An exaggerated sensory component of the gastrocolonic response in patients with irritable bowel syndrome

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Abstract

Background/aims—Visceral hypersensitivity is a feature of the irritable bowel syndrome (IBS). Postprandial symptoms are common in these patients. The effects of nutrients on colonic perception in IBS are incompletely understood.

Subjects—We studied 13 healthy subjects and 16 patients with IBS—eight had diarrhoea predominant (IBS-D) and eight constipation predominant (IBS-C) IBS.

Methods—Colonic perception thresholds to balloon distension and viscerosomatic referral pattern were assessed before and after duodenal infusion of lipid or saline, respectively. At the end of the infusions, plasma levels of gastrointestinal peptides were determined.

Results—Lipids lowered the thresholds for first sensation, gas, discomfort, and pain in the IBS group but only for gas in the control group. The percent reduction in thresholds for gas and pain after lipids was greater in the IBS and IBS-D groups but not in the IBS-C group compared with controls. IBS patients had an increased area of referred discomfort and pain after lipids compared with before infusion whereas the referral area remained unchanged in controls. No group differences in colonic tone or compliance were observed. In both groups higher levels of cholecystokinin, pancreatic polypeptide, peptide YY, vasoactive intestinal polypeptide, and neuropeptide Y were seen after lipids. Motilin levels were higher in patients and differences in the subgroups were observed. Levels of corticotrophin releasing factor were lower in the constipated group than in the diarrhoea group.

Conclusions—Postprandial symptoms in IBS patients may be explained in part by a nutrient dependent exaggerated sensory component of the gastrocolonic response.

Keywords: irritable bowel syndrome; lipids; colonic perception; visceral hypersensitivity; gastrointestinal peptides

Methods

SUBJECTS

We studied 16 patients with IBS (14 women, two men; mean age 39 years (range 22–56)—eight with diarrhoea predominant (IBS-D) and eight with constipation predominant (IBS-C)

Abbreviations used in this paper: IBS, irritable bowel syndrome; IBS-D, diarrhoea predominant irritable bowel syndrome; IBS-C, constipation predominant irritable bowel syndrome; IOP, intraoperative pressure; CCK, cholecystokinin; NPY, neuropeptide Y; PYY, peptide YY; PP, pancreatic polypeptide; CRF, corticotrophin releasing factor; 5-HT, 5-hydroxytryptamine; PVEs, phasic volume events; SP, substance P; VIP, vasoactive intestinal polypeptide.
IBS—referred to our clinic from primary care physicians. All fulfilled the Rome criteria for IBS and organic gastrointestinal disorders were excluded by routine laboratory tests and colonoscopy with biopsies. The control group consisted of 13 healthy individuals (10 women, three men; mean age 30 years (range 20–43)). Mean age in the control group was significantly lower than that in the patient group (p=0.02). All control subjects were free of gastrointestinal symptoms and had no evidence of acute or chronic illnesses. None was taking any medication known to affect the gastrointestinal tract. All subjects gave informed consent and the study was approved by the ethics committee of the University of Göteborg.

STUDY DESIGN

The studies took place on two separate days: subjects were randomised to receive a lipid solution (Calogen 1.5 kcal/ml; Nutricia Nordic, Stockholm, Sweden) via an enteral feeding tube (Meda Polyuretansond; Fresenius AG, Bad Homburg, Germany) on one day and 0.9% saline on the other. Subjects were blinded to the solution they received. The infusion rate was 2 ml/min and the duration was one hour (fig 1).

After an overnight fast the subjects took a tap water enema (1500 ml) at home and arrived at hospital at 0730. Under fluoroscopic guidance the enteral feeding tube was placed with its tip in the descending part of the duodenum. The tube was then secured with tape to the cheek. Thereafter the balloon catheter, consisting of a highly compliant balloon made by polyethylene attached to a double lumen polyvinyl tube (Salem Sump Tube, 18F; Sherwood Medical, Tullamore, Ireland), was placed in the mid sigmoid colon. The balloon was tied at both proximal and distal ends and the distance between the attachment sites was 10 cm. Distension to a maximal volume of 550 ml resulted in a cylindrical balloon. A thin thread was inserted through the biopsy channel of a flexible fibreoptic sigmoidoscope and tied to the distal end of the balloon catheter. With the subject in the left lateral position the balloon catheter was carried along the sigmoidoscope to a level of 30–40 cm from the anal verge. In none of the subjects was retained formed stools observed in the rectum or sigmoid colon during this procedure. The balloon catheter was left in place as the sigmoidoscope was removed. After this procedure the balloon catheter was connected to a computer driven electronic barostat (Dual Drive Barostat, Distender Series II; G&J Electronics Inc., Toronto, Canada). The variation in time for placing the enteral feeding tube and the balloon catheter was small (<10 minutes), and so the experiment was started at the same time of the day in all subjects, approximately 0830. A small number of distensions up to 25 mm Hg were performed to unfold the balloon. Intraoperative pressure (IOP) was set to 2 mm Hg above the minimal distending pressure necessary to record respiratory variations in balloon volume. Thereafter a 30 minute equilibrium period was allowed with the balloon pressure at IOP during which colonic tone was assessed (fig 1). Two series of distensions were performed each day—one before and one immediately after administration of lipid or saline. When subjects received the infusions the balloon pressure was set at IOP. Subjects were told to report if they perceived any symptoms during the infusions. Immediately after the end of the infusions venous blood samples were obtained for analyses of gastrointestinal peptides before the second distension series (fig 1).

DISTENSIONS

We used phasic distensions of 30 seconds’ duration starting at IOP and increasing stepwise by 3 mm Hg until the subject reported pain or when a pressure of 50 mm Hg was reached. Pressure steps were used according to recent international recommendations. The distensions were separated by resting periods of 30 seconds with balloon pressure set to IOP. Subjects were instructed to grade their sensations during distensions using a keypad linked to the main barostat. A grading scale consisting of seven parameters was used: 1, no sensation; 2, fullness; 3, gas; 4, discomfort; 5, pain; 6, severe pain; and 7, intolerable pain. A tracking technique (single random staircase) was used, with tracking beginning when pain (=5) was first reported, after which 10 more distensions were carried out, provided the subject reported pain for at least three of them. Subjects were also instructed to mark the location of their respective sensation on a body map to evaluate the viscerosomatic referral pattern. This was done separately for each distension series.

GASTROINTESTINAL PEPTIDES

Blood samples were obtained for analysis of plasma levels of motilin, vasoactive intestinal polypeptide (VIP), PP, peptide YY (PYY), CCK, neuropeptide Y (NPY), substance P (SP), and corticotrophin releasing factor (CRF). The samples were immediately centrifuged at 3800 g at 4°C for 10 minutes. The supernatant was then aspirated and stored at −20°C until analysis of the peptides by a radio-immunoassay technique performed at the Department of Neurochemistry, Sahlgrenska University Hospital. CCK could only be analysed in the interval 0.4–12.5 pmol/l.

DATA ANALYSIS

The Protocol Plus software package (G&J Electronics Inc., Toronto, Canada) was used for data analysis. The pain threshold was the average pressure of the distensions where pain was reported. If pain was not experienced, the
pain threshold was set to a maximum pressure of 50 mm Hg. The thresholds for first sensation, gas, discomfort, and pain before compared with after saline and lipid, respectively, were compared in each group (healthy controls, IBS, IBS-D, IBS-C). Baseline thresholds—that is, before enteral administration of the solution—and the percent reduction in thresholds after lipids and saline were compared between groups. Colonic balloon volumes, reflecting tone, were assessed with pressure set at IOP. Barostat balloon volumes were averaged over 10 minutes and mean volumes were calculated for 30 minute periods before distensions (fasting tone) and during the first and second 30 minutes of the infusions, respectively. Balloon volumes before compared with during administration of the solutions were then compared. In addition, the number of phasic volume events (PVEs), defined as changes of $\geq 10\%$ compared with baseline volume and occurring at a rate of 1–4/minute, were calculated during the infusions and compared between groups and the two study days (fig 1). Compliance curves, that is pressure-volume relationship, were also created for each distension series by plotting the volume increase ($\Delta V$) against the corresponding pressure level above IOP ($\Delta P$). This was done up to the pressure level which the majority of subjects reached during the distension series both before and after the infusions. Changes within and between groups were then analysed. To assess the viscerosomatic referral pattern we calculated the relative area of referred discomfort on the body maps. The results were then compared before versus after the infusions and also between groups. Plasma levels of gastrointestinal peptides after lipid infusion compared with saline were assessed for each group and differences between groups analysed.

**STATISTICS**

Results are expressed as mean (SEM) with the exception of the gastrointestinal peptides, where median and interquartile range are used because highly skewed data were obtained for some of the peptides. Non-parametric tests were used for analysis of the peptides. The compliance curves were compared using analysis of variance (ANOVA), and the $\chi^2$ test with continuity correction was used for comparisons of symptoms during the infusions. Otherwise the t test was used for statistical comparisons. Significance was accepted at the 0.05 level.
significant for gas (212 (29) vs 203 (29) ml; 

p=0.16), discomfort (230 (15) vs 220 (27) ml; 

p=0.06), or pain (251 (25) vs 230 (26) ml; 

p=0.06). Highly significant reductions in 

perception thresholds for all sensations com-

pared with baseline were observed after lipid 

infusion in IBS patients (fig 2B). Also, volumes 

at the perception thresholds were significantly 

reduced for all sensations after lipid infusion 

compared with baseline in IBS patients—that 

is, for first sensation (159 (12) vs 124 (13) ml; 

p=0.05), gas (186 (15) vs 142 (13) ml; p=0.01), 

discomfort (230 (17) vs 202 (17) ml; p=0.005), 

and pain (264 (19) vs 230 (17) ml; p=0.003). 

The thresholds in the IBS subgroups were also 

evaluated. After lipid infusion, IBS-C 

patients had lower perception thresholds for 

gas (29.8 (2.4) mm Hg vs 23.5 (2.5) mm Hg; 

p=0.01), discomfort (41.9 (2.6) mm Hg vs 35.4 

(4.0) mm Hg; p=0.03), and pain (47.6 (1.5) 

mm Hg vs 40.5 (3.2) mm Hg; p=0.02) and the 

same was also true for the IBS-D group (gas: 

26.6 (3.3) mm Hg vs 19.0 (2.4) mm Hg 

p=0.02); discomfort: 32.9 (3.0) mm Hg vs 

24.5 (2.5) mm Hg (p=0.01); and pain: 41.1 

(3.2) mm Hg vs 29.8 (2.7) mm Hg (p=0.002)). 

Comparisons between patients and controls 

(percent reduction in thresholds) revealed that 

the reduced thresholds for gas and pain in IBS 

patients were significantly greater and trends in 

the same direction were observed for the two 

other thresholds studied. The IBS-D group 

differed in the same way from controls but that 

was not the case for the IBS-C group whose 

reduced thresholds did not differ significantly 

from control subjects (fig 3). We also analysed 

how many subjects in each group had reduced 

perception thresholds ≥10% after lipid infusion. 

For gas and pain it was 13/16 IBS patients 

compared with 4/13 control subjects (p=0.02), 

and for discomfort 12/16 vs 5/13 (p=0.11). 

None of the subjects reported “intolerable pain” 
during distensions and a minority reported “severe pain”; hence these descriptors 

were not used for further analysis. 

COLONIC TONE AND COMPLIANCE 

The minimal distending pressure did not differ 

between patients and controls (9.8 (0.8) 

mm Hg vs 9.7 (0.6) mm Hg; p=0.20). No changes in colonic balloon volumes were 

observed during the saline experiment and 

there were no significant differences between 
groups (data not shown). In four control 

subjects and one patient, adequate tone 

recordings were not obtained during lipid infu-

sion due to technical problems and the results 

of these subjects were excluded. During lipid 

infusion balloon volumes decreased in both 

groups compared with the fasting recording, 

reflecting increased colonic tone, both during 

the first and second 30 minutes of the infusions 

(fig 4). However, only the reduction in the 

patient group reached statistical significance. 

No significant differences in tone between 

patients and controls were observed in the fast-

ing state or during infusions (p>0.20 for all 

comparisons). There was no correlation be-

 tween the increase in colonic tone and percent 

reduction in perception thresholds (p>0.20 for 

all sensations). The number of PVEs was not 

significantly different during the lipid infusion 

compared with the saline infusion in patients 

(31 (5) vs 27 (6); p>0.20) or controls (16 (5) v 

9 (3); p=0.18). The IBS group demonstrated 

more PVEs during both saline (p=0.01) and 

lipid (p=0.04) infusions compared with controls. 

The infusions did not produce any significant changes in compliance (fig 5). In 

control subjects however, a trend towards 

higher compliance after saline (p=0.11) and 

after lipid (p=0.15) infusions was noted but no 
such trend was seen in patients (p=0.20). 

Compliance tended to be higher in patients but 

no significant differences were observed. 

THE VISCEROSOMATIC REFERRAL PATTERN 

In the patient group the relative area of referred 

discomfort increased after lipids compared with 

baseline (15.2 (3.2) cm² vs 9.0 (2.0) cm²; 

p=0.01). The same was also seen for the 

relative area of referred pain in patients (15.2 

(3.5) cm² vs 8.2 (3.2) cm²; p=0.03). In control 

subjects, the relative area did not change 

significantly after lipid infusion compared with 

baseline for discomfort (4.9 (1.2) cm² vs 5.6 

(1.3) cm²; p=0.15) or pain (6.8 (1.7) cm² vs 5.7 

(2.1) cm²; p>0.20). In the IBS group, 69%
(11/16) of subjects had increased abdominal and back areas of referred discomfort after lipids compared with baseline (fig 6) whereas this was seen in only 8% (1/13) of the control group (p<0.01). After saline, no changes in viscerosomatic referral patterns compared with baseline were seen.

GASTROINTESTINAL PEPTIDES
Higher levels of CCK, PP, PYY, VIP, and NPY were seen after lipid compared with after saline infusion in both the IBS group and the control group, but SP and CRF levels did not differ (table 2). Higher motilin levels after saline infusion in the patient groups were seen (p=0.04) and a tendency towards higher levels after lipid infusion (p=0.08). In the IBS group several very high motilin levels were seen, particularly in the IBS-C group. IBS-C patients exhibited higher motilin levels after saline than after lipids (116 (68–147) v 102 (56–136) pmol/l (median (IQR)); p=0.02) while a trend in the opposite direction was noted for IBS-D patients (70 (61–85) v 76 (70–115) pmol/l; p=0.06). Levels of SP tended to be higher in the control group than in the IBS group after lipids (p=0.05). Lower CRF levels were seen in the IBS-C group compared with the IBS-D group after saline (40.5 (35.0–41.5) v 45.0 (42.5–47.8) pmol/l; p=0.02) and lipids (39.0 (34.5–41.0) v 44.5 (43.0–48.0) pmol/l; p=0.005). There were no other differences between groups.

Discussion
We have demonstrated an exaggerated sensory component of the gastrocolonic response in IBS patients, expressed as a marked reduction in colonic perception thresholds and alteration of the viscerosomatic referral pattern after lipid administration in the upper gut. Together with postprandial motor alterations seen in IBS, our findings may explain postprandial symptoms in IBS patients.

As in many other studies, IBS patients in our study demonstrated visceral hypersensitivity compared with healthy individuals in the fasting state. However, pain thresholds did not differ between groups but this may have been for methodological reasons. A significant number of subjects did not experience pain at
Table 2  Gastrointestinal peptides in the control and irritable bowel syndrome (IBS) groups

<table>
<thead>
<tr>
<th></th>
<th>Control subjects</th>
<th>IBS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Saline</td>
<td>Lipid</td>
</tr>
<tr>
<td></td>
<td>Saline</td>
<td>Lipid</td>
</tr>
<tr>
<td>CCK</td>
<td>65 (50–76)</td>
<td>72 (61–78)</td>
</tr>
<tr>
<td>Motilin</td>
<td>108 (94–121)</td>
<td>117** (104–132)</td>
</tr>
<tr>
<td>PP</td>
<td>20 (19–23)</td>
<td>23 (16–25)</td>
</tr>
<tr>
<td>VIP</td>
<td>8.0 (7.0–11.2)</td>
<td>11.0* (8.8–13.8)</td>
</tr>
<tr>
<td>NPY</td>
<td>105 (97–110)</td>
<td>118** (106–127)</td>
</tr>
<tr>
<td>SP</td>
<td>1.4 (1.2–1.8)</td>
<td>1.3 (1.1–1.5)</td>
</tr>
<tr>
<td>CRF</td>
<td>42 (39–48)</td>
<td>42 (39–48)</td>
</tr>
</tbody>
</table>

Plasma levels of gastrointestinal peptides (pmol/l; median, interquartile range (IQR)).

*p<0.05, **p<0.01, plasma levels after lipid compared with after saline.

Baseline up to 50 mm Hg but their pain threshold was set to 50 mm Hg, which is an underestimation, and may have influenced the statistical comparison of pain thresholds between groups.

In our study, IBS patients were slightly older than controls. The small age difference between groups is unlikely to explain the differences in baseline thresholds between groups as recent studies have shown decreased rectal sensitivity with increasing age. After saline infusion the only difference compared with baseline was a small reduction in pain threshold in the patient group. We consider three possible explanations for this observation. Despite using a tracking technique to minimise psychological influences on the results, it is possible that the reduced pain threshold was due to psychological bias. Another possible explanation is that the first distension series, with several painful distensions, caused central sensitisation resulting in lower pain thresholds in the following series of distensions. There are conflicting results regarding the reproducibility of sensory thresholds determined on the same day. Our results, with no threshold changes in healthy controls during the saline experiment, indicate good reproducibility of the threshold determinations in our study. Therefore, we believe that the first two mechanisms are more likely explanations of the reduced pain threshold in patients after saline. However, these cannot explain the observed reduction in perception thresholds following lipid administration as this reduction was of a greater magnitude, particularly in the IBS group.

The small but significant reduction in the threshold for gas following lipid infusion in the control group and the tendency to lower thresholds for discomfort and pain are in agreement with previous reports in healthy controls. There seems to be a sensory component to the gastrocolonic response to food in healthy individuals although in the current study it was rather modest and limited to a small number of individuals. In the IBS group the reduction was more pronounced and a more constant phenomenon as the majority of patients demonstrated reduced thresholds after lipids. Both IBS subgroups had lowered thresholds after lipids but only in the IBS-D group was the reduction clearly different from controls. These findings are in agreement with another study showing differences in sub-groups of IBS patients. However, the relatively small numbers of subjects in the subgroups may explain the lack of significant changes between the IBS-C group and controls (type II error).

Tonic and phasic motor activity was evaluated by measuring colonic balloon volumes (tone) and phasic volume events during and before infusions. Colonic balloon volumes were reduced by about 20% in both groups during lipid infusion, reflecting increased tone. This is comparable with results obtained by Ford and colleagues. However, due to higher variability and a smaller number of subjects in the control group, the reduction in volume was significant only in patients. No differences in tone between groups were observed and no correlation between the increase in colonic tone and reduction in sensory thresholds was demonstrated. Thus it is unlikely that the pronounced reduction in perception thresholds after lipids in patients was solely secondary to the tonic motor response. Furthermore, in previous studies neither fasting nor postprandial colonic tone was found to be different in IBS patients compared with healthy controls. The phasic motor events were more frequent in patients during both the saline and lipid experiments but there were no differences between saline and lipids. Therefore, increased phasic motor activity is not a likely explanation for the observed group differences in perception after lipids compared with saline.

The viscerosomatic referral pattern in IBS patients differs from that in healthy individuals and a conditioning stimulus in the form of repetitive noxious sigmoid stimulation can increase the referral area of sensations provoked by rectal distension in IBS patients. These findings are thought to reflect abnormal spinal processing of viscerosensory information. In our study IBS patients, but not control subjects, had an increased area of referred discomfort and pain following lipid infusion. This may be due to central sensitisation of the dorsal horn neurones. This sensitisation may be secondary to an exaggerated postprandial motor response in IBS patients yielding more high amplitude contractions.

The pattern of increased levels of several gastrointestinal peptides after lipid infusion has previously been reported for CCK, PPY, and PP. The increase was similar in controls and patients. Therefore, our peptide data do not explain the pronounced threshold reduction in IBS patients. However, altered sensitivity to some of the peptides in the IBS group may be one explanation. It has previously been shown that infusion of cholecystokinin octapeptide (CCK-8) can provoke abnormal reactions of the gall bladder in IBS patients and higher pain scores in patients with functional abdominal pain, and unmask dysmotility in IBS patients.

Motilin, known to act primarily in the upper gut but also proposed to stimulate colonic motor activity, and CRF were the only peptides that differed between groups. Higher motilin levels were observed in patients, but
more interesting were levels of motilin in the IBS subgroups. Preston et al have shown lowered motilin levels after oral infused water in constipated patients relative to findings of disturbances in transit time, and patients with IBS-C have prolonged, and IBS-D shorter, whole gut transit time. These findings are to some extent in accordance with ours, as the group where slow transit is more likely (IBS-C) had lowered motilin levels after lipid infusion whereas the opposite was seen in IBS-D patients—that is, patients with probable accelerated transit time. The lower CRF levels in the constipated patients compared with the diarrhoea group were also interesting as CRF has been shown to stimulate colonic motor function. However, one must be cautious when interpreting our peptide data as no release profile of the peptides was analysed which is probably a more relevant approach.

There are several possible mechanisms for the pronounced reduction in thresholds in the IBS group. Reports of higher postprandial 5-HT levels in IBS patients with diarrhoea compared with healthy individuals and promising results in the treatment of patients with IBS with 5-HT, antagonists indicate that 5-HT may be a mediator of the sensory component. Another possibility is that endogenous factors could have influenced the lowered thresholds as the symptoms experienced during lipid infusion could have triggered a stress response, particularly in IBS patients. The altered visceral sensory referral pattern, demonstrated in patients, is thought to reflect abnormal visceral sensitivity and psychological factors may have influenced this through descending inhibitory and facilitatory pathways. However, some patients did not experience any symptoms during lipid infusion but still had an altered visceral sensory referral pattern and reduced perception thresholds after lipid infusion. In our opinion, this argues against the importance of psychological factors for our findings.

In conclusion, we have demonstrated that IBS patients exhibit an exaggerated sensory component of the gastrocolonic response, expressed as lowered colonic perception thresholds and increased visceral sensory referral area after duodenal lipid infusion. This may be an important factor in the postprandial symptoms in IBS patients. However, the mechanism remains to be clarified.

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