to each of the study days. Written informed consent was obtained from all subjects and the study was approved by the University of Nottingham Medical School Ethics Committee and the Department of Health (ARSAC) and carried out according to the guidelines in the Declaration of Helsinki.

STUDY DESIGN

Two studies were performed, separated by two to four weeks, the sequence being randomised. In part 1 of the study, volunteers ingested a solid meal following an overnight fast. The meal consisted of 5 MBq of Technetium-99m labelled amberlite resin baked into a bran supplemented pancake (1.67 MJ; Sainsbury’s Batter Mix, J Sainsbury plc, London (55% carbohydrate, 35% fat, 10% protein); 15 g bran, Natural wheat bran, Holland & Barrett, Hinckley, Leicestershire, UK). Thereafter subjects remained fasted but were allowed fluids, including decaffeinated tea and coffee, until the end of the study. In part 2 of the study, an additional Indium-111 DTPA labelled liquid/solid meal (Clinifeed 400 drink, Clintec Nutrition Ltd, Slough, UK (66% carbohydrate, 16% fat, 18% protein; 1.67 MJ) plus two digestive biscuits (0.61 MJ), McVitie’s Digestive, McVitie, Staines, Middlesex, UK) was consumed three hours after the pancake meal, following which subjects again remained fasted but were allowed fluids freely. The two meals were of very different viscosities, the liquid meal being watery (viscosity 1 Pascal sec) while the pancake was solid and required chewing prior to swallowing.

SCINTIGRAPHIC IMAGING

Anterior and posterior reference markers, consisting of Tc-99m labelled filter paper contained within a sheath of waterproof tape, were taped on to the abdomen over the right lobe of the liver anteriorly and posteriorly, and were used as guides to align successive scintigraphic images during analysis.

Anterior and posterior gamma scintigraphic images of 60 seconds duration were recorded using an IGE Maxicamera II (IGE Ltd, Herts, UK) fitted with a medium energy parallel hole collimator (300 keV maximum energy). Views were acquired using dual 20% symmetrical energy windows centred on the 245 keV of In and the 140 keV photopeak of Tc. The images were stored on a 128 x 128 matrix using a dedicated Nuclear Diagnostics computer system (Gravesend, Kent, UK).

Imaging commenced immediately after ingestion of the radiolabelled pancake, and thereafter at approximately 30 minute intervals until the end of the study 11 hours later.

SCINTIGRAPHIC ANALYSIS

Scintigraphic images for each subject were reviewed, and outlines of the stomach and colon constructed. The activity in these regions was determined using a variable region of interest program. Scintigraphic counts were corrected for background radiation, radioactive decay, and for anteroposterior movement by using the geometric mean of the anterior and posterior counts. Since the higher energy In causes detectable counts in the lower energy Tc window a correction factor was determined by assessing the apparent increase in Technetium counts immediately after the In was ingested. Technetium counts after the In ingestion were then corrected for this “down scatter”.

Gastric emptying time (GE T50) was taken as the time for 50% of the activity to leave the stomach, and small bowel transit (SBT) was calculated by subtracting GE T50 from the time for 50% of the activity to enter the colon. Additionally an assessment of regional SBT was made by drawing a line at 45 degrees through the small bowel region of interest, with its starting point at the hepatic flexure of the colon thereby dividing it into left upper quadrant (LUQ) and right lower quadrant (RLQ) regions. Transit through the LUQ was calculated from the difference between the time for 50% of the activity to accumulate in the RLQ or beyond and the GE T50 whereas transit through the RLQ was calculated by subtracting the time for 50% of the counts to reach the RLQ and beyond from the time for 50% of the activity to enter the colon.

STATISTICAL ANALYSIS

Results are expressed as mean (SEM). The significance of differences between paired data was assessed using the non-parametric Wilcoxon signed rank sum test.

Results

Eleven volunteers completed the study, with one subject dropping out after the fasted study as she was unable to tolerate the bran supplemented pancake. Accurate interpretation of the scintigraphic images of the small and large bowel was possible in all subjects except for one in part 1 of the study, and two in part 2 of the study. This was due to overlap of the small bowel and colon and these studies were omitted from the analysis.

GASTRIC EMPTYING

As expected there was no difference in gastric emptying of the solid pancake meal A in part 1 or part 2 of the study (115 (13) versus 113 (11) minutes, p=0.38, n=11). Gastric emptying of the pancake meal A was however significantly slower than the subsequently ingested liquid/solid meal B in part 2 of the study with a T50 of 113 (11) versus 48 (3) minutes respectively (p=0.004) (fig 1), even though the energy content was slightly less (1.67 MJ versus 2.28 MJ).

COLONIC ARRIVAL

In part 1 of the study, 18 (4%) of the pancake meal A label had reached the colon by the end of the study period (11 hours). Significantly less pancake meal A residue reached the colon by 11 hours in part 2 of the study (5 (3)%, p<0.05) (fig 2).

SMALL BOWEL TRANSIT

Exact calculation of the complete small bowel transit was not possible for solid pancake meal
A, as in no study did 50% of the activity reach the colon, and indeed for most subjects less than 10% of the residue had entered the colon by 11 hours. In notable contrast, 50% or more of the liquid/solid meal B reached the colon in eight of the nine individuals in which analysis was possible, even though it was ingested three hours after the pancake meal A (T50 liquid/solid meal B = 218 (23) minutes versus T50 solid meal A = more than 488 minutes, p<0.01, n=9).

REGIONAL SMALL BOWEL TRANSIT
Division of the small bowel into left upper quadrant (LUQ) and right lower quadrant (RLQ) regions revealed that transit of both the solid meal A and liquid/solid meal B residue was similar and relatively rapid (37 (14) versus 42 (11) minutes respectively, p=0.16, n=9) through the LUQ which emptied as the RLQ filled. By contrast, transit through the RLQ was slower and showed a pronounced hold up of the solid meal residue compared with the liquid meal residue (more than 390 versus 176 (29) minutes, p<0.01, n=9) (fig 3). This sequential transit and striking difference in transit times strongly suggest that the LUQ region was predominantly proximal and the RLQ predominantly distal small bowel.

Discussion
The most striking observation from this study was the selective retention of pancake meal residue in the distal small bowel while the liquid label from the second meal passed through. The two meals were of very different viscosities, the more viscous pancake meal moving more slowly than the liquid meal through the stomach and distal ileum. As the egg, flour, and fat from the pancake in the first meal would have been completely digested and absorbed in the more proximal bowel, the retained residue most likely represents a viscous mass of bran. In contrast, the 111In DTPA marker of the low viscosity liquid/solid meal B, taken three hours after the pancake meal, caught up with and overtook the bran residue, with the majority passing through into the colon. We consistently observed the accumulation of both the liquid phase marker, and to a much greater extent the solid phase marker, in the RLQ region of the small bowel, immediately adjacent to the lower pole of the ascending colon.

Selective emptying of liquids ahead of solids, “sieving”, is well recognised in the stomach, but initial human studies comparing iodine labelled cellulose with water soluble 99mTc DPTA suggested that this did not occur in the small bowel. Whether this does or does not occur may well depend on the type of solid since plainly it will require the physical separation of liquid and solid by mixing movements, which must be able to propel liquid forward while selectively retaining the larger particles as has been described with the canine stomach and rabbit colon. Studies in the dog have shown that the ileocolonic sphincter is able to regulate the exit of liquids from the small bowel, which occurs in discrete boluses in both dog and man. This bolus emptying would not be well suited to sieving, which seems to require passage through a longer segment of ileum, as shown by a study in which resin pellets and a liquid phase marker were infused into the ileum at either 50 or 100 cm proximal to the ileocaecal junction. At 50 cm
both solid and liquid markers were transferred to caecum together, but when infused at 100 cm proximal to the ileocaecal junction, liquids were more rapidly transferred to the colon.20

Although our method of division of the small bowel into proximal and distal segments is undoubtedly only an approximation, it was nevertheless useful in that it allowed a more objective measurement of our visual impression of rapid transit through the LUQ and relative stasis in the RLQ, as previously described by others.21 22 The sequential passage through the LUQ and then RLQ together with the very large differences in transit strongly suggest that the LUQ is predominantly proximal and the RUQ region predominantly distal small bowel. Overlap between the two regions does occur, but cannot be substantial because if this were so the transit times through the two regions would be much closer in value. Using this method we found transit through the proximal small bowel rapid and non-discriminatory for solid and liquid markers. In contrast, passage of both solid and liquid meal markers through the distal small bowel was much slower, with selective retention of the solid meal residue. A similar effect can be seen even if we consider the small bowel transit as a whole, with transit time being more than 488 minutes for the most rapid transit of bran residue compared with 218 (23) minutes for the liquid meal (p<0.01).

There appeared to be almost complete stasis of the solid meal A residue in the distal small bowel in our study, with less than 50% entering the colon by the end of both parts 1 and 2 of the study (11 hours). We were therefore effectively studying the arrival of the “head of the meal” and in retrospect a longer study time would have been better to allow a more exact definition of the arrival time of the bulk of the meal. Thus, although the transit time through the proximal small bowel was approximately 60 minutes, distal small bowel transit time was greater than 390 minutes. The stagnation of meal A residue is probably a result of the high bran content of our pancake meal. Other studies using intubation have found transit of liquid and solid residue through the last 60–100 cm of ileum to be considerably faster20 21 but this may reflect the excitatory effect of an intestinal tube.21 Exact meal composition appears to be important in determining transit in this region, as a previous study using a similar quantity of coarse bran to supplement a rice pudding meal showed a considerably faster small bowel transit time of 333 minutes.24 It is possible that in this previous study the fructose contained in the fruit jam and concentrated orange juice used with the rice pudding acted as an osmotic agent thereby accelerating transit and preventing the development of such a concentrated, viscous mass as was likely to be produced in this present study.

The other main and unexpected finding from the current study was that ingestion of the second liquid/solid meal B slowed terminal ileal emptying of the pancake meal A residue (fig 3). Although earlier reports25 suggested that a second meal would accelerate ileal transit by virtue of the gastroileal response, later studies showed that although there is an initial immediate increase in motor activity this does not translate into concomitant acceleration of a liquid phase marker through the distal 50 cm of ileum. Liquid transit does however accelerate two to four hours after a guar containing mixed solid/liquid meal,7 probably due to the increased flow of chyme that occurs at this time.3 Our study has shown that despite the probable increase in ileal motility induced by the second liquid/solid meal, this does not necessarily result in augmented propulsion of luminal contents, which depends on numerous other factors such as luminal viscosity and flow. We speculate that in the present study the migrating motor complex (MMC) would have reappeared four to six hours after ingesting meal A. However, ingestion of the second liquid/solid meal B would be predicted to delay further the appearance of the MMC thereby depriving the ileum of its most effective clearance mechanism for the solid meal residue. This may explain the reduced clearance of meal A ileal residue into the colon following the second liquid/solid meal, which we observed in part 2 of the study. It is possible that had our liquid meal been of a higher (4.18 MJ) energy content it would have stimulated rather than inhibited ileal emptying.

It was crucial in this study to identify and differentiate the terminal ileum and caecum accurately. This was relatively straightforward in most cases, as the Indium labelled liquid meal label passed through into the wider, readily recognisable ascending colon and so provided a clear landmark for analysis of the distribution of the Technetium label. We excluded from analysis the three studies in which this was not possible.

A further important issue relates to the solid meal A marker, Technetium, and its possible separation from the meal residue. By selectively labelling the bran, we have previously shown that bran remains in close association with meal residue during gastric emptying.24 Furthermore, it is well established that amberlite resin binds Technetium avidly, and it seems likely that the resin (together with label) becomes trapped in the “mesh” of viscous bran residue after digestion of the proteinaceous elements of the pancake meal. This supposition is strengthened by the fact that the water soluble marker5 in DTPA passed through the terminal ileum, overtaking the technetium label rather than flushing it ahead.

In summary, this study has shown that the distal ileum is able to retain solid meal residue selectively while allowing liquids to pass through into the colon. Although this may allow further opportunity for digestion and absorption, it may also cause a degree of partial obstruction, perhaps accounting for the known postprandial bloating which can occur some hours following bran ingestion in the irritable bowel syndrome.22

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