Pain Severity Correlates With Biopsy-Mediated Colonic Afferent Activation But Not Psychological Scores in Patients With IBS-D

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INTRODUCTION: Despite heterogeneity, an increased prevalence of psychological comorbidity and an altered pronociceptive gut microenvironment have repeatedly emerged as causative pathophysiology in patients with irritable bowel syndrome (IBS). Our aim was to study these phenomena by comparing gut-related symptoms, psychological scores, and biopsy samples generated from a detailed diarrhea-predominant IBS patient (IBS-D) cohort before their entry into a previously reported clinical trial.

METHODS: Data were generated from 42 patients with IBS-D who completed a daily 2-week bowel symptom diary, the Hospital Anxiety and Depression score, and the Patient Health Questionnaire-12 Somatic Symptom score and underwent unprepared flexible sigmoidoscopy. Sigmoid mucosal biopsies were separately evaluated using immunohistochemistry and culture supernatants to determine cellularity, mediator levels, and ability to stimulate colonic afferent activity.

RESULTS: Pain severity scores significantly correlated with the daily duration of pain ($r = 0.67$, $P < 0.00001$), urgency ($r = 0.57$, $P < 0.0005$), and bloating ($r = 0.39$, $P < 0.05$), but not with psychological symptom scores for anxiety, depression, or somatization. Furthermore, pain severity scores from individual patients with IBS-D were significantly correlated ($r = 0.40$, $P < 0.008$) with stimulation of colonic afferent activation mediated by their biopsy supernatant, but not with biopsy cell counts nor measured mediator levels.

DISCUSSION: Peripheral pronociceptive changes in the bowel seem more important than psychological factors in determining pain severity within a tightly phenotyped cohort of patients with IBS-D. No individual mediator was identified as the cause of this pronociceptive change, suggesting that nerve targeting therapeutic approaches may be more successful than mediator-driven approaches for the treatment of pain in IBS-D.


INTRODUCTION
Chronic abdominal pain and loose stools is a debilitating condition and one of the most common causes of presentation to a gastroenterologist. After excluding inflammatory or infectious disease, most cases are diagnosed as irritable bowel syndrome with diarrhea (IBS-D) (1). However, the cause of the characteristic pain remains obscure. Pain has been attributed to visceral hypersensitivity to otherwise non-noxious stimuli, found in between 50% and 90% of patients (2). This may arise from a range of abnormalities including enhanced nociception, augmented central pain processing (3), and impaired adaptation to pain (4), which may explain the heterogeneity of pathophysiology within patients with irritable bowel syndrome (IBS).

Although central factors are undoubtedly important, recognition that IBS could arise after acute infectious gastroenteritis, or postinfectious IBS (PI-IBS), and the chronic changes in mucosal...
cellularity and mediator content associated with PI-IBS (5,6) has focused attention on local mucosal abnormalities. Subsequent studies across different subgroups of patients with IBS have broadly documented altered mast cell and endocrine cell numbers and mediators (7). However, such findings are not universal and the link between mucosal changes, visceral hypersensitivity, and pain symptomology in unselected patients with IBS is variable (8–10). These disparities may be due to studying unselected, and hence heterogeneous patients with IBS, rather than specific subtypes.

The commonest subtype of PI-IBS is diarrhea predominant (11), a group in whom several studies have shown alterations in tight junctions (12) and increased permeability (13). These features are linked to visceral hypersensitivity by promoting exposure to luminal content and local immune cell activation (14). Evidence for these changes have come from biopsy studies that have documented mast cell hyperplasia and elevated levels of mediators such as histamine, serotonin, and tryptase, which contribute to activation of enteric nerves by biopsy supernatants (8,15,16). In some studies, these changes have been shown to correlate with pain scores or pain threshold assessed by rectal barostat, thereby providing a link between local changes in the gut mucosa with pain symptomology (17).

We have previously reported the results of a large proof of concept clinical trial in patients with IBS-D (18). Nested within the prescreening period of the trial was a mechanistic study that is reported here. We first examined the correlation between symptoms of pain with related sensory abnormalities of urgency, bloating, and psychological factors known to influence pain processing (19). We then focused on the histology of the sigmoid colonic mucosa and mediator release from incubated biopsies to identify mediators and cell types contributing to IBS pain. Finally, we tested the effect of sigmoid biopsy supernatant on colonic afferent activity and correlated this with pain symptomology. We found considerable heterogeneity in our patient group, with evidence that locally generated mediators are associated with the severity of abdominal pain, with a stronger effect than central psychological factors in this subtype of IBS.

**METHODS**

**Patient details**

Patients with IBS-D were recruited into a multicentered, parallel group, randomized placebo-controlled trial as previously reported (18) (ClinicalTrials.gov ISRCTN76612274). Nested within the main trial was a mechanistic study of 42 patients who were recruited in the Nottingham center and consented for sigmoid biopsy before randomization. In addition to stool consistency and frequency, other bowel-related symptoms comprising pain severity and daily duration, urgency, and bloating were recorded daily over the 14-day screening period. Pain severity was recorded on a 0–10 scale (0 = no pain and 10 = the most severe pain ever experienced) along with daily pain duration in hours per day. Urgency and bloating were also recorded on a similar 0–10 scale, stool frequency as bowel movements/day and stool consistency recorded daily using the Bristol Stool Form scale. Symptoms were recorded each evening documenting the preceding 24 hours.

All randomization patients completed the Hospital Anxiety and Depression Scale (20) along with the Patient Health Questionnaire-12 Somatic Symptom score, a measure of non-gastroenterological somatic symptoms (21).

**Sigmoid biopsy and assessments**

After symptom screening to confirm eligibility for the trial, mucosal biopsies were obtained at 30 cm from the anus during an unprepared, unprocessed sigmoidoscopy in the left lateral position. Two biopsies were taken and processed for immunohistochemistry and 2 were cultured to obtain supernatants. Biopsy processing is detailed in the Supplemental Methods (see Supplementary Digital Content 2, http://links.lww.com/CTG/AS10).

Two biopsies were weighed and placed into 2 mL of Hanks balanced salt solution in a polystyrene organ culture dish and incubated for two 30 minute periods at 37°C, in 5% CO₂. Fresh Hanks was used for the second, 30-minute incubation period. The supernatant collected in the first 30 minutes was used to assess released mediators, whereas the supernatant collected in the second 30-minute period was used for testing in colonic afferent preparations. Supernatants were aliquoted and stored at −80°C. Samples were transported on dry ice and aliquots were thawed on the day of use in electrophysiological or laboratory studies.

Biopsy supernatant levels of histamine, tryptase, chymase, and carboxypeptidase 3 (CPA3) were measured using sandwich ELISA assays provided by the Immunopharmacology Research Group, the University of Southampton, as described previously (22,23). Histamine was measured using a commercially available enzyme immunoassay kit (Neogen, Lexington, KY) as directed by the manufacturer.

**Ex-vivo recordings of colonic afferent fiber activity**

Few fiber afferent activities were recorded from teased lumbar splanchnic nerve bundles in a flat sheet colorectal preparation (male 12-weeks old, wild-type C57BL/6 mice, or Naᵥ1.9 −/− mice, as previously described (24)) using suction electrodes. Receptive fields were identified and characterized based on the criteria developed by Brierley et al. (25–27). Experiments were only performed on the receptive fields of vascular (or serosal) afferents. Once characterized mechanosensitivity was determined by probing with 0.6 g and 1.0 g von Frey hairs (vFh). Thereafter, a brass ring was placed around the receptive field and the indwelling buffer replaced with biopsy supernatant (100 μL) for 12 minutes, and mechanosensitivity retested after removal of the ring and supernatant. The individual single unit discharge of the receptive field tested was discriminated using template matching software within Spike 2 software (Cambridge Electronic Design, Cambridge, UK) performed over the period of vFh probing. Mechanosensitivity was determined for each weight of vFh before

**Table 1. Clinical details of patients**

<table>
<thead>
<tr>
<th></th>
<th>Mean</th>
<th>SD</th>
<th>N</th>
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</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>39.3</td>
<td>13.5</td>
<td>42</td>
</tr>
<tr>
<td><strong>Gender</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male/female</td>
<td>15/27</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>HAD anxiety score</strong></td>
<td>8.1</td>
<td>4.2</td>
<td>42</td>
</tr>
<tr>
<td><strong>HAD depression score</strong></td>
<td>4.8</td>
<td>4.4</td>
<td>42</td>
</tr>
<tr>
<td><strong>PHQ-12 SS score</strong></td>
<td>7.0</td>
<td>4.2</td>
<td>42</td>
</tr>
<tr>
<td><strong>HAD = Hospital Anxiety &amp; Depression Scale; PHQ-12 SS= Patient Health Questionnaire-12 Somatic Symptom.</strong></td>
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</table>
and after supernatant application, and the difference was calculated and expressed in action potentials (spikes) per second. Chemosensitivity to biopsy supernatant was expressed as the increase in afferent discharge over the 12-minute application period (see Supplemental Methods, Supplementary Digital Content 2, http://links.lww.com/CTG/A510).

Power and statistical analysis
All analysis was performed using Graphpad Prism Version 7 and above (GraphPad Software, San Diego, CA). Unless otherwise stated, data are expressed as mean ± SD. Normality of data was tested by using the D’Agostino and Pearson omnibus normality tests, and comparisons were made between parameters for individual patients using Pearson or Spearman correlation coefficients or between group data using a Student t-test or Mann-Whitney U test for parametric and nonparametric data, respectively. Adjustments were made to significant data sets for greater than 3 multiple comparisons using a Bonferroni correction and false discoveries highlighted. Significance was set at $P < 0.05$ or smaller.

RESULTS

Patient details
Symptom scores, psychological tests, and sigmoid biopsies were obtained from 42 patients with IBS-D. Clinical details are shown in Table 1.

Symptom scores
**Bowel-related symptoms and their correlation with pain severity.** As required for trial entry, patients recorded frequent loose stools (Table 1). Pain severity and bloating scores lay within the mild-to-moderate ranges with slightly higher urgency scores. The mean daily duration of pain experienced was 2.9 ± 3.2 hours, with the wide SD highlighting the heterogeneous nature of pain experienced by patients, ranging from brief periods of pain to more prolonged periods of pain. Pain severity was strongly correlated with daily pain duration and urgency and to a lesser degree with bloating and stool frequency, but not with stool consistency (Table 1 and Figure 1).

As expected, patient symptom scores for urgency also correlated significantly with stool consistency (Figure 2), but not with stool frequency or bloating. There was no significant correlation between symptom scores for bloating, stool frequency, or stool consistency (Table 2).

**Psychological scores and their correlation with pain severity and bowel-related symptoms.** Anxiety, depression, and Patient Health Questionnaire-12 Somatic Symptom scores showed significant psychological distress with 24/42 (57%), 10/42 (24%), and 13/42 (31%) of patients having values above the normal cutoffs (7, 7, and 6, respectively) (20,21). However, no significant correlation was found between these and pain severity (Table 3) or other bowel-related symptoms (see Table S1, Supplementary Digital content 1, http://links.lww.com/CTG/A509); although as expected, these psychological measures were highly correlated with each other (see Figure S1, Supplementary Digital content 3, http://links.lww.com/CTG/A508 and Table S2, Supplementary Digital content 1, http://links.lww.com/CTG/A509).
**Effect of age and sex on bowel-related and psychological symptom scores.** No difference was observed in the magnitude of bowel-related symptoms or psychological scores between male and female patients. Furthermore, no correlation was observed between any symptom score and patient age (see Table S3, Supplementary Digital content 1, http://links.lww.com/CTG/A509). No significant correlation was found between biopsy mediator levels or histology, and no significance was observed in the magnitude of bowel-related symptoms or psychological scores between male and female patients. Furthermore, no correlation was observed between any symptom score and patient age (see Table S3, Supplementary Digital content 1, http://links.lww.com/CTG/A509).

**Sigmoid biopsies**

**Correlation of pain severity scores with biopsy mediator levels and histology.** No significant correlation was observed between pain severity scores and biopsy supernatant levels of histamine, tryptase, chymase, and CPA3 nor with biopsy mast cell or CD68 positive cell counts (see Table S4, Supplementary Digital content 1, http://links.lww.com/CTG/A509). Furthermore, no significant correlation was found between biopsy mediator levels or histology and other bowel-related symptoms (see Table S5, Supplementary Digital content 1, http://links.lww.com/CTG/A509) or psychological scores (see Table S6, Supplementary Digital content 1, http://links.lww.com/CTG/A509). It is perhaps worth noting that there was a correlation between CPA3 and urgency and stool consistency (Table S5, Supplementary Digital content 1, http://links.lww.com/CTG/A509), but after correction for multiple comparisons, this failed to reach conventional significance.

**Correlation of biopsy-evoked colonic afferent activity with pain severity scores.** By contrast, application of biopsy supernatant to the receptive field of colonic afferents produced colonic afferent responses that correlated significantly \((P < 0.008)\) with the severity of pain experienced by the patient from which the biopsy was obtained (Figure 3, Table 4). No correlation was found between the change in colonic afferent mechanosensitivity \((0.6 \, g \text{ and } 1.0 \, g \text{ vFh})\) after supernatant application and pain severity scores (Table 4).

**Correlation of colonic afferent activity with biopsy mediator levels.** No significant correlation was observed between biopsy supernatants mediator levels or biopsy histology findings and respective magnitudes of colonic afferent activation or change in colonic afferent mechanosensitivity (see Table S7, Supplementary Digital content 1, http://links.lww.com/CTG/A509). This suggests that multiple mediators may be responsible for the pronociceptive potential of biopsy supernatants, and these mediators may vary from patient to patient.

**Effect of age and sex on biopsy responses.** Furthermore, no significant difference was found in the colonic afferent response to the application of biopsy supernatant or subsequent change in mechanosensitivity based on the sex of the patient from which the biopsy was taken (see Table S8, Supplementary Digital content 1, http://links.lww.com/CTG/A509). In addition, no effect of patient sex was found on biopsy mediator levels or histology, and no correlation was found between the patients’ age and the effect of biopsy supernatant on colonic afferent activity and mechanosensitivity or age and biopsy mediator release or histology (see Table S8, Supplementary Digital content 1, http://links.lww.com/CTG/A509).

### Table 2. Bowel-related symptom scores and their correlation with pain severity

<table>
<thead>
<tr>
<th>Symptom Score</th>
<th>Mean**</th>
<th>SD</th>
<th>N</th>
<th>Pain severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain severity score (0–10)</td>
<td>3.9</td>
<td>2.0</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>Daily pain duration h/d</td>
<td>2.9</td>
<td>3.2</td>
<td>42</td>
<td>(r = 0.67, P &lt; 0.00001^b)</td>
</tr>
<tr>
<td>Urgency score (0–10)</td>
<td>5.6</td>
<td>1.7</td>
<td>42</td>
<td>(r = 0.57, P &lt; 0.0005^c)</td>
</tr>
<tr>
<td>Bloating score (0–10)</td>
<td>3.7</td>
<td>2.5</td>
<td>42</td>
<td>(r = 0.40, P &lt; 0.05^d)</td>
</tr>
<tr>
<td>Stool frequency BM/d</td>
<td>3.8</td>
<td>1.8</td>
<td>42</td>
<td>(r = 0.36, \text{N.S.}^c)</td>
</tr>
<tr>
<td>Stool consistency (BSF score)</td>
<td>5.4</td>
<td>0.7</td>
<td>42</td>
<td>(r = 0.02, \text{N.S.}^c)</td>
</tr>
</tbody>
</table>

**P** values are adjusted for multiple comparisons using a Bonferroni correction and significance was set at \(P < 0.05\). \(\text{N.S.}\) denotes nonsignificant comparisons with false discoveries significant before correction highlighted by \(\text{N.S.}\).

**b**Denotes statistical comparison using Pearson.

**a**Denotes statistical comparison using Spearman.

**Significant results shown in bold text.**

### Table 3. Correlation of patient symptom scores for urgency, bloating, stool frequency, or stool consistency

<table>
<thead>
<tr>
<th>Symptom Score</th>
<th>Urgency</th>
<th>Bloating</th>
<th>Stool frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stool consistency</td>
<td>(r = 0.45, P &lt; 0.02^a)</td>
<td>(r = -0.10, \text{N.S.}^a)</td>
<td>(r = 0.23, \text{N.S.}^b)</td>
</tr>
<tr>
<td>Stool frequency</td>
<td>(r = 0.22, \text{N.S.}^b)</td>
<td>(r = -0.20, \text{N.S.}^b)</td>
<td>—</td>
</tr>
<tr>
<td>Bloating</td>
<td>(r = 0.17, \text{N.S.}^a)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

**Significant results shown in bold text.**

**a**Denotes statistical comparison using Pearson.

**b**Spearman correlation coefficient \((r)\). \(P\) values are adjusted for multiple comparisons using a Bonferroni correction, and significance was set at \(P < 0.05\). \(\text{N.S.}\) denotes nonsignificant comparisons.
Effect of NaV1.9 deletion on biopsy-mediated colonic afferent responses. Given the lack of correlation of nerve response with individual mediators in the supernatant, an alternative strategy to the treatment of abdominal pain in IBS-D may be to target ion channels responsible for the activation of colonic afferents by multiple mediators. NaV1.9 is one such channel that is responsible for the sensitization of colonic afferents in response to inflammatory and algogenic mediators (24,28) consistent with the observation that gain of function human NaV1.9 mutants display episodic abdominal pain and diarrhea (28). To highlight the therapeutic potential of NaV1.9, we also evaluated the effect of biopsy supernatants from patients with the highest pain scores (severity score of 5 or greater) on colonic afferent activity in tissue from NaV1.9 −/− mice. We demonstrated a reduced afferent response by comparison to the responses observed when supernatants were tested in wild type tissue (Figure 4a). Furthermore, although colonic afferent mechanosensitivity to von Frey probing was comparable in tissue from C7B6 mice or NaV1.9 −/− mice (Figure 4b), the change in mechanosensitivity after supernatant application was also significantly reduced in tissue from NaV1.9 −/− mice (Figure 4c).

DISCUSSION
The aim of this study was to gain insight into putative mechanisms of visceral pain in IBS-D. Although previous studies have shown separately that central psychological factors such as mood and somatization (2) and peripheral mediators (8,9) can contribute to visceral pain, we have assessed these factors within a single study. An additional strength of our study is that it has been conducted in a single subtype of IBS, using bowel symptom scores recorded in a daily diary rather than retrospective symptom scores across a mixture of IBS subtypes. We have been able to show that although variable, reported pain severity was highly correlated with the overall daily duration of pain. Urgency, a key feature of IBS-D, was also found to be strongly correlated with pain severity, which is perhaps unsurprising, given that urgency is also believed to be driven by colorectal hypersensitivity and points toward a common pathology.

However, we found no correlation between pain severity scores and either anxiety, depression, nor somatic sensitivity as assessed by the PHQ-12SS. Our sample was representative of all patients with IBS because the mean scores and the proportion scoring above the upper limit of normal for anxiety and depression were very similar to the published data from a much larger IBS-D patient cohort (21). As a consequence, our findings indicate that such psychological factors are not the major determinant of pain severity in our IBS-D patient cohort. This does not however exclude a contribution from psychological factors to pain in some patients with more marked psychological disturbances who may not be selected for clinical trials.

<p>| Table 4. Colonic afferent activity and evoked mechanosensitivity after supernatant application and their correlation with respective pain severity scores |
|----------------------------------|-----------------|-----------------|-----------------|-----------------|------------------|</p>
<table>
<thead>
<tr>
<th></th>
<th>Mean ± SD</th>
<th>Change</th>
<th>N</th>
<th>Pain severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colonic afferent activity (spikes per application)</td>
<td>116.3 ± 155.1</td>
<td>333.0 ± 333.2</td>
<td>216.7 ± 247.7</td>
<td>42</td>
</tr>
<tr>
<td>0.6g vFh (spikes/s)</td>
<td>8.4 ± 5.0</td>
<td>9.9 ± 6.2</td>
<td>1.5 ± 4.3</td>
<td>37</td>
</tr>
<tr>
<td>1.0g vFh (spikes/s)</td>
<td>9.8 ± 3.0</td>
<td>10.5 ± 4.0</td>
<td>0.7 ± 3.6</td>
<td>34</td>
</tr>
</tbody>
</table>

Significant results shown in bold text.
Distribution of the data is indicated by a for normally and b for non-normally distributed.
N.S., nonsignificant; r, Pearson correlation coefficient; vFh, von Frey hair.
Our data instead point to a consistent contribution of peripheral factors to pain severity in patients with IBS-D. We examined the effect of biopsy supernatants on colonic afferent activity in a population of lumbar splanchnic afferents classified as vascular afferents and previously shown to display a nociceptor phenotype (26). Recordings were performed from the lumbar splanchnic nerve because this pathway has previously been shown to be responsible for the transmission of pain from the sigmoid colon (29), our site of biopsy collection. Consistent with our hypothesis, we found a strong correlation between biopsy-mediated colonic afferent activation and patient pain severity. Although we found no correlation between individual biopsy mediator levels and pain scores, we speculate that this reflects the range of possible mediators (e.g., histamine, serotonin, PGE2, and tryptase) (8,9,15) that may differ from patient to patient. An alternative explanation for the lack of correlation would be that other unmeasured mediators are important, such as bile acid derivatives, short-chain fatty acids, lipopolysaccharide, or other microbial metabolites. One possible therapeutic approach would be to generally suppress neural activation in the periphery while avoiding the side effects associated with actions on higher centers (30). To illustrate the utility of this approach, we also examined the effect of biopsy supernatants from patients with high pain severity scores on colonic afferent activity in tissue from NaV1.9 /− mice, a channel highly expressed in colonic afferents (31), and possibly implicated in IBS-D visceral nociception because episodic abdominal pain and diarrhea has been reported in the gain of function human mutants (28). Consistent with our previous findings that colonic afferent response to algogenic mediators and supernatants generated from inflammatory bowel disease (IBD) patient tissue are attenuated in tissue from NaV1.9 /− mice (24,28), we also observed a significant reduction in the response to IBS-D biopsy supernatants in NaV1.9 mouse tissue.

A further observation from this study was the significant correlation between urgency scores and stool consistency, suggesting that looser stools may contain mediators such as bile acids (32) and fecal proteases (33,34) that could promote urgency (35) by stimulating colorectal afferents (36,37). This concept is supported by previous reports that urgency correlates with fecal tryptase (38). Future studies are now warranted to explore these possibilities and the effect of biopsy supernatants on afferent fiber subtypes responsible for the perception of urgency such as pelvic afferent fibers within the colorectum.

In conclusion, using a tightly defined patient group, we were able to show a strong correlation between pain severity and the stimulation of colonic afferent activity by biopsy supernatants, suggesting that in this patient group, the peripheral influences are more significant than central ones. However, we found no evidence for the dominant role of any one of the mediators examined suggesting that either there is another, as yet unmeasured, mediator or that targeting multiple mediator pathways may be a better strategy than targeting a single specific pathway. Future studies should include larger panels of potential mediators that may be present in the stool of patients with IBS.

CONFLICTS OF INTEREST

Guarantor of the article: Robin Spiller, FRCP.

Specific author contributions: Vincent Cibert-Goton, PhD, and Ching Lam, PhD, MRCP, are joint first author. David C. Bulmer, PhD, and Robin Spiller, FRCP, are joint senior author. All authors contributed important intellectual content during manuscript drafting and revision and approved the final draft. Research idea and study design C.L., R.S., D.C.B., and V.C.-G. Data collection and generation of transgenic mice C.L., V.C.-G., M.Y., Y.F., and J.N.W. Data analysis C.L., R.S., D.C.B., and V.C.-G.

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Study Highlights

WHAT IS KNOWN
- Peripheral and psychological factors contribute to symptomology in irritable bowel syndrome (IBS).
- Peripheral factors contribute to pain in IBS by stimulating sensory nerves.
- Psychological factors contribute to pain in IBS by promoting hypervigilance and increasing stress responses.

WHAT IS NEW HERE
- Pain severity scores correlated with daily duration of pain, and urgency, but not with anxiety, depression, or somatization in patients with IBS-D.
- Pain severity scores correlated with biopsy-mediated colonic afferent firing.
- Individual biopsy mediator levels did not predict pain severity.

TRANSLATIONAL IMPACT
- Nerve targeting therapies may be more effective than specific mediator antagonists.

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