A MATTER OF OPINION

Bacterial Overgrowth and Irritable Bowel Syndrome: Unifying Hypothesis or a Spurious Consequence of Proton Pump Inhibitors?

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Some studies indicate that small intestinal bacterial overgrowth (SIBO), as measured by hydrogen breath tests (HBT), is more prevalent in patients with irritable bowel syndrome (IBS) vs. matched controls without IBS. Although the data are conflicting, this observation has led to the hypothesis that SIBO may be a primary cause of IBS. Yet, it remains unclear whether SIBO is truly fundamental to the pathophysiology of IBS, or is instead a mere epiphenomenon or bystander of something else altogether. We hypothesize that SIBO might be a byproduct of the disproportionate use of proton pump inhibitors (PPIs) in IBS, as follows: (1) IBS patients are more likely than controls to receive PPI therapy; (2) PPI therapy may promote varying forms of SIBO by eliminating gastric acid; and (3) existing studies linking SIBO to IBS have not adjusted for or excluded the use of PPI therapy. When linked together, these premises form the basis for a simple and testable hypothesis: the relationship between SIBO and IBS may be confounded by PPIs. Our article explores these premises, lays out the argument supporting this “PPI hypothesis,” discusses potential implications, and outlines next steps to further investigate this possibility.

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SMALL INTESTINAL BACTERIAL OVERGROWTH (SIBO) AND IRRITABLE BOWEL SYNDROME (IBS)

IBS is a condition of unknown etiology that presents with a recurrent abdominal pain or discomfort along with abnormalities in stool frequency or form (1). Because the symptoms of IBS overlap with SIBO (diarrhea, constipation, bloating, gas, and pain), it has been hypothesized that many patients with IBS have underlying SIBO (2–6). In fact, some believe that the diagnosis of IBS should be questioned in patients not found to have SIBO by diagnostic testing, or in those patients failing to respond to appropriate antibiotic therapy (7). This causal theory has gained traction with the publication of studies indicating that SIBO, as measured by imperfect surrogate tests (8) (e.g., glucose hydrogen breath test [GHBT], lactulose HBT, jejunal aspirate), is more prevalent in patients with IBS than matched controls. For example, Pimentel et al. reported that 84% of IBS patients are LHBT-positive compared to only 20% of healthy controls (6). Notably, other investigators have not detected a significant difference in LHBT positivity between the groups (9, 10). More recently, Posserud and colleagues found no difference in jejunal aspirate yield between IBS and control patients when adopting the 10^5 colony forming units (CFU) threshold, but did find that mildly increased bacterial counts (using a lower 10^3 CFU cut off) were more common in IBS than controls (10). Kassinen et al. found that fecal microbiota of IBS subjects (as measured by DNA fingerprinting) differed significantly from healthy subjects, although there was no reported difference in the overall bacterial count (11). The relationship between IBS and SIBO is further supported by the data that IBS patients treated with a short course of antibiotics are more likely to experience symptom improvement versus patients receiving placebo (12). Although some investigators contend that this effect is based on the eradication of SIBO in IBS, the theory remains to be definitively proven.

COULD SIBO BE AN EPICENPHENOMENON OF ANOTHER UNMEASURED FACTOR IN IBS?

Although there are data to support the relationship between SIBO and IBS, the SIBO theory is potentially limited in its ability to fully explain other proposed models of IBS, such as the biopsychosocial, visceral sensitivity, inflammatory, or neurohormonal models, among others (13). For example, it remains unclear how IBS symptoms could improve with nonpharmacological interventions (14–26), yet at the same time, be a fundamentally infectious disease. The observation that a subset of IBS patients consistently benefits from the nonpharmacological therapies distinguishes it from a condition like pneumonia and suggests that a cause-and-effect disease paradigm remains elusive. In other words, it seems unlikely that antibiotics alone will provide an answer for more than a
subgroup of IBS patients. The lack of a strong relationship between SIBO and IBS is also supported by the observations that an antibiotic treatment in patients with IBS can relieve symptoms in a subset of patients without SIBO (27). Moreover, there are conflicting data that the eradication of SIBO correlates with symptom relief. Thus, the beneficial effects of antibiotics in some patients may not be related to the eradication of SIBO, but instead may be from another mechanism such as an antimicrobial effect on pathogenic organisms in the gut (27). As the SIBO hypothesis does not appear to unify multiple competing hypotheses, it raises the question of whether SIBO is truly fundamental to the pathophysiologic basis of IBS, or whether SIBO is a mere bystander, or even an epiphenomenon of other processes. The lack of consistency in the data linking SIBO to IBS raises the possibility that some other factor may be operating in the background. In other words, when data fail to converge in support of a hypothesis, it is reasonable to consider whether the null hypothesis is true instead, and that variations in the data merely reflect variations in other factors extrinsic to the relationship being tested. Although the inconsistent results linking SIBO to IBS could be a consequence of varying study methodologies, different local SIBO prevalence or disparate definitions of IBS, it may also simply reflect the presence of an external risk factor for SIBO that travels along with IBS but is not, in fact, intrinsic to IBS at all. In other words, some as yet unmeasured factor might confound the relationship between IBS and SIBO.

**COULD IBS BE LINKED TO SIBO THROUGH PROTON PUMP INHIBITORS (PPIs)?**

One simple and prevalent variable may fit the bill: PPIs. We hypothesize that the relationship between IBS and SIBO could potentially be confounded by the use of PPIs, as follows: (1) IBS patients are more likely than controls to receive PPI therapy (2), PPI therapy may promote varying forms of SIBO by eliminating gastric acid, and (3) the existing studies linking SIBO to IBS have not adjusted for or excluded the use of PPI therapy. When linked together, these premises form the basis for a simple and testable hypothesis: the relationship between SIBO and IBS may be confounded by PPIs (Fig. 1). We explore each premise in more detail below.

**IBS Patients Are More Likely Than Controls to Receive Long-Term PPI Therapy**

Up to 40% of patients with IBS have comorbid gastroesophageal reflux disease (GERD) (28), and 30–50% have overlapping dyspepsia (29, 30). Conversely, one-half of patients with GERD have comorbid IBS (28, 31). Because patients with IBS are more likely to have GERD and dyspepsia versus matched controls, they are also more likely to receive PPI therapy. Moreover, overuse of PPI therapy is common, and is often triggered by an unexplained abdominal pain. Because IBS patients have a long-standing and often difficult-to-treat abdominal pain, coupled with the fact that GERD and dyspepsia commonly overlap, chronic PPI therapy in IBS patients is extremely common in everyday clinical practice. This is supported by a recent cohort study revealing that 44% of patients with IBS were receiving a PPI (32)—a percentage that is much higher than non-IBS healthy controls, most of whom do not take regular PPI therapy. Thus, it is notable that IBS and PPI use are inexorably linked because IBS patients are highly enriched with PPI users.

**PPI Therapy Can Promote SIBO**

The high prevalence of PPI use in IBS would be irrelevant if PPI use were not, in fact, related to the outcome of interest—SIBO. However, PPIs are potent antisecretories, and hypochlorhydria is a risk factor for SIBO (33). The existence of gastric acid has a teleological explanation in that it serves as the primary defense against enteric infection. Thus, it comes as no surprise that removing this natural defense inevitably leads to perturbations in enteric flora—some clinically significant, some not. It has long been established that PPI therapy can alter gastric, duodenal, and intestinal bacterial profiles. For example, Thorens et al. randomized 47 patients with peptic ulcer to receive 4 wk of cimetidine versus omeprazole, and subsequently, cultured duodenal juice obtained during follow-up endoscopy (34). The authors found a higher incidence of bacterial overgrowth in the omeprazole arm (53% vs 17%). This finding was duplicated by Fried et al., who further demonstrated that PPI-related SIBO was due to both oral and colonic-type bacteria—not merely oral flora alone (35). Theisen and colleagues found that the suppression of gastric acid with omeprazole led to a high prevalence of SIBO, which, in turn, led to a markedly increased concentration of unconjugated bile acids (36). Moreover, Lewis et al. documented that omeprazole-related SIBO was associated with shorter intestinal transit times (37). These studies suggest that PPI-related SIBO could potentially lead to symptoms of IBS, such as diarrhea, as a result of an increased osmotic load from bile acids coupled with more rapid intestinal transit. It is notable that the most common side effects...
of PPIs include abdominal pain, bloating, flatulence, constipation, and diarrhea—symptoms that overlap with IBS and occur in up to 5% of PPI users.

Very few studies have investigated the relationship between PPI use and SIBO in IBS. Recently, Majewski and colleagues reported data on a cohort of 204 patients with IBS undergoing GHBT for SIBO, some of whom were receiving concurrent PPI therapy (32). The authors found that PPI use was higher in GHBT-positive patients (48% on a PPI) compared to GHBT-negative patients (39% on a PPI). Although this difference was not statistically significant ($P = 0.2$), the study was not powered to measure the impact of PPI therapy on GHBT results, nor did it measure the dose–response relationship to compare amount and duration of PPI exposure to GHBT positivity. However, the study provides initial pilot data, with a numerical trend supporting a potential relationship, and emphasizes the need to perform a larger study to overcome a potential type II error. Moreover, recent data indicate that, among patients with GHBT positivity (including patients with IBS) receiving rifaximin for eradication, the re-growth of SIBO is independently predicted by the use of concurrent PPI therapy (38). Thus, not only might PPI therapy lead to SIBO in some patients with IBS, but also the recurrence of SIBO following antibiotic therapy might be accelerated in the setting of PPI therapy. In other words, so long as the risk factor for SIBO is present, the condition may recur despite temporary removal with antibiotics.

In considering this line of inquiry, it is important to distinguish the varying types of enteric infections related to PPI therapy. Skeptics might contend that PPIs are unlikely to confound the relationship between IBS and SIBO, chiefly because infectious complications are rare events. Indeed, the evidence-based reviews conclude that PPI-related bacterial overgrowth infrequently leads to clinically important disease (34, 39). However, these reviews have focused on overt infections such as Shigella, Salmonella, Yersinia, and Clostridium difficile (C. difficile) colitis—not merely HBT positivity. It is possible that a broader spectrum of PPI effects exists, akin to an “iceberg phenomenon,” with rare but observable events above the waterline and common yet covert events below the waterline (Fig. 2). According to this model, PPIs can cause rare yet clinically dramatic enteric infections (e.g., C. difficile colitis) that are detected above the waterline. These “JAMA-worthy events” (40) are truly rare (far below 1%), but are proof of principle that profound acid suppression can meaningfully alter the enteric flora in susceptible individuals. But what about the 5% of patients that develop IBS-type symptoms after initiation of PPI therapy? These patients may have underlying SIBO with resulting dyspepsia, abdominal pain, bloating, flatulence, diarrhea, and/or constipation—still a rare group, but an intriguing group because of the overlap between IBS and PPI-related symptoms. And below this group might reside a much larger population of PPI users with altered intestinal flora, but without clinically overt symptoms. This group might only be detected with HBT, or with highly sensitive tests for altered enteric flora (e.g., stool DNA fingerprinting). Given what we know about the profound impact of PPI therapy on gastric acid secretion, coupled with the knowledge that hypochlorhydria can alter the enteric flora, it is not hard to imagine that a highly tuned assay like stool DNA fingerprinting might detect minor differences in the flora between PPI users and non-PPI users.

**Studies Linking SIBO to IBS Have Not Excluded PPI Users**

This line of inquiry would be moot if the existing studies linking SIBO to IBS had excluded PPI users. That is, if the linkage were found in the absence of PPI use, then it would systematically exclude PPI exposure as a confounding influence. However, the studies reporting higher rates of SIBO in IBS, including those using HBT (3–6), jejunal aspirate (9), and DNA fingerprinting (10), have not explicitly excluded or adjusted for PPI users. Moreover, none of the studies report the prevalence of PPI exposure in the IBS versus healthy control groups, making it impossible to judge whether PPI exposure could have played any role in influencing the results. This appears to be an important oversight as PPI use is highly prevalent in patients with IBS and is a known risk factor for SIBO. Yet, the published studies are meticulous about excluding other risk factors that are rare in IBS, such as cirrhosis, inflammatory bowel disease, and connective tissue disorders, among others (11). It could be argued that of all the potential confounders, short of previous antibiotic therapy, the use of PPIs should be considered among the most important potential confounders, given its high prevalence in the target population and its association with the development of SIBO.
The “PPI hypothesis” remains untested. Nonetheless, if it were true, then it would suggest that SIBO may not be fundamental to the pathophysiology of IBS, and instead may sometimes be a mere byproduct of treatment with PPIs. It would not indicate that PPIs cause IBS. Instead, it would suggest that PPIs might exacerbate IBS or merely alter the intestinal flora in a subclinical manner. This, in turn, could yield a “red herring” of HBT positivity, which might be falsely interpreted as causal of IBS when it is instead a mere bystander. In addition, it would not definitively prove that IBS is unrelated to SIBO, but would suggest that the studies demonstrating higher rates of SIBO in IBS versus healthy controls would need repeating with careful exclusion or adjustment for PPI status. This would also apply to randomized controlled trials of antibiotic therapy in IBS, where the benefits of active treatment might simply reflect, at least in part, a temporary reversal of PPI-related symptoms superimposed on underlying IBS.

Future research should include a prospective evaluation to measure the dose–response relationship between PPI exposure and SIBO in patients with IBS. If that were positive, the additional work might also include a randomized withdrawal study in IBS patients on PPI therapy (in the absence of concurrent acid-peptic disorders otherwise warranting PPIs). If there were a meaningful change in the bowel symptoms between patients switching to placebo versus those staying on active PPI, then it would indicate that PPIs play some role in exacerbating or propagating IBS symptoms and suggest that PPI withdrawal should be considered prior to initiating antibiotics. Although this PPI hypothesis might seem naïve or overly simplistic, it is worth recalling the wisdom of Sir William Ockham, who surmised even in the 14th century that, among competing solutions to a problem, the solution with the fewest steps, postulates, or entities is generally preferred. In other words, as complexity rises, so does burden of proof. We believe this theory is both tenable and testable.


REFERENCES


CONFLICT OF INTEREST

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