Effect of drink temperature on antropyloroduodenal motility and gastric electrical activity in humans

W M Sun, R Penagini, G Hebbard, C Malbert, K L Jones, S Emery, J Dent, M Horowitz

Abstract
There is little information on the motor mechanisms underlying the effects of meal temperature on gastric emptying. The effects on antropyloric pressures and the surface electrogastrogram of ingesting drinks at 4°C, 37°C, and 50°C (350 ml normal saline and 50 ml low calorie (7 kJ) orange cordial) given in randomised order were measured over 60 minutes in 12 normal volunteers (10 men and 2 women, aged 18–55 years). The warm and cold drinks suppressed antral pressure waves (p<0.05), altered the organisation of antropyloric pressure waves (p<0.05), stimulated isolated pyloric pressure waves (p<0.05), and increased electrogastrogram frequency (p<0.05) when compared with the 37°C drink. These changes were greatest in the first 30 minutes after ingestion and greater (p<0.05) with the 4°C drink. Temperature has major effects on post-prandial antropyloroduodenal motility in normal subjects. Both cold and warm drinks stimulate a pattern of motility associated with retardation of transpyloric flow.

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The rate of gastric emptying is influenced by the chemical and physical characteristics of food. Food rich in nutrients or of high osmolality leaves the stomach slowly, primarily as a result of feedback on gastric motility from receptors in the small intestine.1–3 Meal temperature also influences gastric emptying. We have reported that a low nutrient liquid ingested either cold (4°C) or warm (50°C) empties from the stomach more slowly than a drink ingested at body temperature.4 The effects of meal temperature on gastric motility have been studied in humans5 and in monkeys.6 Temperature was not found to have any effect on antral motility but the measurement techniques and experimental protocols used in these studies had major limitations. One of the possible mechanisms of action of meal temperature on emptying is via the gastric pacemaker which determines the timing and maximum rate of gastric contractions.7 The discharge of the gastric pacemaker can be recorded by surface electrogastrography (EGG). This non-invasive technique can detect gastric arrhythmias (tachygastria and bradygastria) in physiological and pathological situations.8–11 There are no data on the effects of meal temperature on the EGG. It is also possible that meal temperature has an effect on the spatial organisation of gastric contractions, independent of any effect on the pacemaker frequency but the available studies are insufficient to give any indication on this possibility. Localised pyloric contractions probably have a major role in the regulation of nutrient liquid emptying by acting as a brake.12

Our aim in this study was to evaluate as definitively as possible the effects of the temperature of a low nutrient drink on antropyloroduodenal motility and the EGG in humans. To do this we used manometric methods, which enabled detailed analysis of the occurrence and spatial organisation of antral and pyloric pressure waves, concurrent with cutaneous recordings of the EGG.

Methods

SUBJECTS
Studies were carried out in 12 normal volunteers, (10 male, two female; aged 18–55 years). None had a previous history of gastrointestinal disease or was taking medication. Smoking was prohibited from the evening before the study. The protocol was approved by the Human Ethics Committee of the Royal Adelaide Hospital in 1992 and each subject gave written informed consent.

PROTOCOL
The study began at approximately 10 am, after an overnight fast. A manometric assembly incorporating a sleeve sensor was passed into the stomach via an anaesthetised nostril. The sleeve was positioned across the pylorus by monitoring the antral and duodenal transmucosal potential difference (TMDP).10,11 Two adhesive electrodes were attached to the skin between the umbilicus and xiphoid to record gastric electrical activity (electrogastrogram – EGG).12 A third adhesive electrode served as a reference (Fig 1). The experiments were performed with the subject in a semi-recumbent position and the upper body at 45° from the bed. Each 400 ml drink consisted of 350 ml saline and 50 ml low calorie (7 kJ) orange cordial (Cottee’s, NSW, Australia), 290 mOsm/l, pH 3.5, and was given three times at 4°C, 37°C, or 50°C during phase I of the migrating motor complex. The order of the temperatures was randomised and each drink

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was consumed within three minutes. These drinks and temperatures were essentially identical to those used in our previous study.4

Antropyloroduodenal pressures and EGG were recorded for 60 minutes following each of the drinks and drinks were separated by at a minimum time interval of 90 minutes. Time zero was considered to be the time of completion of drink ingestion.

MEASUREMENTS

Antropyloric pressures
The 10 lumen manometric assembly was similar to that used in previous studies (Fig 1).12,14

The nine manometric sideholes were arranged at 1-5 cm intervals spanning the antrum and pylorus. A 4-5 cm sleeve sensor monitored pyloric contractions and was located between the sixth and ninth sideholes, counting from the proximal end of the manometric assembly. In order to define sleeve position, TMPD was monitored using the manometric sideholes at either end of the sleeve.15 All manometric channels were perfused with degassed distilled water at a rate of 0-3 ml/min, except the TMPD channels, which were perfused with degassed normal saline at the same rate.13

Manometric signals were amplified by a 16 channel polygraph (Synectics, Sweden). The signals were subsequently digitised at 10 Hz using an A-D card (NB-MIO16, National Instruments, Texas, USA) board, and then processed and stored in an Apple Macintosh Quadra 700 computer using proprietary software (MAD 16, Synectics/Royal Adelaide Hospital/C H Malbert) developed using the Lab View package (National Instruments).

Electrogastrography
To ensure optimal electrical contact of the surface electrodes, the abdominal skin was cleaned with ethanol and lightly abraded. The electrical signals were preamplified with a purpose designed preamplifier (Synectics, Sweden) (Fig 1). A Butterworth band-pass filter (attenuation 24 dB per octave) was used. The low and high cut off frequencies were 0-02 Hz and 0-28 Hz (~3 dB) respectively. EGG signals were sampled at 10 Hz and also stored on the Apple Macintosh Quadra 700 computer.

DATA ANALYSIS
Recordings were only analysed when the sleeve sensor was positioned correctly according to the following criteria: when the antral TMPD was equal to or more negative than –20 mV, the duodenal TMPD was equal to or more positive than –15 mV, and the difference between the two readings was at least 15 mV13. The first minute of recording after completion of the drink was excluded from data analysis (see below).

Terminology and definition of pressure waves
Phasic pressure changes which had an amplitude ≥10 mm Hg, and lasted for between 5–25 seconds were considered to be lumen occlusive pressure waves.12 An antropyloric pressure wave (APW) recorded by two or more sideholes was defined as a sequence when it occurred in at least one antral sidehole as well as the pylorus, and had onset times in adjacent channels within 5–10 seconds of each.16 Isolated pyloric pressure waves (IPPWs) were
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Antropyloric pressures and gastric electrical activity (EGG) after drinks at 4°C, 37°C, and 50°C.

<table>
<thead>
<tr>
<th>Temperature of drinks</th>
<th>4°C</th>
<th>37°C</th>
<th>50°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>First APW after drinking (min)</td>
<td>22 (5)*</td>
<td>13 (4)</td>
<td>19 (5)</td>
</tr>
<tr>
<td>No of APWs/60 min</td>
<td>21 (3)**</td>
<td>33 (5)</td>
<td>29 (7)*</td>
</tr>
<tr>
<td>No of IPPWs/60 min</td>
<td>30 (8)*</td>
<td>19 (7)</td>
<td>26 (7)*</td>
</tr>
<tr>
<td>Change in basal pyloric pressure (mmHg) 10 min after the drink</td>
<td>+0.1 (0.1)*</td>
<td>+0.8 (0.5)</td>
<td>+0.9 (0.05)*</td>
</tr>
<tr>
<td>Mean EGG frequency (Hz)</td>
<td>0.000 (0.000)*</td>
<td>0.000 (0.000)</td>
<td>0.000 (0.000)*</td>
</tr>
</tbody>
</table>

*p<0.05 cf 37°C; **p<0.05 cf 50°C; acompared with 2 minutes before the drink.

APW = antropyloric pressure wave; IPPW = isolated pyloric pressure wave.

defined as pressure waves that were detected by the sleeve sensor and by no more than one sidehole within the sleeve length, provided there was also absence of a pressure wave within ±5 seconds of any magnitude that was ascribable to either gastric or duodenal contraction. Basal pyloric pressure was defined as the difference between the basal pressure recorded by the sleeve sensor and the distal antral (antral TMPD) pressure after editing of phasic pressure, and was scored between 2–3 minutes before and 5–6, 10–11, and 30–31 minutes after each drink.

Electrogastrography (EGG)

EGG signals were processed by fast Fourier transformation (FFT) in order to obtain individual spectra. A Hanning window for the signal was used, as this method allows optimal definition of amplitude. The running spectral analysis was updated every minute from the preceding four minutes with an overlap period of three minutes. The mean frequency of the EGG was derived from the highest frequency of each spectrum.

STATISTICAL ANALYSIS

The number of pressure waves for either five or 10 minute recording periods are expressed as median values and interquartile range. Other data are presented as mean (SD). Statistical significance was examined using the quasi-likelihood method, a p value <0.05 being considered significant in all analyses.

Results

The experiments were well tolerated by all subjects. There were no significant differences in the time taken for subjects to consume the different drinks (1–8 (1.4) minutes for 4°C; 17 (1.1) minutes for 37°C, and 19 (1.2) minutes for 50°C). The sleeve sensor was positioned correctly >90% of the time. Satisfactory EGG signals were obtained for 88% of the time during cold drink, 86% of the time during warm drink, and 91% of the time after a drink at body temperature. After ingestion of the drinks there was no change in the quality of the signal.

During consumption of each drink, there were some pressure waves in the distal antrum and pylorus (Fig 2), but directly after completion of the drink there was a period of motor quiescence (Fig 2). The first APW occurred earlier after the quiescent period (p<0.05) following the 37°C drink than after the 4°C drink (Table). The effects of temperature on the number and organisation of APWs, IPPWs, and EGG frequency are summarised in the Table. There were fewer APWs (p<0.05) after the 4°C drink than after both the 37°C and the 50°C drinks. For all three drinks, the number of APWs increased gradually during the first 30 minutes and did not change after that time (p<0.05) (Fig 3). For the first 30 minutes following the 37°C drink the number of APWs in each five minute period was greater (p<0.05) when compared to both the warm and cold drinks (p<0.05) (Fig 3).

After each of the drinks most APWs were confined to the distal antrum (Fig 4). For both the cold and the warm drinks, there were fewer APWs in the proximal three antral sideholes in the first 30 minutes than in the second 30 minutes (Fig 4), whereas there was no difference with the 37°C drink. In the first 30 minutes, the number of APWs in both the proximal and the distal antral sideholes was greater (p<0.05) after the 37°C drink than after.
Figure 4: Number of antropyloric pressure waves (APW) originating in the proximal and distal antrum after the drinks. The effect of drink temperature on the number of APW was greatest for the distal antrum. The number of APW was greater after the 37°C drink than the other drinks for the first 30 minutes (*p<0.05). Proximal and distal antrum were defined as regions between sideholes that were located at 7.5–4.5 cm and 3 cm – antral transmucosal potential difference channel respectively.

Figure 5: Effect of drink temperature on the number of isolated pyloric pressure waves (IPPW) over time. The increase over the first 30 minutes was less following the 37°C drink compared to the other two drinks (*p<0.05). There were less after the 50°C drink than the 4°C drink (†p<0.05).

both the warm and cold drinks (Fig 4). This effect of temperature on the more proximal antrum was not apparent in the distal antrum, as with all three drinks there were no significant differences in the number of APWs in the distal three sideholes between the first and the second 30 minutes (Fig 4; p>0.05).

There were fewer IPPWs (p<0.05) in the first 30 minutes with the 37°C drink than with both the cold and warm drinks (Table) and there were more IPPWs (p<0.05) after the 4°C drink than after both the 50°C and 37°C drinks (Fig 5). There was a gradual increase in the number of IPPWs for up to about 25 minutes after each drink (Fig 5, p<0.05). The rate of increase of IPPWs in the first 30 minutes was less (p<0.05) after the 37°C drink than after both the 4°C and the 50°C drinks. Beyond 25 minutes, the number of IPPWs decreased (p<0.05) for all three drinks. Drink temperature had no effect on basal pyloric pressure (Table).

After the 37°C drink, the mean EGG frequency was similar to the predrink values (0.049 ± 0.018 Hz v 0.047 ± 0.01 Hz). Both the 4°C and the 50°C drinks were associated with a higher (p<0.05) mean EGG frequency than the 37°C drink (Table). This increase was due to periods of 6–10 cpm (0-1–0-17 Hz) EGG oscillation (p<0.05). There was no evidence of 3 cpm (0-05 Hz) activity during 6–10 cpm activity (Fig 6). The 6–10 cpm periods occurred most frequently within the first 30 minutes after the cold and warm drinks (Fig 7).

Discussion

We have reported previously that changes in meal temperature influence gastric emptying. The temperatures tested in our study represent the greatest range of normal drink temperature. These, and the content of the drink, were virtually identical to those examined in our previous study, so that comparable effects on gastric emptying would be anticipated. The mechanics of gastric emptying are complex, being dependent on the interaction of mechanical effects of the variable patterns of contraction of the proximal stomach, antrum, pylorus, and proximal small intestine. It does not seem that there is normally a single dominant mechanism which controls gastric emptying. The spatial/temporal organisation of gastric pressure waves may be a more important determinant of their mechanical consequences than their absolute number.

The present study, which is the first detailed investigation of the effects of meal temperature on gastric motility, shows that the cold and warm drinks cause suppression of antral pressure waves, alterations in the organisation of antral pressure waves and stimulation of pressure waves localised to the pylorus. These changes are likely to contribute to the retardation of gastric emptying produced by warm and cold drinks. Because of the significant impact of drink temperature on gastric emptying and antropyloric motility, this variable needs to be controlled for in studies of gastric motility and emptying.

At first glance our findings seem to be at variance with two previous studies, one in monkeys and the other in humans, which found no effect of temperature on antral motility. However, the rate of heat transfer to the thermoreceptors in the gastric wall is dependent on both the intragastric volume and the temperature gradient, and the negative observations in previous studies may reflect both the lower volume of the liquid meals (100–250 ml) used and the narrower range of temperatures that was evaluated (5–39°C). Our observation that both the magnitude of antral suppression and stimulation of isolated pyloric pressure waves were greater with the cold (4°C) than the warm (50°C) drink is not unexpected, as the temperature gradient was larger with the cold drink (33°C v 13°C). It should also be recognised that in previous studies the methods used to measure gastric motility were suboptimal. For example, in the human studies, pressures were recorded at only one recording site in the antrum and in the animal study,
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Figure 6: Pseudo-3D plot of the running spectrum analysis of the electrogastrogram (BOG), before, during, and after a 4°C drink. After the drink 6–10 activity is apparent.

Figure 7: The number of minutes that the 6–10 cpm (0.1–1.17 Hz) component of electrogastrogram (BOG) was present in each 10 minute period. The amount of 6–10 cpm activity was less after the 37°C drink than the other two drinks (*p<0.05).

Antral motility was evaluated with only one extraluminal strain gauge. These methods could not give information about pyloric contractions or the spatial patterning of antral contractions, variables that could be assessed fully in our study because of the use of multiple, closely spaced, manometric side holes. Unlike the present study, previous studies did not control for the effect of the phase of the interdigestive motor complex on gastric emptying and gastric motility. This is likely to have been a significant confounder, given the larger variations of antral motility that occur during the interdigestive cycle, which persist after ingestion of non-nutrient liquids in volumes in excess of those tested. Less direct studies by El and Mei in cats, have found evidence for temperature effects in that both cold (10–12°C) and warm (46–49°C) solutions infused into the duodenum inhibited antral spike activity, a finding which is compatible with our observation of reduced antral pressure waves.

Cold and warm drinks changed the patterns of antral pressure waves in that antral motility was totally suppressed for about 10 minutes. We have previously reported that the effects of temperature on gastric emptying are greatest during this time. While it is possible that the reduced number of pressure waves in the antrum could reflect larger antral volumes as a result of slow gastric emptying, this seems unlikely as the effects of temperature were evident soon after ingestion of the drinks, when differences in antral area are likely to be minimal, and were similar in the proximal and distal antrum.

Our data are the first to evaluate the effects of drink temperature on pyloric motility and suggest that the pylorus also plays a role in the temperature induced slowing of gastric emptying. Both cold and warm drinks stimulated IPPWs, a contraction pattern that has been shown to prevent transpyloric flow. The association of IPPW stimulation with slowing of gastric outflow is strong, since previous studies have shown that the retardant effects in gastric emptying of small intestinal nutrient infusion, cold stress, and induced hyperglycaemia are also associated with stimulation of IPPWs. These previous studies also found that basal pyloric pressure was increased concurrent with IPPWs. We cannot determine from the current experiment why pyloric tone was not stimulated by the cold and warm drinks. But have observed a similar disassociation between the tonic and phasic localised pyloric motor responses to duodenal distension in humans. There are other indications that pyloric tone and IPPWs are controlled separately, or at least have differing stimulation thresholds. Our data do not exclude an effect of temperature on proximal gastric tone and proximal small intestinal motility, which might also contribute to effects on gastric emptying. To our knowledge no studies have addressed this possibility.

The changes in EGG frequency observed after the cold and warm drinks are of uncertain significance. The 6–10 cpm frequency of EGG oscillation produced by the cold and warm drinks has previously been defined as tachygastria. It is seen with glaucogen induced antral motor inhibition and is associated with nausea and delayed gastric emptying. It is possible that bursts of tachygastria lead to antral motor inhibition following the warm and cold drinks and thereby contribute to slowing of gastric emptying. The prevalence of tachygastria after the 37°C drink was higher than expected. While we have no definitive explanation to account for this observation, it is unlikely to represent a harmonic frequency of 3 cpm as the observed frequency was never precisely twice the fundamental frequency.

Our study was not designed to identify the mechanisms that mediate the effects of temperature on gastric motility but other studies suggest several possibilities. Three types of thermoreceptors which are silent at normal body temperature have been demonstrated in the cat stomach and duodenum. Cold receptors start to respond at temperatures below 36°C, peaking at 10–12°C; warm receptors respond most intensely at 46–49°C; and mixed thermoreceptors respond to both warming and cooling of the mucosal surface. Cottrell has described cold sensitive

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mechanoreceptors in the sheep duodenum when recording from afferent C fibres, suggesting that thermoreceptors may respond to multiple stimuli. The time effects observed in our present study give some indication of possible mechanisms, as these mirror the changes in intragastric temperature that we have observed after ingestion of cold or warm drinks. After ingestion of 400 ml drinks at 4°C or 50°C, intragastric temperature returns to within 1°C of body temperature after 20–30 minutes. While the observation that differences in antpyloric pressures were still apparent at 15 minutes can be interpreted as indicating that minor changes in intragastric (proximal small intestinal) temperature may affect antpyloric motility, it is perhaps more likely that these effects are prolonged after the initial stimulation of these receptors when the temperature differences (from body temperature) are greatest.

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