Interrogating the Gut-Brain Axis in the Context of Inflammatory Bowel Disease: A Translational Approach

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This review examines preclinical and clinical studies relevant to our understanding of how the bidirectional gut-brain axis influences the natural history of inflammatory bowel disease. Preclinical studies provide proof of concept that preexisting behavioral illness, such as depression, results in increased susceptibility to inflammatory stimuli and that commonly used classes of antidepressants protect against this vulnerability. However, clinical studies suggesting behavioral illness as a risk factor for IBD and a protective role for antidepressants have relied primarily on symptom-reporting rather than objective measurements of inflammation. In terms of gut-to-brain signaling, there is emerging evidence from preclinical and clinical observation that intestinal inflammation alters brain functions, including the induction of mood disorders, alteration of circadian rhythm both centrally and peripherally, and changes in appetitive behaviors. Furthermore, preclinical studies suggest that effective treatment of intestinal inflammation improves associated behavioral impairment. Taken together, the findings of this review encourage a holistic approach to the management of patients with IBD, accommodating lifestyle issues that include the avoidance of sleep deprivation, optimized nutrition, and the monitoring and appropriate management of behavioral disorders. The review also acknowledges the need for better-designed clinical studies evaluating the impact of behavioral disorders and their treatments on the natural history of IBD, utilizing hard end points to assess changes in the inflammatory process as opposed to reliance on symptom-based assessments. The findings of the review also encourage a better understanding of changes in brain function and circadian rhythm induced by intestinal inflammation.

Key Words: depression, antidepressants, circadian rhythm, inflammation, vagus nerve, cytokines

BACKGROUND

From a historical perspective, our understanding of the natures of ulcerative colitis and Crohn’s disease have followed quite different trajectories. Ulcerative colitis was recognized much earlier than Crohn’s disease and was originally considered to be a psychosomatic disorder, or one in which behavioral issues played a significant etiological role. This line of thinking persisted until the latter half of the 20th century. In contrast, “regional ileitis” or Crohn’s disease was considered an organic disease, due in part to the fact that surgeons managed most cases in the early years. Nevertheless, as our understanding of the nature and extent of brain-gut communication

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autonomic nervous system, and the limbic system. In the recent past, the intestinal microbiota has become incorporated into the bidirectional gut-brain axis and provides a pathway that links intestinal bacteria and the brain.

Since the discovery of the inflammatory reflex by Tracy and colleagues, there has been considerable interest in the role of the vagus nerve in IBD. A preclinical study by Ghia and colleagues demonstrated the protective effect of the vagus nerve in a model of intestinal inflammation in the absence or presence of psychiatric comorbidity. In addition, preclinical studies of the efficacy of vagal nerve stimulation have shown therapeutic benefit in models of colitis. The first pilot clinical trial has demonstrated symptomatic and endoscopic improvement in 5 of 7 patients with Crohn’s disease involving the terminal ileum; this was accompanied by a reduction in C-reactive protein of 7 patients with Crohn’s disease involving the terminal ileum; this was accompanied by a reduction in C-reactive protein of 7 patients with Crohn’s disease involving the terminal ileum; this was accompanied by a reduction in C-reactive protein of 7 patients with Crohn’s disease involving the terminal ileum; this was accompanied by a reduction in C-reactive protein of 7 patients with Crohn’s disease involving the terminal ileum; this was accompanied by a reduction in C-reactive protein.

Studies demonstrating that patients with IBD exhibit evidence of anxiety and/or depression more frequently than healthy controls have rekindled interest in the role of behavioral factors in these conditions. For example, the Manitoba IBD cohort study was the first population-based study to examine the prevalence of common psychiatric disorders among IBD patients and showed that the 12-month prevalence of major depression in a population-based cohort of subjects with IBD was 9.1% vs 5.5% in community controls. In addition, the same group identified that the lifetime prevalence of major depression in the subjects with IBD was 27.2% vs 12.3% in controls. A systematic review showed prevalence rates of 19.1% vs 9.6% for anxiety and 28.2% vs 13.4% for depression, and the rates were positively influenced by the activity of IBD. Other studies have found higher than expected prevalence rates for anxiety and depression in IBD patients, more commonly often in Crohn’s disease than ulcerative colitis, and in patients with active disease. Placebo response rates average 19% in clinical trials evaluating a broad spectrum of treatments, including biological agents and this also prompts interest in the role of psychiatric comorbidity. In addition, preclinical studies of the efficacy of vagal nerve stimulation have shown therapeutic benefit in models of colitis.

Several models have demonstrated that preexisting depression increases vulnerability to intestinal inflammatory stimuli. Maternal separation among rodents is a well-established model in which the maternally deprived offspring exhibit depression-like and anxiety-like behaviors in adulthood and increased susceptibility to stress as a result of hyperresponsiveness of the Corticotrophin Releasing Factor (CRF)-mediated hypothalamic-pituitary axis. A study showed that maternally separated mice exhibited depression-like behavior at 8 weeks of age and developed increased intestinal permeability but no evidence of spontaneous inflammation in the gut. Colitis was then induced by dextran sodium sulphate (DSS), and maternally separated mice experienced a more severe inflammatory response than nonseparated control mice. In terms of underlying mechanisms, it is known from preclinical studies that maternal separation alters the composition of the intestinal microbiota and that this, along with host factors contribute to the model’s behavioral phenotype, but the effect of this altered microbiota on susceptibility to colitis has not yet been investigated. In a model of depression induced in mice by intracerebral ventricular injection of low-dose reserpine, this was accompanied by a reduction in C-reactive protein.

BRAIN TO GUT SIGNALING

1. Do emotional disorders predispose individuals to IBD?
2. Do stress and emotional disorders precipitate a relapse of pre-existing IBD?
3. Does effective treatment of emotional disorders prevent or improve IBD?

GUT TO BRAIN SIGNALING

4. Does intestinal inflammation influence the brain?
5. Does intestinal inflammation alter brain function relevant to IBD?
6. What is the impact of IBD-directed therapy on brain & behavior?

FIGURE 1. The structure of this review is illustrated. It will address 6 questions pertaining to the role of the bidirectional gut-brain axis in the natural history of IBD. Each question is addressed by first examining evidence of biologic feasibility based on preclinical studies, followed by a review of the existing clinical literature pertinent to the question.
depressed mice were more susceptible to the DSS-induced colitis. A study in rats has also demonstrated the vulnerability to experimental colitis in depressed rodents. Taken together, these results provide proof of concept that preexisting depression-like behavior increases vulnerability inflammatory stimuli and implicate alterations in intestinal permeability and attenuation of the vagal anti-inflammatory pathways among underlying mechanisms. If these results can be extrapolated to humans, they predict that depression is a risk factor for the development of intestinal inflammatory conditions.

Observations from clinical studies

The Manitoba IBD cohort study demonstrated that 64% of IBD patients reporting a lifetime anxiety or mood disorder received a psychiatric diagnosis at least 2 years before the onset of IBD; 54% of patients with a mood disorder experienced the first episode of depression at least 2 years before the diagnosis of IBD. A subsequent study utilizing the Manitoba Population Research Data Repository examined the risk of developing immune-mediated inflammatory disease (IMID), including IBD, following a previous diagnosis of a psychiatric disorder. In the IBD cohort of 3766 patients, the incidence of depression, anxiety, and bipolar disorder was increased compared with a matched cohort, and the onset of the psychiatric disorder preceded diagnosis of IBD by as much as 5 years. Taken together, these studies support the view that common psychiatric disorders increase the risk of developing IBD. A similar conclusion was drawn from a recent study investigating the incidence of IBD in 403,665 patients with a new onset diagnosis of depression established between 1986 and 2012. The study found that patients with a history of depression had an increased risk of developing Crohn’s disease (adjusted hazard ratio [HR], 2.11; 95% CI, 1.65–2.70) or ulcerative colitis (HR, 2.23; 95% CI, 1.92–2.60).

Do Stress or Mood Disorders Precipitate Relapse of Preexisting IBD?

Proof of concept from preclinical studies

Preclinical studies show that psychological stress reactivates quiescent inflammation in the gastrointestinal (GI) tract. In one study, restraint stress was applied to mice in histological remission after the induction of acute colitis with dinitrobenzene sulfonic acid (DNBS). Stress alone failed to reactivate colitis. However, when stress was applied together with a subthreshold dose of DNBS, colitis was reactivated. Moreover, this susceptibility to colitis could be adaptively transferred into naïve mice via CD+ve lymphocytes from stressed mice. These findings suggest that stress alone is insufficient to induce relapse in the absence of an overt inflammatory stimulus. Other work has shown that stress increases intestinal permeability via pathways involving cholinergic nerves and the release of corticotropin releasing factor (CRF), and this may be a mechanism whereby stress increases susceptibility to inflammatory stimuli. Experimentally induced depression has shown to reanimate quiescent colitis by attenuating vagally mediated inhibition of pro-inflammatory cytokine secretion from macrophages. In this study, acute colitis was induced by either DSS or DNBS and allowed to become quiescent. Colitis was reactivated by depression, induced by intra-cerebro-ventricular administration of reserpine, and was associated with increased secretion of pro-inflammatory cytokines from peritoneal macrophages mediated via the α-7 subunit of the nicotinic acetylcholine receptor. These findings invoke vagal dysfunction as a mechanism whereby depression influences intestinal inflammatory processes, and tricyclic antidepressants (TCAs) were shown to restore vagal function and reduce intestinal inflammation. Interestingly, vagal dysfunction is well recognized in depression and is the basis for the use of vagal stimulation in treatment-resistant depression.

Do stress or mood disorders precipitate relapse of preexisting IBD? Observations from clinical studies

Depression has been associated with increased unplanned 30-day hospital readmission rates in ulcerative colitis, symptomatic exacerbations, and increased hospitalization and reduced responsiveness to anti-TNF therapy. A prospective observational longitudinal study involving 60 IBD patients followed for 18 months correlated the severity of baseline depression scores with the number of subsequent relapses and a shorter interval to the first clinical recurrence of IBD. A recent prospective cohort study found a significant association between symptoms of depression and clinical evidence of disease recurrence in patients with Crohn’s disease or ulcerative colitis. Symptoms of anxiety also correlated with clinical recurrence in that study. Depression has been associated with poor compliance with anti-inflammatory or immunosuppressive or biological therapies and may be a contributing factor in these reports. Narula et al found that anxiety—but not depression—predicted increased hospitalizations, ER visits, and the need for corticosteroids. A more recent prospective study showed that high anxiety scores during documented remission predicted a more severe disease outcome in patients with ulcerative colitis or Crohn’s disease. The question arises as to whether these results simply reflect increased symptom reporting by patients with depression or anxiety or actual exacerbation of the underlying inflammatory disease. Gracie et al found a poor correlation between symptoms or the Harvey-Bradshaw Index and evidence of mucosal inflammation using fecal calprotectin as a biomarker of the inflammatory process in Crohn’s disease patients but not those with ulcerative colitis. Though preclinical studies link psychopathology with deterioration of IBD-associated symptoms, further prospective studies involving objective measures of inflammation are required to resolve the issue of whether depression or anxiety exacerbate intestinal inflammation or simply promote symptom reporting in IBD patients.
Although clinical experience strongly suggests an association between stressful life events and exacerbations of IBD, support for this association from existing clinical studies is weak. A study by Bernstein et al\textsuperscript{47} demonstrated that perceived stress, but not major stressful life events or low mood, were significantly associated with symptomatic flare of IBD. A subsequent study demonstrated that the association between perceived stress and symptomatic activity did not reflect worsening inflammation as reflected by fecal calprotectin levels of >250 µg/g.\textsuperscript{48} Thus, although the association of stress with exacerbation of the inflammatory process in IBD remains clinically plausible and biologically feasible,\textsuperscript{49} further studies involving objective measurements of the inflammatory process are much needed.

**Does Effective Treatment of Emotional Disorders Prevent or Improve IBD?**

**Proof of concept from preclinical studies**

Preclinical studies have focused on models of depression-like behavior and on the ability of antidepressants to influence the inflammatory process. Varghese et al used a model of maternal separation to induce depression-like behavior in mice. Maternally separated mice exhibited increased intestinal permeability and developed a more severe colitis induced by DSS compared with nonmaternally separated mice. The antidepressant desipramine attenuated depression-like behavior in maternally separated mice, and colitis was less severe in these mice. Importantly, desipramine had no effect on the severity of colitis in nonmaternally separated mice.\textsuperscript{26} These results are consistent with the view that desipramine attenuated the severity of colitis by reducing depression-like behavior, rather than via a direct effect of the drug on the colonic inflammatory process. Desmethylinipramine, a closely related tricyclic antidepressant, was also shown to reduce depression-induced relapses of quiescent colitis induced by previous exposure to DSS. Depression was induced by intracerebro-ventricular administration of reserpine once the DSS colitis became quiescent, and this was accompanied by reactivation of the colitis. Treatment with the antidepressant attenuated both the depression-like behavior and reactivation of colitis. The protective effect of the antidepressant was dependent on the integrity of the vagus nerve, and thus the proposed mechanism of action was the restoration of vagally mediated suppression on cytokine release by macrophages.\textsuperscript{12, 33} Thus, although at least 1 mechanism of action underlying the ability of tricyclic antidepressants to attenuate depression-induced exacerbation of inflammation is via the restoration of vagal modulation of cytokine release from macrophages, other studies also reveal a direct anti-inflammatory effect. In a study involving rodents without induced depression, the tricyclic antidepressant amitriptyline stimulated production of interleukin (IL)-10 and directly attenuated inflammatory responses to abdominal sepsis induced by cecal ligation or lipopolysaccharide (LPS).\textsuperscript{36} Studies on the ability serotonin-modulating antidepressants to attenuate depression-induced exacerbation of experimental colitis in rodents suggest a direct inhibitory effect on the intestinal inflammatory process. Studies using fluvoxamine (a selective serotonin reuptake inhibitor [SSRI]) or venlafaxine (a serotonin-norepinephrine reuptake inhibitor [SNRI]) showed that although these agents attenuated exacerbation of acetic acid–induced colitis in rats depressed by peripheral administration of reserpine, the antidepressants also reduced inflammation in rats not treated with nonreserpine.\textsuperscript{51} Under the same conditions, these investigators also demonstrated that the tricyclic amitriptyline reduced colitis in nonreserpine-treated rodents.\textsuperscript{28} These results differ from others in which TCAs failed to attenuate colitis in the absence of depression.\textsuperscript{50}

Taken together, these preclinical results indicate that several classes of antidepressants are able to attenuate the increased severity of de novo colitis or exacerbations of preexisting colitis induced by depression, but mechanisms of action differ depending on the model under study and may include a direct anti-inflammatory effect.

**Clinical observations on the impact of treatment of behavioral disorders on IBD**

Up to 30% of IBD patients receive antidepressant therapy, with selective serotonin reuptake inhibitors and tricyclic antidepressants being the most commonly prescribed treatments.\textsuperscript{52} Indeed, the use of antidepressants is more common in patients suffering from IBD compared with a control population, and this is seen in both adolescents and adults.\textsuperscript{53, 54} Despite this usage of antidepressants in IBD, surveyed gastroenterologists expressed skepticism regarding the impact of this drug category on the natural history of the disease.\textsuperscript{55} It is not surprising, therefore, to find few studies that examine this relationship. A retrospective review of the records of 29 IBD patients 1 year before and 1 year after commencing antidepressant therapy showed that there were fewer steroid courses, endoscopies, and relapses during antidepressant therapy compared with the year prior.\textsuperscript{56} A review of records of 42,890 IBD patients in the Danish national database found that the incident rate of disease activity was lower among patients on antidepressants, and this was prominent in those patients with no history of antidepressant use before the onset of IBD.\textsuperscript{57} Interestingly, a showing an increased risk of IBD in over 400,000 patients with recent onset found that the risk was reduced in those patients receiving antidepressant treatment. Specifically, selective seroton reuptake inhibitors (SSRIs) and tricyclic antidepressants appeared protective against the risk of developing Crohn's disease, whereas selective serotonin and norepinephrine reuptake inhibitors (SNRIs) in addition to SSRIs and TCAs seemed protective against ulcerative colitis.\textsuperscript{58} Three systematic reviews evaluating the impact of antidepressants on the course of IBD were unable to draw firm conclusions regarding the benefit of
antidepressants in this context, primarily because of the lack of prospective and randomized trials and particularly those with inflammation-based objective end points. Each of the previously cited papers identified this as the major obstacle to determine whether antidepressants impact the natural history of these chronic intestinal inflammatory diseases or simply attenuate symptom reporting.

There have been 3 randomized controlled trials of cognitive behavioral therapy (CBT) in IBD patients. One study demonstrated an improvement in quality of life, but another study failed to confirm this or an effect on the course of IBD over 24 months. A more recent study found no effect of CBT on disease activity in young patients with mild anxiety or depression. At this time, one cannot conclude that CBT influences the natural history of IBD. There is a limited number of studies on the benefit of hypnosis in the management of IBD. These studies are limited by small sample size but suggest that courses of hypnotherapy can induce remission lasting from 2.5 months to up to 5 years. Interestingly, reduction in the expression of inflammatory cytokines including TNF-α have been observed in the intestine after a single session of hypnotherapy. Many IBD patients use complementary and alternative approaches to manage their condition, and these include herbal therapy, acupuncture, exercise, and mind-body therapy, but the efficacy of these approaches has yet to be determined in well-designed clinical studies.

GUT-TO-BRAIN SIGNALING

The influence of intestinal inflammation of the brain has received relatively little clinical attention in comparison with the brain-to-gut communication in IBD. The interrogation of this axis will, therefore, rely more on an emerging body of preclinical studies rather than clinical observations.

**Question 4: Does Intestinal Inflammation Influence Brain Function and Promote Psychiatric Morbidity?**

*Preclinical studies*

Preclinical studies support the view that intestinal inflammation produces changes in brain functions and behavior. Colitis induced by DNBS induced anxiety and depression-like behaviors, and this was associated with increased hippocampal expression of genes related to inflammation and innate immunity, in addition to a reduction in mitochondrial function. Dextran sodium sulphate–induced colitis resulted in reduced brain excitability and seizure thresholds, anxiety-like behavior in all mice, and hyperalgesia to painful somatic stimuli in male mice. In a study of chronic low-grade colitis induced by a noninvasive nematode infection, anxiety-like behavior was associated with a reduction in brain-derived neurotrophic factor (BDNF) and increased circulating levels of TNF-α and kynurenine, a tryptophan metabolite. Administration of anti-TNF-α normalized behavior but did not normalize BDNF expression. Anxiety induced by colitis was evident in vagotomised mice, suggesting that behavioral changes were a result of circulating TNF-α rather than activation of afferent vagal pathways. Interestingly, interferon-α induced changes in behavior also persisted in vagotomised mice. Other cytokines including interleukin-6 and interferon-γ have also been implicated in inflammation-induced depression or anxiety.

A relative newcomer to the bidirectional brain-gut axis is the intestinal microbiota, and experimentally induced changes in microbiota composition have been shown to alter brain function. The introduction of a bacterial pathogen, Campylobacter jejuni, into the rodent gut was shown to activate the vagus and induce anxiety-like behavior before a local inflammatory response was established. A causal link between anxiety and the microbiota has been suggested in a recent paper in which colonization of germ-free mice with the microbiota of IBS patients with high levels of anxiety induced anxiety-like behavior in the recipient mice, and this was associated with innate immune activation. Similar studies involving the microbiota of IBD patients with high levels of anxiety or depression have not yet been published. Taken together, these preclinical studies demonstrate that intestinal inflammation induces changes in brain function and behavior.

**Clinical observations**

Changes in T cell activation, circulating cytokines, acute phase reactants, oxidative and nitrosative stress, and the serotonin-depleting catabolism of tryptophan have been implicated in the pathogenesis of depression; similar changes have also been documented in IBD. Thus, depression and IBD may share a common inflammatory-based pathophysiology to account for the increased coexistence of these conditions. This is supported by the observations that (1) depression may precede the onset of IBD and (2) clinical disease activity was the only independent risk factor for depression in Crohn's disease patients. Several changes have been described in the brains of IBD patients and include structural changes in white matter and functional changes on MRI. For example, young patients with IBD showed increased activation in brain regions associated with pain perception compared with healthy controls or patients with irritable bowel syndrome. In addition, an MRI study of the brain in patients with stable Crohn's disease and no psychiatric comorbidity demonstrated changes in brain connectivity compared with healthy controls; this included the region involved in processing of emotion and visceral stimuli including pain. However, the relationship between brain changes and the activity of the intestinal inflammatory process has not yet been established.
Question 5: Does Intestinal Inflammation Alter Other Behaviors Relevant to IBD?

Disruption of circadian rhythm and IBD

Preclinical studies. Circadian rhythm governs both behavior and physiology and is determined by a central oscillator in the suprachiasmatic nucleus (SCN) of the hypothalamus where it is cued by exposure to light. The central oscillator synchronizes peripheral clocks via neurohumeral pathways. Several mammalian clock genes have been identified that govern positive and negative transcriptional feedback control loops. There is increasing interest in the circadian control of immune function and inflammatory responses in IBD. Genetic ablation of circadian clock function or environmental circadian rhythm disruption (CRD) in mice disrupted intestinal barrier function and increased susceptibility to DSS-induced colitis. In a recent study, C57BL/6J mice were exposed to circadian shifts for 3 months before the induction of DSS colitis. Circadian shifts induced changes in intestinal permeability aggravated by DSS-induced colitis; this was accompanied and increased by circulating TNF-α, interleukin-1β, interleukin 6, C-reactive protein, and fecal calprotectin. Conversely, marked changes in circadian gene profiles were observed in mice after induction of experimental colitis induced by DSS or TNBS. Thus, results of preclinical studies reveal a vicious cycle in which primary changes in circadian rhythm worsen intestinal inflammatory responses that, in turn, alter the expression of circadian genes. The circadian clock governs other functions that impact IBD patients, and these include sleep and feeding behavior.

Clinical observations

Circadian clock gene expression in IBD. A study in young, recently diagnosed, untreated patients with IBD demonstrated reductions in expression levels of all clock genes, except for PER2, in noninflamed intestinal mucosal tissues from patients compared with controls (P < 0.05). The expression of clock genes (eg, CLOCK, BMAL1, CRY1, CRY2, PER1, and PER2) were lower in white blood cells from recently diagnosed young patients with IBD compared with controls, suggesting that circadian clock gene disruption is a very early feature of IBD. Furthermore, a cohort study suggested that sleep deprivation is a risk factor for the development of ulcerative colitis. Liu et al found that the expression of circadian genes was downregulated in peripheral blood monocytes and in intestinal biopsies from IBD patients, and the findings were more marked in ulcerative colitis patients. Circadian gene expression was inversely related with assessments of disease activity and acute phase reactants. Another study linked polymorphism of the PERIOD3 (PER3) clock gene with a more aggressive form of Crohn’s disease; the rs2797685 variant of the PER3 gene was associated with early onset of disease, higher use of immunosuppressants, and more frequent stricturing and fistulizing disease. Taken together, these observations suggest that circadian disruption is a risk factor for IBD. However, it is also clear from preclinical studies that inflammatory mediators can disrupt circadian function. It is possible that a vicious cycle exists in some IBD patients in whom disruption of circadian clock function increases susceptibility to or enhances intestinal inflammation, and the products of the inflammatory process further disrupt circadian function both centrally and in the periphery, as illustrated in Figure 2.

Sleep deprivation and IBD

Preclinical studies. Forced acute or chronic intermittent sleep deprivation results in worsening of colitis induced in mice by DSS and also delays recovery from the colitis. Sleep deprivation using a water-immersion technique also aggravated DSS colitis in mice. This was accompanied by downregulation of adiponectin and aquaporin 8 and upregulation of of E2F transcription factor (E2F2) mRNA, histocompatibility class II antigen A, and beta 1 (H2-Ab1) mRNA in the colon. Administration of melatonin reversed these microRNA profiles and attenuated the severity of colitis in sleep-deprived mice. Another study confirmed the ability of melatonin to attenuate the severe colitis induced by sleep deprivation, and this was associated with restoration of the normal expression of adiponectin and the cytokines IL-6 and IL-17.

Clinical observations. Results of surveys have indicated that IBD patients often experience fragmentation of sleep, prolonged sleep latency, and poor overall sleep quality that correlates with low scores using the IBD Quality of Life (QoL) questionnaire. Psychiatric comorbidity is associated with poor quality of sleep in IBD, and patients with depression had a 2-fold higher risk of poor quality sleep compared with nondepressed patients. The question arises as to the extent to which poor quality sleep is a consequence of a primary sleep disorder or whether it is secondary to immune activation.
associated with chronic intestinal inflammation and its attendant morbidities. Graff et al found that poor quality of sleep and subsequent fatigue were highly prevalent in not only IBD patients with active disease but also those in remission. Quantitative and qualitative impairment of sleep is therefore associated with poor quality of life and worse disease outcomes and may be a risk factor for IBD.

Altered feeding behavior in IBD

**Preclinical studies.** It is well established in animal studies that administration of pro-inflammatory cytokines including TNF-α and IL-1β results in reduced food intake and weight loss in animals. Induction of experimental colitis using acetic acid resulted in anorexia and weight loss that was prevented by centrally administered interleukin-1β antagonist. Interleukin-1β also mediated anorexia in a model of TNBS-colitis. The action is mediated, in part, by IL-1β-induced production of 5-HT in the hypothalamus. In another study in TNBS-induced colitis, the severity of colonic inflammation correlated with plasma leptin concentrations. As the colitis resolved, food intake normalized and plasma leptin levels fell below normal values, suggesting that the adipokine contributed to the anorexia.

**Clinical studies.** Reduced appetite is common in IBD, particularly in Crohn’s disease, and is likely multifactorial. It is an important factor in weight loss and growth retardation in young patients. According to a systematized review, there may also be a link between eating disorders, particularly anorexia nervosa and Crohn’s disease, associated with a poor prognosis due in part to noncompliance with therapy. Mechanisms underlying inflammation-associated anorexia are poorly understood, but interest has focused on the anorexigenic properties of pro-inflammatory cytokines and adipokines such as leptin, produced in part by mesenteric white adipose tissue. For example, it is postulated, primarily on the basis of preclinical studies, that pro-inflammatory cytokines release leptin from adipose tissue. Circulating leptin would then signal the hypothalamus to reduce appetite. Although data regarding leptin levels and food intake in IBD have been conflicting, recent attention has focused on the ability of adipokines to modulate the inflammatory process in IBD.

**Clinical observations.** A retrospective analysis of 69 patients with active IBD receiving anti-TNF-α therapy showed a significant improvement in depression-like symptoms as assessed by the Patient Health Questionnaire (PHQ-9). In a prospective cohort study, IBD patients with moderate to severe disease starting biologic therapy with vedolizumab or anti-TNF reported improvement in sleep and elevation of mood by 6 weeks. A recent study evaluated cognitive and affective biases before and after anti-TNF-α treatment in 9 patients with Crohn’s disease. Treatment was associated with an improved sense of well-being and postprandial fullness. Importantly, these subjective improvements were associated with post-treatment changes in the limbic system as assessed by fMRI.

**Question 6: What Is the Impact of Anti-Inflammatory Therapy on Brain and Behavior?**

**Preclinical studies.** In a model of chronic colitis induced by nematode infection, administration of anti-TNF by intraperitoneal injection resulted in an improvement in colitis-associated anxiety-like behavior. In a model of depression induced by chronic restraint stress, peripheral administration of anti-TNF improved depression scores but, interestingly, not the attendant weight loss. Clinical data indicate that preexisting behavioral disturbance increases susceptibility to inflammatory stimuli or exacerbates quiescent inflammation and that effective treatment of altered behavior mitigates this vulnerability. Studies also indicate that antidepressants may have direct anti-inflammatory effects. From clinical studies, it is evident from epidemiological studies that psychiatric comorbidity, particularly depression or anxiety, is more common in IBD patients compared with non-IBD controls, may precede the onset of IBD, and is associated with clinical relapse of IBD. However, there is no good evidence at this time that effective treatment of psychiatric comorbidity attenuates the inflammatory process; virtually all studies rely on symptom-based evaluations of IBD disease activity. Absence of evidence does not, of course, imply absence of an effect, and there is general agreement that well-designed randomized controlled studies of antidepressants are urgently needed and must utilize objective measurements of inflammatory disease activity. If psychotropic drugs are shown to impact disease activity, their use in depressed IBD patients should improve response rates of existing immunomodulatory therapies and further improve quality of life.

In terms of gut-to-brain signaling in IBD, it is evident from preclinical models that mediators generated by intestinal inflammation alter brain function and modify behavior. Clinical studies suggest that effective counterinflammatory therapy directed at the gut can improve mood, a general sense of well-being, and sleep quality. This may, in part, reflect the reduction in intestinal symptomatology. It may also reflect changes in brain function. Investigation of the impact of disrupted circadian rhythm in the brain and the periphery on the natural history of IBD is in its infancy and may be subject to novel therapeutic approaches; recent preclinical studies suggest that activation of melatonin receptors may restore disrupted circadian function in the intestine and improve inflammatory responses.
The findings summarized in this review prompt consideration for a holistic approach to the management and perhaps the prevention of IBD by optimizing bidirectional gut-brain interactions (see Fig. 2). Such an approach may involve the appropriate use of antidepressant therapy, chronotherapy, or other approaches to restored circadian rhythm, in conjunction with pharmacotherapeutic or biologic anti-inflammatory therapies.

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