Hyperglycaemia stimulates pyloric motility in normal subjects

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Abstract
The motor correlates of the delay in gastric emptying produced by hyperglycaemia were investigated in 11 healthy volunteers. Fasting gastroduodenal motility was measured during euglycaemia (blood glucose concentration 3–5 mmol/l) and during hyperglycaemia induced by intravenous dextrose (blood glucose concentration 12–16 mmol/l). Antral, pyloric, and proximal duodenal pressures were recorded by a sleeve/sidehole venous infusion assembly positioned across the pylorus, with the aid of measurements of transmucosal potential difference. During hyperglycaemia there was stimulation of isolated pyloric pressure waves when compared with the euglycaemia period (p<0.05). This was associated with inhibition of antral pressure waves (p<0.05). In nine of the 11 subjects an episode of duodenal 'phase III like' activity occurred within 15 minutes of the onset of hyperglycaemia. It is proposed that the stimulation of localised pyloric contractions and inhibition of antral contractions contribute to the delayed gastric emptying induced by hyperglycaemia. Abnormal gastric motility in patients with diabetes mellitus may be the result of hyperglycaemia itself, rather than irreversible autonomic neuropathy.

In normal subjects, hyperglycaemia induced by intravenous glucose infusion has been associated with slowing of gastric emptying of nutrient liquid meals 1 2 and a reduction in the rate of absorption of the sulphonylurea drug glipizide. 3 Furthermore, a relation between delayed gastric emptying and poor glycemic control in patients with diabetes mellitus has been shown. 4 5 There is little information about the motor mechanisms by which hyperglycaemia slows gastric emptying. Barnett and Owyang reported that a modest increase in the blood glucose concentration is associated with suppression of antral pressure waves and antral phase III motor activity in healthy fasted volunteers. 6 There are no data on the effects of hyperglycaemia on pyloric motility. Recent studies suggest that the pylorus plays an important role in the regulation of gastric emptying. 7 8 Accordingly, we investigated whether hyperglycaemia stimulates pyloric motility in healthy people.

Methods
Studies were performed on 11 healthy volunteers (eight women and three men) aged between 18 and 30 years. No subject had a history of diabetes mellitus, previous gastric surgery, or was taking medication at the time of the study. Informed consent was obtained in each case and the study protocol was approved by the Human Ethics Committee of the Royal Adelaide Hospital.

PROTOCOL
Antropyloroduodenal manometry was performed in each subject with a sleeve/sidehole manometric assembly during an initial 30 minute control period of euglycaemia followed by 60 minutes of hyperglycaemia induced by intravenous infusion of 20% dextrose. No adverse effects were reported by any subject during either infusion.

Before the start of each study a cannula was inserted into an antecubital vein. Each study began in phase I or phase IIA of the interdigestive motor complex. Once the manometric assembly was correctly positioned, 150 ml normal saline was given intravenously over five minutes: thereafter normal saline was infused at a rate of 150 ml/hour. These volumes were chosen to control for the volume of intravenous dextrose required to maintain hyperglycaemia. After 30 minutes an intravenous bolus of 150 ml of 20% dextrose was given over five minutes, followed by an infusion of 20% dextrose. Blood was drawn from a cannula placed in an antecubital vein of the arm not used for glucose infusion, and venous blood glucose concentrations were measured with BM-Test-Glycemia 20–800 strips (Boehringer-Mannheim) and a portable blood glucose meter (Refloflux IIM, Boehringer-Mannheim) at least every 10 minutes. The rate of dextrose infusion was adjusted to maintain a blood glucose concentration of approximately 14 mmol/l for 60 minutes. The accuracy of the blood glucose measurements was subsequently confirmed with a hexokinase technique.

The transnasally introduced manometric catheter was similar to that used in recent studies 9 and incorporated a 5 cm long sleeve sensor that was placed astride the pylorus to monitor pyloric pressures (Fig 1). Sideholes at 1 cm intervals along the sleeve allowed better definition of contractions around the pylorus. Manometric sideholes at each end of the sleeve also monitored transmucosal potential difference (TMPD) in order to verify assembly position. 10 Pressures and the TMPD values were recorded continuously throughout the experiment on a 12 channel chart recorder (model 7D; Grass Instrument Co, Quincy, MA, USA).

Sideholes were also located in the distal antrum 2 and 4 cm proximal to the sleeve and in the proximal duodenum 3 and 6 cm distal to the sleeve.

ANALYSIS OF MANOMETRIC TRACINGS
Tracings were only analysed if the sleeve sensor
was correctly positioned across the pylorus according to previously published criteria—that is if the antral transmucosal potential difference was more negative than −20 mV, the duodenal value was more positive than −15 mV, and the difference between the antral and duodenal measurements was at least 15 mV. On these criteria, the assembly was positioned correctly for more than 95% of the total recording time during the experimental period.

As in previous studies9 pressure waves were scored if their amplitude was greater than or equal to 10 mm Hg. Waves recorded by the sleeve sensor were classified as isolated pyloric pressure waves when they occurred in the absence of a wave of any amplitude in the antral or duodenal transmucosal potential difference sideholes. Basal pyloric pressure was measured for each minute of the study by determining the difference between the basal pressure recorded by the sleeve and that in the most distal antral sidehole. A basal pyloric pressure of >2 mm Hg was considered to indicate pyloric tone.10

In addition, duodenal contractions were assessed qualitatively to determine the phase of the interdigestive motor cycle.11

STATISTICAL ANALYSIS
Values were compared for each 30 minute interval (30 minutes during euglycaemia and 60 minutes during hyperglycaemia) of the study using the two tailed Wilcoxon matched signed rank test. A p value of <0.05 was considered significant in all analyses. All manometric measurements are expressed as median values and interquartile range. Blood glucose concentrations are expressed as mean (SEM) values.

Results
Dextrose infusion started a median (range) of 44 (3–77) minutes after completion of the preceding duodenal phase III contractions. The blood glucose concentrations are shown in Figure 2. There was a prompt increase in the mean blood glucose concentration from 4 mmol/l to 13 mmol/l after the initial bolus of dextrose. Blood glucose concentrations were subsequently maintained at approximately 14 mmol/l.

An example of a tracing from one subject showing the changes in motility patterns associated with hyperglycaemia is shown schematically in Figure 3.

The number of isolated pyloric pressure waves was significantly greater during hyperglycaemia than euglycaemia (Fig 4). There was no significant difference between the number of waves during the first 30 minutes of hyperglycaemia and the second 30 minutes of hyperglycaemia. Neither during hyperglycaemia nor euglycaemia was there evidence of pyloric tone in any subject.

There was a prompt reduction in the number of antral waves after the onset of hyperglycaemia. This reduction was sustained, almost no antral activity being observed during the second 30 minute period of hyperglycaemia (Fig 5). No antral phase III activity was seen after the induction of hyperglycaemia, but in nine of the 11 subjects an episode of high amplitude duodenal pressure waves with a rate of 10–14 per minute and a duration of three to five minutes was observed within mean (range) 15 (4–15) minutes of the dextrose bolus (Fig 6). This motor pattern was considered as 'phase III like' as it was indistinguishable from the duodenal phase III complex of the interdigestive motor cycle. The median (range) time of appearance of this activity was 55 (11–92) minutes from the previous phase III episode of duodenal activity. Before the infusion of dextrose duodenal phase III activity was preceded by phase III activity in the gastric antrum in all 11 subjects. The duodenal phase III like activity was followed by duodenal quiescence (phase I) for between 20 and 40 minutes.
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Figure 4: Number of isolated pyloric pressure waves during hyperglycaemia and euglycaemia shown as median values and interquartile ranges for 30 minute segments of recording. There is a significant and sustained increase in the number of IPPWs during hyperglycaemia.

Figure 5: Number of antral pressure waves during euglycaemia and hyperglycaemia shown as median values and interquartile ranges for 30 minute segments of recording. There is a significant and sustained decrease in antral pressure waves during hyperglycaemia.

when sporadic duodenal contractions (phase II) were again recorded.

Discussion
This study adds considerably to knowledge about the effects of hyperglycaemia on upper gastrointestinal motility. Firstly, we have shown that hyperglycaemia stimulates localised pyloric contractions. Secondly, we have confirmed the potent suppressive effect of hyperglycaemia on antral motility. Lastly, we have found that duodenal phase III like activity can be stimulated by hyperglycaemia.

The intermittent obstruction to gastric outflow by pyloric contractions and the suppression of antral motility are likely to contribute to the delay in gastric emptying caused by hyperglycaemia in healthy volunteers and in patients with diabetes mellitus. We did not assess the effect of hyperglycaemia on the motility of the proximal stomach, which is likely to be important in gastric emptying and may be abnormal in diabetic patients with gastroparesis. Further studies are also appropriate to examine the effect of lower blood glucose concentrations (within the range observed postprandially in normal subjects) on pyloric and duodenal motility. Barnett and Ouyang observed suppression of antral motility at a blood glucose concentration of 7.8 mmol/l, suggesting that changes in motor function may occur at concentrations that are seen after meals. Similar changes in antral and pyloric motility have been observed during intraduodenal dextrose infusion. Intraduodenal dextrose also results in an increase in basal pyloric pressure, which was not observed in the present study. This may reflect a difference in stimulus potency or possibly a difference in mechanisms responsible for the mediation of pyloric tone.

Both neural and hormonal mechanisms may be responsible for the antral inhibition and pyloric stimulation associated with hyperglycaemia. Raised blood glucose concentrations may directly stimulate small intestinal receptors and hence lead to similar motor patterns to those seen during intraduodenal dextrose infusion. The central nervous system may modulate antropic pyloric motor activity via altered vagal tone, and studies in animals suggest the pylorus is under tonic neural inhibition. As hyperglycaemia suppresses efferent vagal activity, increased pyloric activity may result from interference to vagal outflow. Ascending intramuscular pathways have been shown to mediate pyloric activity and raised blood glucose concentrations to impair nerve conduction. Intravenous glucose administration is also associated with changes in plasma concentrations of various hormones such as insulin, glucagon, and somatostatin which may influence antral and pyloric motility.

The unusual duodenal phase III like activity noted during hyperglycaemia has not been reported previously. While the interdigestive motor cycle is subject to considerable inter- and intraindividual variation, the close relation between the onset of these duodenal contractions and the establishment of hyperglycaemia, and the apparent absence of a relation with the time of the previous phase III, suggests that the triggering of this activity is related to hyperglycaemia. Previous workers did not detect any change in the mean cycle length of the duodenal phase III activity, but did not comment on the timing of these contractions. Duodenal phase III like activity has previously been shown in response to a variety of stimuli such as intraduodenal infusion of dextrose and lipid and intravenous infusion of β endorphin, and during stress induced by cold, pain, and labyrinthe.

![Figure 6: Timing of duodenal phase III like activity during each study. In 9 of the 11 subjects an episode of activity was seen within 15 minutes of the induction of hyperglycaemia. Length of solid bars represents length of contraction burst.](http://gut.bmj.com/first-published-as-10.1136/gut.32.5.475-on-1-May-1991/downloaded-from-http://gut.bmj.com/)}
stimulation. In healthy volunteers plasma insulin concentrations rise promptly in response to an intravenous glucose load. Furthermore, intravenous administration of insulin stimulates jejunal phase III activity before any significant change in blood glucose concentration. Hyperinsulinaemia may therefore be important in the mediation of the duodenal phase III like activity observed in the study.

Although the blood glucose concentrations achieved in our study are not seen in normal people, they are frequently observed in patients with type 1 and type 2 diabetes mellitus. Hyperglycaemia delays gastric emptying significantly and may lead to poor glycaemic control in diabetic patients by causing discrepancies between the onset of insulin action and the release of nutrients into the intestine. These effects may be partly explained by our observations on antral, pyloric, and duodenal motility. It is of interest that a recent study reported an increased frequency of phasic pyloric contractions in diabetic patients with gastroparesis. The results of our study suggest that this and other manometric abnormalities reported in diabetic gastroparesis — for example reduced antral motility and absent antral phase III activity with normal duodenal phase III activity may also be partly due to hyperglycaemia itself rather than irreversible autonomic neuropathy.

This work has been published in abstract form in the Australian and New Zealand Journal of Medicine and Gastroenterology. Dr Fraser was supported by a Royal Adelaide Hospital Dawes Postgraduate Research Scholarship. The study was supported by grants from the National Health and Medical Research Council of Australia and the Rebecca L Cooper Medical Research Foundation Pty Ltd. The authors thank Ms S Graham for expert technical assistance.


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