Cholecystokinin in transient lower oesophageal sphincter relaxation due to gastric distension in humans

J Boulan, S Mathieu, M D’Amato, A Abergel, M Dapoigny, G Bommelaer

Abstract

Background and aims—Transient lower oesophageal sphincter relaxations (TLOSRs) has been found to be the main mechanism of gastro-oesophageal reflux. In dogs, cholecystokinin (CCK) is involved in their occurrence. The aim was to evaluate the role of endogenous and exogenous CCK in the occurrence of TLOSRs induced by gastric distension at constant pressure in humans.

Methods—Ten healthy volunteers were studied. Lower oesophageal sphincter pressure was monitored with a sleeve device and gastric distension was performed via an intragastric bag monitored by a barostat. During distensions, saline, CCK (30 ng/kg/h) or the CCK-A receptor antagonist loxiglumide (10 mg/kg/h) was perfused in a random double blind order.

Results—There was no significant difference between the number of TLOSRs during the different distensions with saline; CCK increased the number of TLOSRs at a mean rate of 1.31 compared with 9.1 with saline (p<0.001). Loxiglumide significantly decreased the number of relaxations to 5.3 versus 8.3 under paired saline infusion (p<0.001).

Conclusions—In humans, CCK-A receptor subtype is involved in the occurrence of transient lower oesophageal sphincter relaxations induced by gastric distension.

Keywords: lower oesophageal sphincter, transient relaxations, gastro-oesophageal reflux, cholecystokinin, loxiglumide.

Transient lower oesophageal sphincter relaxation (TLOSR), unrelated to swallowing, has been found to be the main mechanism of gastro-oesophageal reflux both in healthy subjects and in patients with gastro-oesophageal reflux disease. Research is now directed toward identifying the mechanisms involved in the occurrence of TLOSR and developing drugs to decrease its rate in patients with gastro-oesophageal reflux disease.

The frequency of TLOSR is greatly increased in humans and dogs by distension of the stomach, which probably triggers gastric mechanoreceptors mainly located in the subcardiac region. In dogs, cholecystokinin (CCK) is involved in the occurrence of TLOSR induced by gastric distension through peripheral CCK-A receptors.

In normal subjects, CCK-8 infusion and meals have been reported to reduce the lower oesophageal sphincter (LOS) pressure and both these effects were blocked by loxiglumide, a CCK-A receptor antagonist.

The aim of the present study was to evaluate the role of CCK-8 and loxiglumide in the occurrence of TLOSR induced by gastric distension in humans.

Subjects

We studied 10 healthy volunteers (five women and five men; age range 25-39 years). Subjects were free of any gastrointestinal symptoms and had no history of upper gastrointestinal surgery. They did not take any medication known to alter oesophageal motor function. Each volunteer gave written consent, and the study was approved by the human ethics committee of Clermont-Ferrand Hospital, France (Comité Consultatif et de Protection des Personnes dans la Recherche Biomédicale de la Région Auvergne).

Materials and Measurements

Oesophageal manometry was performed with a multilumen assembly incorporating a 6 cm sleeve device (ESM3DS Arndorfer Medical Specialties, Greendale, WI, USA). The sleeve sensor monitored LOS pressure. Side hole catheters recorded pressure in the gastric fundus and oesophageal body at 5, 10, and 15 cm above the LOS. Another side hole detected pharyngeal pressure to monitor swallowing. Catheters were perfused with gas free distilled water by a low compliance capillary pneumatic pump (Arndorfer Medical Specialties, Greendale, WI, USA). The perfusion rates were 0.1 ml/min for the pharyngeal catheter and 0.5 ml/min for the gastric and oesophageal body catheters and for the sleeve device. Output from the pressure transducers was processed by an eight channel polygraph (Polygraph HR Synectics Medical, Stockholm,
Studied. Due to the half life ofloxiglumide (>six hours), it was not possible to randomise its administration against placebo on a single day. Therefore the experiment was run on two separate days, saline being perfused during the first distension as a control and loxiglumide or saline in a random double blind order (loxiglumide and saline were delivered in similar blinded ampoules) during the second distension. Infusions of loxiglumide were started 20 minutes before gastric distension with a bolus of 30 mg/kg/h during 10 minutes and continued at a dose of 10 mg/kg/h until the end of distension.

**DATA ANALYSIS**

The reader was blinded as to whether CCK, saline, or loxiglumide had been infused. Manometric traces were analysed for basal LOS pressure, and for the occurrence of TLOSRs. Mean end expiratory LOS pressure was estimated with reference to gastric pressure at end expiration defined as zero.

Based on the analysis and according to Holloway et al., TLOSR was defined as: (1) the absence of swallowing four seconds before and two seconds after the onset of TLOSR, (2) a relaxation rate of 1 mm Hg/s, (3) time from onset to complete relaxation of 10 seconds, and (4) nadir pressure of 2 mm Hg. 

Excluding TLOSR associated with multiple rapid swallowing, falls in LOS pressure that fulfilled the last three criteria but had a duration >10 s were also judged as TLOSR, irrespective of the timing of the onset of the fall in LOS pressure in relation to swallowing.

The maximal distension volume was reported for each distension and for each subject.

**STATISTICAL ANALYSIS**

Data were compared using analysis of variance (ANOVA) and Student’s t test for paired values. Statistical significance was accepted if p<0.05. A statistical analysis of order and treatment effect was calculated. Values are presented as means (SD).

**Results**

**OESOPHAGEAL MANOMETRY**

As we always found the same LOS pressure or number of TLOSRs under saline whatever the order or day (no order effect), data obtained with CCK or loxiglumide were compared with the paired saline infusion (Fig 2).

**Transient lower oesophageal sphincter relaxations**

Without distension, at the basal state, TLOSRs occurred at a rate of 1-4 (1-0)/30 min.

Gastric distension with the barostat increased the number of TLOSRs. This increase was reproducible as the number of TLOSRs under saline infusion was similar during all series of distensions with saline (Fig 2).
During the first experiment, gastric distension induced the occurrence of TLOSRs at a mean rate of 9.1 (4.0)/30 min under saline infusion and 13.1 (5.5) min with CCK-8 infusion (p<0.001; Fig 3).

During the second experiment, loxiglumide infusion significantly decreased the number of TLOSRs to a mean rate of 5.3 (2.5) compared with 8.3 (1.7) under paired saline infusion (p<0.001; Fig 4).

No significant difference was found between the mean duration of TLOSR during saline, CCK-8, or loxiglumide infusion (26.9 (12.5) s, 23.6 (10.5) s, and 26.5 (6.7) s respectively).

**LOS pressure**

Table 1 shows the individual values of mean LOS pressure under gastric distension with either saline, CCK, or loxiglumide. Under saline infusion, from a basal value of 11.5 (3.0) mm Hg before distension, the LOS pressure was significantly increased by gastric distension to mean values ranging from 13.6 (5.0) to 14.2 (7.0) mm Hg (p<0.01) according to the various distension periods. Infusion of CCK significantly decreased the LOS pressure compared with paired saline infusion to a mean value of 8.3 (5.0) mm Hg (p<0.001). Loxiglumide did not significantly alter the mean resting LOS pressure after gastric
distension, compared with saline (11.6 (4.0) v 13.6 (5.0) mm Hg (p>0.05).

BAROSTAT

Intragastric pressure
From day to day, in each subject the constant pressure chosen for distension never varied by more than 2 mm Hg, the mean being 17.5 (0.5) mm Hg.

Intragastric distension volumes
At the constant distension pressure chosen, the average maximal value of gastric volume was significantly higher during CCK infusion (881 (235) ml) than with saline (763 (215) ml) (p=0.001).

On the contrary, the average maximal volumes decreased during loxiglumide infusion (631 (185) ml) compared with paired saline infusion (813 (134) ml) (p=0.001; Table II).

There was a slight but significant correlation (r=0.4, p<0.05) between the gastric volume and the number of TLOSRs, whatever the infusion (Fig 5).

Discussion
The results of the present study suggest that in humans gastric distension elicits TLOSRs and that endogenous CCK is involved via a CCK-A receptor dependent mechanism, confirming our previous findings in dogs.

In the first part of the study, our aim was to validate a model of induction of TLOSRs in humans via gastric distension at a constant pressure as gastric gaseous distension has been found to be a potent and consistent trigger of TLOSRs in dogs. In the same way, in humans stomach distension with carbon

Figure 2: All distensions with saline. There were no significant differences between these series.

Figure 4: Number of TLOSR/30 min. Numbers in parentheses are SD. Mean data with loxiglumide infusion compared with paired saline infusion p<0.001.

Table 1: LOS pressure: individual data and mean values (SD) in experiments 1 and 2.

<table>
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<th>Subjects</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
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</tr>
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<tr>
<td>10</td>
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<td>22.0</td>
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Mean (SD) (14.2 (7)) (8.3 (5)) (13.6 (5)) (11.6 (4)) *p<0.05 vs. paired saline.
dioxide induced TLOSRs similar to relaxations associated with acid gastro-oesophageal reflux. In our dog model, we used an electronic barostat to maintain constant intragastric pressure during distension. With constant insufflation, an eventual action of a drug on gastric or pyloric motility may lead to changes in intragastric pressure that will indirectly modify the occurrence of TLOSRs. Moreover, in dogs gas insufflation produces a distension of the gut, discomfort, and some vomiting. Consequently, and to avoid these problems in humans we preferred to use gastric distension with a bag. Other workers have used a fatty meal as a more physiological stimulant of TLOSRs; however, this usually led to a low rate of TLOSRs, insufficient to display a possible pharmacological reduction of their occurrence rate by loxigulamide. Moreover, the rate of TLOSRs elicited by gastric distension was not maximal as CCK was still able to increase their number further.

To obtain a similar number of TLOSRs at either control distension, we performed at the beginning of each experiment and for each subject a stepwise increase in gastric distensions with 2 mm Hg increments until we obtained a constant painful sensation. Then, for each subject and for all the distensions we chose a constant pressure corresponding to 75% of the pain threshold pressure. With this level of distension, we found a mean TLOSR number of eight to nine in humans, which is comparable with the mean of seven found in dogs. No significant difference was recorded from day to day concerning the pressure distensions in the same subject as defined above.

Table II: Intragastric distension volumes: individual data and mean values (SD) in experiments 1 and 2

<table>
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<th>Subjects</th>
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<th>Experiment 2</th>
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</tr>
<tr>
<td>10</td>
<td>390</td>
<td>380</td>
</tr>
</tbody>
</table>

Mean (SD) | 763 (215) | 881 (235)* | 813 (134) | 631 (185)* |

*p<0.05 v. paired saline.

Neither did we obtain any significant difference in the number of TLOSRs occurring the same day between the first and the second distension with saline so no order effect interfered with our results. Moreover, this number of TLOSRs did not change between different distensions with saline from day to day. This shows the reproducibility of this model in the triggering of TLOSRs.

By contrast with a recent study which found that CCK-33 infusion did not affect the occurrence of TLOSR in humans, we chose CCK-8, which can be considered as a neuronal form of CCK synthesised in nerve cell bodies and released at nerve endings. Due to the short half life of CCK-8 (sincalide; 1-5 minutes), we were able to run the first experiment in one day with a wash out period of 90 minutes between the two distensions, and saline or sincalide was perfused in a random order. By contrast, this procedure was impossible in the second experiment due to the longer half life of loxigulamide. So we subsequently performed the second experiment on two different days on each subject with perfusion of loxigulamide only during the second distension period in a double blind random order with placebo. Sincalide was perfused at a rate of 30 ng/kg/h as used medically to induce contraction of the gall bladder and to induce oesophageal motility. Loxigulamide (CR-1505) is the only CCK-A antagonist available for human use. Previous studies have shown that loxigulamide given intravenously in a dose we used in our experiments, completely inhibits CCK induced effects on the gall bladder and the pancreas at physiological concentrations.

The major finding of our study is the involvement of CCK receptors in the occurrence of TLOSRs induced by gastric distension in humans. This is in agreement with the postprandial release of CCK and the postprandial occurrence of TLOSRs. The selectivity of loxigulamide for CCK-A receptors indicates that TLOSRs produced by gastric distension are mainly controlled by CCK-A receptors. Due to the lack of specific CCK-B receptor antagonists widely available for human use, we could not study the possible and partial control of CCK-B receptors. Our results did not allow us to determine whether CCK-A receptors are directly or indirectly involved in the control of TLOSRs. Neither did they enable us to differentiate between a peripheral or central location of the CCK subtypes of receptors involved in the control of TLOSRs. Nevertheless, according to our results on dogs, we speculate that those receptors would be peripheral. Their location on vagal afferent fibres can be postulated as CCK has been found to activate these fibres and as CCK binding sites have been shown in the cervical and subdiaphragmatic vagus nerve. Moreover, our results suggest that gastric distension could trigger TLOSRs via stimulation of stretch fundic receptors as the number of TLOSRs paralleled the variation of gastric volumes, whatever the drug used. A neuronal or muscular location of the CCK-A...
receptors cannot be distinguished from our results. In this study, fundic distension induced a small but significant increase in resting LOS pressure as already found in dogs and in humans with such distension volumes.\textsuperscript{8} \textsuperscript{11} \textsuperscript{12} Infusion of CCK-8 induced a reduction in LOS pressure in humans.\textsuperscript{13} \textsuperscript{21} \textsuperscript{32} \textsuperscript{55} However, under our experimental conditions, loxiglumide did not modify the resting LOS pressure under gastric distension. It is worth noting that our subjects were sitting, which is not the usual position for studying LOS pressure. Moreover, the increase in LOS pressure afforded by the gastric distension could preclude a further effect of loxiglumide. On the other hand, our results do not conflict with those obtained in dogs, in which only CCK-B and not CCK-A receptors controlled the resting LOS pressure under gastric distension.\textsuperscript{12} Several authors have shown the inhibitory effect of a CCK-A receptor antagonist on the decrease of the basal LOS pressure induced by CCK infusion,\textsuperscript{13} oral ingestion of cholestyramine,\textsuperscript{30} or a fatty meal\textsuperscript{16} in humans. These results, taken together, allow us to speculate that, in humans, as in dogs, CCK-B receptors might play a part in the reflex involved in the increase in LOS pressure induced by gastric distension whereas CCK-A receptors are involved in the control of LOS pressure under endogenous release or infusion of CCK. However, the aim of our study with this model was not to evaluate the effect of CCK or CCK receptor antagonists on basal LOS pressure.

The finding that loxiglumide inhibits TLOSRs induced by gastric distension may have clinical implications as TLOSR is regarded as the main mechanism responsible for gastro-esophageal reflux in healthy humans as well as patients with gastro-esophageal reflux disease. It has been proposed that manipulating the afferent pathway, either peripherally or within the CNS, that controls and mediates TLOSRs would be the best way of targeting a pharmacological approach of the problem.\textsuperscript{57} Whether CCK antagonists will prove to be useful agents in reducing the frequency of TLOSR in reflux disease awaits further studies.

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