Superior mesenteric artery blood flow and gastric emptying in humans and the differential effects of high fat and high carbohydrate meals

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Abstract
This study was designed to determine if the differential effect of high fat and high carbohydrate meals on mesenteric blood flow is a result of changed gastric emptying rate. Eight healthy men were studied twice. Superior mesenteric artery blood flow (Doppler ultrasound) was measured before and after a 2-5 MJ meal (either 74% of the energy as carbohydrate or 71% as fat). Emptying of meals was followed by γ-scintigraphy. The pattern of the superior mesenteric artery blood flow response was different after the two meals (interaction effect p<0.001 analysis of variance), with a far more sustained response after fat. The time by which half the meal had emptied (t_50) was also significantly greater after fat (p<0.02). Superior mesenteric artery blood flow corresponding to t_50 was 449 ml/min after carbohydrate and 592 ml/min after fat. There was a significant curvilinear relation between the superior mesenteric artery blood flow response and gastric emptying after carbohydrate (r^2=0.94) and no relation at all after fat. This study confirms the finding that ingestion of meals with a high fat content slows gastric emptying compared with meals with a high carbohydrate content in healthy volunteers. A more sustained mesenteric hyperaemia was also recorded after the fat meal compared with the carbohydrate meal. The relation, however, between the volume of meal remaining in the stomach and the mesenteric response was considerably different after the two meals. Further study is required to elucidate the mechanism behind the vascular responses recorded in the mesenteric bed after food in humans.

Subjects and methods
Eight healthy men (22–27 years, body mass index 21.8–24.8 kg/m²) were recruited for the study. The subjects were screened for any cardiovascular or gastrointestinal disorder and none were smokers. The subjects gave written, informed consent to take part in the study, which was approved by the University of Nottingham Medical School ethical committee and the British Department of Health (ARSAC). The subjects were studied on two occasions, with the experiment starting at the same time on both occasions (12.30 pm) to avoid the effect of diurnal variation on gastric emptying rate. The subjects were requested to eat a light breakfast before 8.00 am (caffeine free) and to avoid strenuous exercise on the morning of the studies. On entering the laboratory low energy radioactive anatomical markers (technetium-99m Tc) were attached to the surface of the skin anteriorly and posteriorly. When visualised on the computer screen linked to the gammacamera, these prevent inaccurate repositioning of the subject and also make any subject movement during data acquisition obvious. Once the markers were in place the subjects rested supine for 30 minutes.

After the rest period, fasting superior mesenteric artery blood flow velocity measurements were made using transcutaneous Doppler ultrasound with a 5 MHz curvilinear pulsed wave imaging system with a 2 MHz offset Doppler. On occasions the vessel was obscured by the presence of abdominal gas, particularly in the postprandial stage of the study. In such cases, the angle at which the subject rested was carefully adjusted to a semisupine position, having the effect of displacing the gas and permitting measurement to proceed. The subject was then immediately returned to the supine position. The angle of insonation was recorded and used to convert the Doppler shift values (kHz) into blood flow velocity (cm/s). Care was taken to
ensure that wherever possible the same angle of insonation was used in each subject (mean angle of insonation was 29°, range 18–38°). Error brought into the flow calculation due to an angle of insonation of 29° would be in the range of 4–5%. Doppler signals were recorded, and analysed by spectral analysis using a 2 channel analyser that displays forward and reverse signals separately. Eighty separate frequency measurements were made in the forward and the reverse channel and the input spectrum sampled at 5 ms intervals. Maximum pulse repetition frequency was 5 MHz. Recordings were made with the subjects' breath held in mid inspiration, and stored on video tape. Mean values of time averaged mean velocity (TAM calculated using the intensity weighted mean frequency) and peak systolic velocity were taken from at least 10 Doppler waveform complexes. The time averaged mean velocities were used in the calculation of absolute flow (ml/min). Peak systolic velocity is the maximal velocity reached during each cardiac cycle, and occurs during systole.

The manually operated calipers used to estimate vessel diameter on the Duplex ultrasound machine give diameter readings with an accuracy of 1 mm and cannot resolve changes during the cardiac cycle. At each time point the diameter was determined from a mean of four measurements of the proximal portion of the vessel.

After the fasted measurements the subjects sat up and ingested, in a randomised order, either a high carbohydrate or high fat meal. Both meals had a total energy content of 2·5 MJ. They were based on meals of chicken, gravy, potato, carrots, and banana custard or ice cream and crème. The percentage energy as carbohydrate was 74% in the high carbohydrate meal and 14% in the high fat. The energy content as fat was respectively 7 and 71%. Sodium and potassium contents were around 330 mg and 1650 mg in the two different meals. Volume occupied by the meals did differ, with the high carbohydrate meal being approximately 600 ml compared with the high fat meal which was approximately 350 ml. The weight of the high carbohydrate meal was 581 g and the high fat 325 g.

The vegetables and gravy were mashed and 300 mg of amberlite ion exchange resin BDH (IRA 416) (milled to form a coarse mixture of variable diameter particles), were mixed with the mash. The particles were previously labelled by addition of TcO4⁻ calculated to be 3 MBq at the time of dosing. No attempt was made to selectively label any single component of the meals and it was assumed that the labelled particles would be distributed throughout the stomach contents once ingested. Techniques are available that permit radioisotopic labelling of fat, but the aim of this study was to investigate the effect of fat on the emptying of the entire contents of the stomach.

No longer than 15 minutes were taken to eat the meal on any occasion. Apart from one subject who could not complete the high carbohydrate meal (about 10% remained uneaten), no subject voiced any feelings of discomfort or nausea after either the high fat or high carbohydrate meals and the opinion tended to be that the high carbohydrate meal caused a greater feeling of fulness compared with the fat meal. Inclusion of the data from the subject who failed to complete the meal had no effect on the overall results.

On completion of the meal, the subjects immediately returned to the supine position and imaging was started. Although the tendency to redistribute blood away from the mesenteric bed during exercise is less in the postprandial state, movement of the subjects over the remainder of the experimental period was kept to a minimum. Posture has an effect on the rate of gastric emptying, and in this study the subjects remained quiescent and in the supine position.

Gastric emptying data were acquired with 30 second anterior and posterior images of the stomach every 20 minutes using a IGE gamma camera 11 gammacamera fitted with a low energy general purpose collimator. The gammacamera was linked to a dedicated nuclear diagnostics nuclear medicine computer system. Regions of interest were created around the computer generated image of the stomach for both anterior and posterior images, and counts recorded. The geometric mean of the anterior and posterior measurements was calculated and counts were corrected for background radiation and isotope decay.

Data were collected every 20 minutes, with radioisotope data acquisition being followed by superior mesenteric artery blood flow velocity measurements until at least 70% of the meal had left the stomach or 180 minutes had passed since ingestion of the meal. After this time the subjects tended to become restless.

STATISTICAL ANALYSIS

Statistical analysis of the results was performed by two way analysis of variance with repeated measures (ANOVA) using the package BMDP (BMDP Statistical Software, Los Angeles, California). A paired t test was used for comparison of the t50 data. Volume of food remaining in the stomach (ml) was calculated by multiplying the percentage of the meal remaining in the stomach by the total volume of the meal.

Data are presented in the figures as means (standard errors of the means) except where shown. The data reported in the text are absolute values with standard errors in brackets, and changes from baseline with the 95% confidence intervals of the change unless stated otherwise.

Results

SUPERIOR MESENTERIC ARTERY BLOOD FLOW

Fasted superior mesenteric artery blood flow was 346 ml/min (27·1) before the high carbohydrate meal and 351 ml/min (20·4) before the high fat meal. A peak blood flow of 611 ml/min (80) was achieved 20 minutes after ingestion of the high carbohydrate meal (95% confidence intervals of the increase above baseline 106 to 424 ml/min). In contrast, a peak blood flow of 715 ml/min (49) was reached 40 minutes after fat (95% CI of the increase above baseline 237 to 491 ml/min). The peak blood flows recorded after the two meals were not significantly different (Fig 1).
After the early blood flow peak with carbohydrate, superior mesenteric artery blood flow began to fall towards fasted values, and was not significantly different from baseline 160 minutes postprandially. In contrast, mesenteric blood flow was still 277 ml/min above pre food values 180 minutes after fat (95% CI of the increase 144 to 410 ml/min). The pattern of the mesenteric blood flow response was significantly different after the two meals (interaction effect p<0·001, ANOVA).

**PEAK SYSTOLIC VELOCITY**

Peak systolic velocity in the fasted state was 93 cm/sec (7·3) before the high carbohydrate meal and 89 cm/sec (4·8) before the high fat meal. The highest peak flow velocity after the high carbohydrate meal was reached at 60 minutes (141 cm/s (4·6)) (95% CI of the increase above baseline 33·8 to 61·2 cm/s), and after the high fat was 148 cm/s (8·3) at 100 minutes (95% CI of the increase above baseline 37·6 to 81·4 cm/s). There was no statistical difference between the maximum peak systolic velocities after the two meals. There was a significant difference in the pattern of change after the two meals (interaction effect p<0·02, ANOVA) (Fig 1).

**GASTRIC EMPTYING**

No substantial lag phase between ingestion and gastric emptying was seen after either meal – that is, emptying had started within the first 20

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**Figure 1:** Superior mesenteric artery blood flow (ml/min) and peak systolic velocity (cm/s) during the baseline period (time 0) and after the ingestion of the high carbohydrate (C) and high fat (F) meals. Values are means with their standard errors represented by vertical lines. The pattern of blood flow response and peak systolic velocity was significantly different (interaction effect p<0·001, ANOVA), with blood flow returning to a value not significantly different from baseline 160 minutes after the carbohydrate meal and flow remaining significantly raised throughout the experimental period after the high fat meal.

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**Figure 3:** Shows the individual t50 data after the ingestion of the high carbohydrate (C) and high fat (F) meals. t50 was greater after the high fat meal compared with the high carbohydrate meal in all but one of the subjects. The mean t50 values after the two meals (shown as horizontal bars) are significantly different (p<0·02).
minutes. There was a significant difference in the pattern of emptying after the two meals (interaction effect p<0.05, ANOVA) (Fig 2). There was a significant difference between the times at which 50% of the meals had left the stomach (t50), with t50 after the carbohydrate meal = 97.3 minutes (95% CI 87-107) and t50 after the high fat meal = 145.4 minutes (21.6) (t test p=0.02) (Fig 3).

Superior mesenteric artery blood flow at these time points was 449 ml/min after the high carbohydrate meal and 592 ml/min after the fat (95% CI of the difference 75 to 212 ml/min) (Fig 1). There is a relation between the volume of the meal remaining in the stomach and superior mesenteric artery blood flow after the high carbohydrate meal (Fig 4). The relation is described by an exponential line (r²=0.94). There seems to be no relation between mesenteric blood flow and volume remaining in the stomach for the high fat meal.

Discussion
In this study there was a significant difference between the t50 of the high fat and high carbohydrate meals. No attempt was made to specifically label any single component of the meals and this result confirms findings that fat does slow the rate of gastric emptying.11,12 The delay that exists between ingestion and emptying is referred to as the lag phase.13 This phase is short in length (approximately eight minutes), with fat increasing the lag period, possibly as a result of the redistribution of the stomach contents back to the proximal stomach.22 In this study a lag phase was not seen, possibly because of the discontinuous nature by which gastric emptying data were acquired,23 and it is feasible that this transient phase was missed.

The Hunt and Stubbs hypothesis states that gastric emptying of liquid meals is controlled by nutrient density (total kcal/g or ml of food), with the degree of slowing directly related to the nutrient density of the stomach contents entering the duodenum.1 In our investigation the two study meals were of identical energy content, t50 after the high carbohydrate meal was 97.3 minutes and nutrient density 2.5 Mj/581 g=4.3 kJ/g. Nutrient density of the high fat meal is 7.7 kJ/g. According to the above hypothesis, t50 after the high fat meal should be 151 minutes, which is surprisingly similar to the recorded t50 of 145 minutes.

Increasing weight of solid meals increases the absolute rate of emptying. It has been shown, however, that with meals of the same energy, a 300% increase in meal weight increases emptying rate by 388%, while a 304% increase in energy content reduces emptying rate by 43% in meals of the same weight.1 In our study, the meals consisted of similar food stuffs, with the high fat meal weighing 325 g and the high carbohydrate meal weighing 581 g. It is therefore possible that the delay in emptying seen after the high fat meal in this study is overestimated as a consequence of the difference in weight between the two meals.

The pattern of superior mesenteric artery blood flow response after ingestion of the two meals was significantly different. This confirms findings of a similar nature after both solid11 and liquid meals.10,11 The onset of the mesenteric response is slightly delayed after a high fat compared with a high carbohydrate meal.11,13 In this study there was no significant difference between the size of the mesenteric response 20 minutes after the two meals, although the peak value for the high fat meal occurred later.

In this study there was no significant difference between maximal hyperaemia after the two meals. Similar maximal flows after meals high in fat and high in carbohydrate have been shown by some11 and not others.24,25 All studies are in agreement that the mesenteric blood flow response is more prolonged after fat meals compared with carbohydrate, but postprandial measurements have not been continued beyond 90 minutes.13 In this study mesenteric blood flow was 79% above baseline 180 minutes after the fat meal. In contrast, after the high carbohydrate meal blood flow had returned to baseline values between 140 and 160 minutes. This may in part be accounted for by the delayed emptying of fat seen. Our results also suggest that fat is a more powerful stimulant of mesenteric blood flow than carbohydrate – that is, joule for joule, fat incurs a greater hyperaemia than carbohydrate.

It seems that there may be a relation between the rate of emptying of food from the stomach and the blood flow response. This would confirm reports that increased blood flow in the gut is associated with the absorption of nutrients.13 Calculation (from gastric emptying data) of the volume of food remaining in the stomach (Fig 4), however, shows a relation between volume emptied and mesenteric blood flow only after the high carbohydrate meal. No such relation exists after the high fat meal.

The role of gastrointestinal peptides in the vascular changes seen in the mesenteric bed on food ingestion is unclear. Gastric inhibitory polypeptide would respond to both meals, but does not have vasoactive properties. Pancreatic polypeptide probably responds to both meals, in particular the high fat meal. Neurotensin, the tachykinins, vasoactive intestinal polypeptide, and insulin will all respond to different degrees to the carbohydrate meal but do not have vasoactive properties.24 Of the peptides that have been investigated more fully, however, there is evidence that at physiological concentrations they play only an indirect part in the vascular changes associated with food ingestion.23,24 In this study no measurements were made of plasma concentrations of gastrointestinal peptides after the two meals.

In conclusion, this study has confirmed reports that fat slows the rate at which meals leave the stomach. Considering the effect of the weight of meals on gastric emptying,3 the effect of fat on gastric emptying may have been overestimated. This study also confirms findings that the mesenteric response after the ingestion of a high fat meal is slightly delayed and is more prolonged than after a meal low in fat content. Fat is also a more potent stimulator of mesenteric blood flow compared with carbohydrate. Finally, the relation between gastric emptying of a meal and the mesenteric response to that meal is different for meals of different composition. Further studies are necessary to clarify the
mechanisms by which these changes are mediated.

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