Effect of intravenous amino acids on interdigestive antroduodenal motility and small bowel transit time

H A J Gielkens, A van den Biggelaar, J Vecht, W Onkenhout, C B H W Lamers, A A M Masclee

Abstract

Background—Patients on total parenteral nutrition have an increased risk of developing gallstones because of gall bladder hypomotility. High dose amino acids may prevent biliary stasis by stimulating gall bladder emptying.

Aims—To investigate whether intravenous amino acids also influence antroduodenal motility.

Methods—Eight healthy volunteers received, on three separate occasions, intravenous saline (control), low dose amino acids (LDA), or high dose amino acids (HDA). Antroduodenal motility was recorded by perfusion manometry and duodenocaecal transit time (DCTT) using the lactulose breath hydrogen test.

Results—DCTT was significantly prolonged during LDA and HDA treatment compared with control. The interdigestive motor pattern was maintained and migrating motor complex (MMC) cycle length was significantly reduced during HDA compared with control and LDA due to a significant reduction in phase II duration. Significantly fewer phase IIIs originated in the gastric antrum during LDA and HDA compared with control. Duodenal phase II motility index was significantly reduced during HDA, but not during LDA, compared with control.

Conclusions—Separate intravenous infusion of high doses of amino acids in healthy volunteers: (1) modulates interdigestive antroduodenal motility; (2) shortens MMC cycle length due to a reduced duration of phase II with a lower contractile incidence both in the antrum and duodenum (phase I remains unchanged whereas the effect on phase III is diverse: in the antrum phase III is suppressed and in the duodenum the frequency is increased); and (3) prolongs interdigestive DCTT.

Keywords: amino acids; antroduodenal motility; small bowel transit time; total parenteral nutrition

Patients receiving total parenteral nutrition (TPN) have an increased risk of developing gall bladder sludge and stones. Bile stasis in the gall bladder during prolonged TPN is the major risk factor in the development of gall stones in these patients. The occurrence of gall bladder disease during TPN is prevented by intermittent intravenous injections of cholecystokinin (CCK), leading to profound gall bladder contraction. However, recent studies have shown that separate intravenous infusion of amino acids at high doses also induces gall bladder emptying. It has been suggested that this may be of clinical relevance as an alternative means, by modification of the nutrient regimen, of preventing TPN associated cholelithiasis.

In addition to inducing gall bladder emptying, intravenous amino acids stimulate gastric acid secretion, may increase pancreatic enzyme secretion, and delay gastric emptying of liquids. However, little is known about the possible effects of separate intravenous infusion of high doses of amino acids on gastrointestinal motility. Alterations in gastrointestinal motility may influence the risk of developing gall bladder sludge and stones: slow intestinal transit increases the risk of gallstone formation by influencing biliary lipid composition. We have studied the effect of two different doses of a commercially available mixed amino acids solution on interdigestive antroduodenal motility and small bowel transit time in healthy volunteers.

Methods

SUBJECTS

Eight healthy volunteers (five men, three women; mean age 23 years, range 20–34 years) participated in the study. None of the subjects had a history of gastrointestinal disease or surgery and none was taking any medication. Informed consent was obtained from each individual and the protocol was approved by the ethics committee.

STUDY PROTOCOL

Each subject participated in three experiments performed on separate days in random order with an interval of at least seven days during continuous infusion of either saline (control) or amino acids (Vamin 18EF, Kabi Pharmacia BV, Woerden, The Netherlands), given at a low dose (125 mg protein/kg/h; LDA) and at a high dose (250 mg protein/kg/h; HDA). The experiments were started at 8.00 am after an overnight fast. The manometric assembly was
positioned as described below. Two intravenous cannulas—one for blood sampling, the other for infusion—were inserted into the antecubital veins of each arm. The subjects were studied in a semirecumbent position. Antroduodenal motility was recorded continuously. Fifteen minutes after the onset of spontaneous phase III motor activity in the proximal duodenum, infusion of either saline or amino acids was started (defined as time 15 minutes) and continued for 375 minutes. At time 0 minutes, 6 g of lactulose (Legendal, Inpharzam, Amersfoort, The Netherlands), dissolved in 60 ml distilled water (osmolality 292 mmol) was administered slowly over 10 minutes into the distal duodenum through the central lumen of the manometric assembly in order to determine duodenocaecal transit time. Antroduodenal motility was recorded continuously; breath samples were obtained at 10 minute intervals. Blood samples for measurement of plasma CCK concentrations were obtained at times −30, −15, 0, 15, 30, 45, 60, 90, 120, 180, 240, 300, and 360 minutes. In addition, blood samples for measurement of plasma amino acid concentrations were obtained at time −15 minutes and at time 120 minutes.

ANTRODUODENAL MANOMETRY
Antroduodenal pressures were recorded using a water perfused stationary manometry system. A polyvinyl multilumen tube of 6 mm outer diameter incorporating six catheters with side holes located at 0, 5, 10, 15, 25, and 30 cm from the distal tip (Arndorfer Medical Systems, Greendale, Wisconsin, USA) was introduced transnasally and placed into the duodenum using a guide wire. Correct positioning of each of the pressure ports, the two side holes in the antrum and the four in the duodenum, was verified by fluoroscopy. The distal tip of the tube was located near the ligament of Treitz. Each lumen was connected to a pressure transducer (Medex, Hilliard, Ohio, USA) and perfused with gas-free distilled water by a low compliance pneumohydraulic infusion pump (Arndorfer Medical Supplies, Greendale, Wisconsin, USA) at a rate of 0.5 ml/min. The outputs from the pressure transducers were processed by an eight channel polygraph (Synectics Medical, Stockholm, Sweden), displayed on a monitor screen and stored on a personal computer system pending analysis. At the end of each experiment correct positioning of the manometric assembly was again verified by fluoroscopy.

SMALL BOWEL TRANSIT TIME
Lactulose breath hydrogen analysis was used to measure duodenocaecal transit time, as described by Bond and Levitt.19 End expiratory breath samples were collected under basal conditions and every 10 minutes after the lactulose had been administered intraduodenally until a sustained rise in breath hydrogen excretion occurred. Breath samples were collected in 25 ml plastic syringes and analysed immediately using a hydrogen breath test unit (Lactoscreen, Hoekloos, Amsterdam, The Netherlands). Duodenocaecal transit time was defined as the time between administration of lactulose (time 30 minutes) and a sustained rise in breath H₂ concentration of at least 10 parts per million (ppm) over basal.

ASSAYS OF AMINO ACIDS AND CCK
Plasma levels of amino acids were measured as described previously.19 Plasma CCK was measured using a sensitive and specific radioimmunoassay using antibody T204.

ANALYSIS OF MOTILITY DATA
Antroduodenal motility recordings were analysed both visually and by computer. The individual tracings were processed by specialised software (Polygram, Synectics Medical, Stockholm, Sweden) for adjusting baselines and extracting respiratory artefacts. However, the computer program does not recognise simultaneous pressure events as artefacts. Therefore, remaining artefacts obviously due to increases in intra-abdominal pressure were identified visually and excluded from analysis. Antral motility was analysed using the pressure tracings recorded from the side hole located most distal in the antrum (25 cm from the distal tip); duodenal motor characteristics were analysed using the pressure tracings recorded from the side hole in the proximal duodenum (10 cm from the distal tip) and from the most distal side hole (at the tip). Antral phase III of the migrating motor complex (MMC) was defined as rhythmic contractile activity at maximum frequency (three contractions per minute) for at least one minute in temporal relation with duodenal phase III activity. Duodenal phases of the MMC were defined as follows: phase I, motor quiescence (no more than two contractions per 10 minutes) for at least five minutes and preceded by phase III; phase II, irregular contractile activity at a frequency of more than two contractions per 10 minutes; phase III, regular rhythmic contractile activity at a frequency of 10–12 contractions per minute for at least two minutes. Phase IIIIs had to be propagated over at least two recording sites. MMC cycle length was defined as the time between the end of phase III in the proximal duodenum and the end of the next phase III. Origin (antrum versus duodenum), duration, and propagation velocity of each duodenal phase III were analysed visually. Propagation velocity was defined as the time interval between the onset of phase III activity in the proximal site and that in the distal site. Frequency, amplitude, and duration of the individual contractions were measured for both phase II and phase III using the computer program. Only pressure waves with an amplitude of at least 10 mm Hg and duration at least 1.5 seconds were considered as true contractions. Additionally, motility indices of the distal antrum and the proximal duodenum were calculated as area under the contraction curves for the last 30 minutes preceding each duodenal phase III activity.
**Statistical Analysis**

Results are expressed as mean (SEM). Differences in plasma CCK levels were analysed for statistical significance by multiple analysis of variance (MANOVA). Differences in plasma amino acid levels were analysed for statistical significance by paired analysis of variance. Data on phase III characteristics were pooled and analysed for statistical significance by unpaired analysis of variance. All other motility data were analysed for statistical significance by paired analysis of variance. Normal distribution of data was verified using the Levine test. When analysis of variance indicated a probability of less than 0.05 for the null hypothesis, Student-Newman-Keuls analyses were performed to determine which values between or within the experiments differed significantly. The significance level was set at p<0.05.

**PLASMA CCK**

Plasma CCK concentrations at time −15 minutes were not significantly different among the three experiments (control: 0.5 (0.1) pmol/l; LDA: 0.5 (0.1) pmol/l; HDA: 0.6 (0.2) pmol/l). No significant changes in plasma CCK levels were observed during either control or LDA. However, during HDA plasma CCK levels showed a small but significant (p<0.05) increase over basal levels during the first 30 minutes of infusion (fig 1).

**PLASMA AMINO ACIDS**

Table 1 presents plasma amino acid concentrations before and after LDA or HDA. No significant changes in plasma amino acid levels were observed in the control experiment. Intravenous infusion of amino acids dose dependently increased plasma levels of most amino acids, except glutamic acid.

**Antroduodenal Motility**

After the start of the intravenous infusion phase III recurred after 124 (34) minutes in the control experiment, after 114 (23) minutes in the LDA experiment, and after 53 (10) minutes in the HDA experiment (p<0.05 versus LDA and control). A total of 19 complete MMC cycles were recorded during the control experiment, 27 during LDA and 38 during HDA. The number of complete MMC cycles recorded for each subject was significantly (p<0.05) higher during HDA (4.8 (0.7)) compared with control (2.4 (0.3)) and LDA (3.4 (0.5)). Individual mean MMC cycle length was significantly (p<0.05) reduced during HDA compared with control and LDA (table 2). The shorter duration of the MMC cycles during HDA resulted from a significant (p<0.05) reduction in phase II duration compared with both control and LDA (fig 2). During LDA individual mean MMC cycle length and phase II duration were not significantly different compared with control. Duration of phase I was not significantly different among the three experiments (table 2).
Table 3  Characteristics of phase III of the migrating motor complex in the proximal duodenum while on LDA or HDA compared with control

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>LDA</th>
<th>HDA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antral origin/total</td>
<td>8/19 (37%)</td>
<td>4/27 (15%)</td>
<td>3/38 (8%)</td>
</tr>
<tr>
<td>Duration (s)</td>
<td>321 (23)</td>
<td>285 (17)</td>
<td>260 (11)*</td>
</tr>
<tr>
<td>Velocity (cm/min)</td>
<td>15.1 (2.3)</td>
<td>16.5 (3.2)</td>
<td>11.5 (1.5)</td>
</tr>
<tr>
<td>Frequency (contractions/min)</td>
<td>11.3 (0.1)</td>
<td>11.4 (0.1)</td>
<td>11.1 (0.2)</td>
</tr>
<tr>
<td>Mean amplitude (mm Hg)</td>
<td>27.9 (1.9)</td>
<td>28.8 (1.2)</td>
<td>25.5 (0.8)</td>
</tr>
<tr>
<td>Mean area under contractions (mm Hg.s)</td>
<td>47.8 (2.2)</td>
<td>50.3 (2.7)</td>
<td>43.5 (1.9)</td>
</tr>
</tbody>
</table>

Results expressed as mean (SEM). *p<0.05 compared with control.

Table 4  Motility characteristics for the last 30 minutes of phase II preceding duodenal phase III activity of the distal antrum and proximal duodenum while on LDA or HDA compared with control

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>LDA</th>
<th>HDA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antrum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frequency (contractions/min)</td>
<td>0.39 (0.10)</td>
<td>0.26 (0.09)</td>
<td>0.09 (0.03)*</td>
</tr>
<tr>
<td>Mean amplitude (mm Hg)</td>
<td>126 (14)</td>
<td>124 (14)</td>
<td>130 (15)</td>
</tr>
<tr>
<td>Motility index (mm Hg.s)</td>
<td>5100 (1585)</td>
<td>3345 (1669)</td>
<td>881 (335)*</td>
</tr>
<tr>
<td>Duodenum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frequency (contractions/min)</td>
<td>1.00 (0.17)</td>
<td>0.81 (0.11)</td>
<td>0.56 (0.08)*</td>
</tr>
<tr>
<td>Mean amplitude (mm Hg)</td>
<td>24.0 (1.2)</td>
<td>25.3 (1.5)</td>
<td>23.8 (1.5)</td>
</tr>
<tr>
<td>Motility index (mm Hg.s)</td>
<td>1209 (179)</td>
<td>1025 (137)</td>
<td>699 (141)*</td>
</tr>
</tbody>
</table>

Results expressed as mean (SEM). *p<0.05 compared with control.

Table 3 presents phase III characteristics of the proximal duodenum. During HDAs phase III duration was significantly (p<0.05) shorter compared with control and its propagation velocity tended to be reduced. During both LDA and HDA, significantly (p<0.05) fewer phase IIIIs originated in the gastric antrum.

Table 4 presents motility parameters of the last 30 minutes preceding duodenal phase II, especially late phase II. During high dose intravenous amino acids, the contraction frequency, and the motility index of late phase II were reduced. Therefore, intravenous amino acids may prolong intestinal transit time by reducing late phase II activity of the small intestine.

Discussion

The present study is the first to investigate the effects of separate intravenous infusions of amino acids on antroduodenal motility. Intravenous amino acids did not induce a fed-like motor pattern by disrupting the cyclic occurrence of the MMC. On the contrary, intravenous amino acids at high doses significantly increased duodenal phase III frequency. However, antral as well as duodenal phase II contractile activity was dose dependently reduced. Theoretically, the finding of a reduction in antral motility may result from technical shortcomings (e.g. catheter dislocation, number of antral recording sites). Correct positioning of the catheter, however, was checked before and after recording. Antral hypomotility was found only during HDA while the order of the experiments was randomised. Reduced antral motility may explain the delay in gastric emptying seen during infusion of intravenous amino acids. A recent study showed that intraduodenal L-tryptophan stimulates pyloric motility and may induce premature duodenal phase III activity. Therefore, the present study shows that not only intraluminal but also intravenous amino acids affect antroduodenal motility.

Intravenous amino acids at both doses significantly prolonged DCTT despite increased duodenal phase III frequency. Luminal flow is known to be more rapid during phase III activity. A tendency towards a decreased propagation velocity of the phase III activity fronts was only seen during high dose intravenous amino acids and therefore does not offer a full explanation. However, physiological studies have shown that few phase IIIIs eventually reach the caecum. Perhaps propagation distances of phase III are reduced during intravenous amino acids but the present study design does not allow comment on this possibility as motility recordings of the small intestine were restricted to the duodenum. Transit of luminal contents also occurs during phase II, especially late phase II. During high dose intravenous amino acids, duodenal phase II duration, the contraction frequency, and the motility index of late phase II were reduced. Therefore, intravenous amino acids may prolong intestinal transit time by reducing late phase II activity of the small intestine.

The mechanisms by which intravenous amino acids may affect gastrointestinal motility are largely unknown. Intravenous amino acids stimulate gastric acid secretion. Gastric acid inhibits antral phase III activity and decreases MMC cycle frequency due to the increased duration of duodenal phase II. Therefore, the effect of intravenous amino acids on intestinal motility is probably not mediated by increased gastric acid secretion, but it may, at least partially, mediate the observed inhibition of antral motility.
Intravenous amino acids induce gall bladder emptying and may increase pancreatic enzyme secretion. The delivery of bile and pancreatic enzymes into the duodenum may affect gastrointestinal motility. Intraduodenal administration of pancreatic enzymes has no effect on interdigestive antroduodenal motility. Intraduodenal bile stimulates motilin release. In humans, intravenous infusion of motilin initiates antral phase III activity. However, a recent study showed that although duodenal perfusion of pancreaticobiliary juice during phase I of the MMC stimulates plasma motilin release it does not change MMC cycle length. Penagini and colleagues studied the effect of intrajejunal bile acids, infused at concentrations resembling postprandial levels, on interdigestive jejunal motility. Duration of the MMC and of its phases was not significantly altered, although there was a trend towards reduced duration of phase II during jejunal infusion of cholesterylamine. However, bile acids significantly reduced phase II contraction frequency and prolonged small bowel transit time. Therefore, the effect of intravenous amino acids on intestinal motility and transit may be partially mediated by increased intraluminal concentrations of bile acids as a result of significant gall bladder emptying.

In the present study intravenous infusion of amino acids at the high dose slightly but still significantly increased plasma CCK levels (0.5–1.5 pmol/l) for a short period of time. Infusion of CCK resulting in plasma CCK levels comparable with those seen after ingestion of a fatty meal (5–6 pmol/l) disrupts the MMC, inducing a fed-like motor pattern and accelerates small intestinal transit. Furthermore, loxiglumide, a CCK-A receptor antagonist, reduces meal stimulated antroduodenal motility and delays small intestinal transit; suggesting that CCK is involved in stimulatory regulation. Therefore, it seems unlikely that the effect of intravenous amino acids on antroduodenal motility and intestinal transit is mediated by CCK.

The mixed amino acids solution used in the present study contains about 11 g/l l-arginine, a nitric oxide (NO) donor. Even when infused at a much higher rate than in the present study l-arginine has no effect on fasted antral motility in humans. In rats intravenous infusion of an NO donor during fasting disrupts the MMC and induces a fed-like motor pattern, whereas inhibition of NO synthase during feeding induces an MMC-like pattern. These data suggest that NO has no part in mediating the effect of intravenous amino acids on gastrointestinal motility.

Apart from l-arginine, commercially available mixed amino acids solutions contain other important amino acids. In the present study the intravenous amino acid mixture dose dependently increased plasma levels of many amino acids. The aromatic amino acids phenylalanine (Phe) and tryptophan (Trp) are potent stimuli of gastric acid secretion. It has been suggested that Phe and Trp may exert a direct effect on or near the parietal cell as parietal cell vagotomy does not inhibit their stimulatory effect on gastric acid secretion. Furthermore, in vitro studies in animals show that several amino acids (alanine, glycine, histidine, methionine, serine) are able to stimulate the colonic smooth muscle directly. However, whether amino acids have any direct effect on human small intestinal smooth muscle in vivo is not known.

Single administration of high doses of intravenous amino acids may be used as a prophylactic against gallstones. One should be aware of the fact that intravenous amino acids, by significantly delaying intestinal transit, also influence biliary bile acid composition (increase in biliary deoxycholic acid) and thus may increase the risk of gallstone formation during TPN.

In summary, the present study shows that in healthy volunteers high dose intravenous amino acids: (1) modulate interdigestive antroduodenal motility; (2) shorten MMC cycle length due to the reduced duration of phase II with a lower contractile incidence both in the antrum and duodenum (phase I remains unchanged whereas the effect on phase III is diverse: in the antrum phase III is suppressed and in the duodenum the frequency is increased); and (3) prolong interdigestive MMC. The question arises whether these effects of high dose intravenous amino acids may interfere with the “housekeeper” function of the MMC.

Part of this work has been published in abstract form (Gastroenterology 1995;108:A726).

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