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Effective colonic motility involves an intricate pattern of excitatory and inhibitory neuromuscular signals that arise from the enteric neural circuitry of the colon. Recent investigations have demonstrated that inflammation leads to a variety of changes in the physiological properties of the neurons in this circuitry, including hyperexcitability of neurons at the afferent end of the peristaltic reflex, synaptic facilitation, and attenuated inhibitory neuromuscular transmission. Furthermore, links have been established between these changes and disrupted motor activity in the colon, and we now know that some of these changes persist long after recovery from inflammation. It is highly likely that inflammation-induced neuroplasticity, which is not detectable by clinical diagnostics, contributes to disrupted motility in active and quiescent inflammatory bowel disease and in functional gastrointestinal disorders.

Introduction

The nervous and immune systems represent two of the final frontiers in our attempts to understand how complex organisms function, and it is becoming clear that these two systems interact extensively in health and disease. For example, neurons have the ability to upregulate or downregulate the extent of inflammatory responses, and pro-inflammatory mediators can act on nearby nerves to mediate short- and long-term changes in neuronal function.

Intestinal functions are regulated by the most complex system of nerves outside of the CNS. During the past decade, significant research efforts have been directed toward unraveling the relationship between the colonic inflammatory response and neuronal functions in the gut. Inflammation-induced changes have been reported in neurons in dorsal root ganglia, sympathetic prevertebral ganglia, and enteric ganglia. This Review concentrates on those components of the enteric nervous system (ENS) that regulate propulsive motility and explores how propulsive motility is altered through inflammation-induced neuromuscular changes in the colon secondary to intestinal inflammatory disorders such as inflammatory bowel disease (IBD) and enterocolitis.

Neuronal circuitry that underlies propulsive motility

Over a century ago, Bayliss and Starling demonstrated that the innervation of the intestines is unique compared with other organ systems (1). They showed that stimulation of the mucosal surface in one location results in contraction proximal and relaxation distal to the site of stimulation. They called this phenomenon the “law of the intestine” and attributed this complicated reflex response to the actions of the “local nervous mechanism.” Based on this knowledge that intrinsic reflex circuits can regulate the functions of the intestine, Langley subsequently divided the autonomic nervous system into three parts — parasympathetic, sympathetic, and enteric, with the enteric division comprising the nerves housed within the wall of the gastrointestinal tract (2).

During the last two decades of the twentieth century, significant effort was directed toward identifying the neuronal elements of the reflex circuits that are responsible for peristaltic propulsion of luminal contents and active secretion of water in the intestines (3–5). Compared with the peristaltic reflex circuit, the circuitry responsible for secretion is relatively simple. Active secretion is housed in the submucosal plexus and involves a two-neuron reflex that includes an afferent neuron, which can be activated by secretory products of enteroendocrine cells (e.g., serotonin secreted by enterochromaffin [EC] cells), and a secretomotor neuron that projects into the lamina propria.

The reflex circuitry that mediates propulsive motility includes ascending and descending limbs that lead to proximal contraction and distal relaxation, as described by Bayliss and Starling (Figure 1 and ref. 1). Like the secretory reflex, the motor reflex is initiated by the activation of an afferent neuron by enteroendocrine secretions and/or by stretch. Interneurons projecting upstream in the intestine selectively form synapses with excitatory motor neurons that cause contraction of the smooth muscle via the release of acetylcholine and tachykinins. Interneurons that project downstream in the gut cause relaxation of the smooth muscle via the release of nitric oxide and purines. Collectively, the upstream contraction and downstream relaxation results in a pressure gradient that propels the luminal contents in an aboral direction.

The neurons that serve at the afferent end of intrinsic reflex circuits in the gut are called AH neurons because one of their characteristic electrical properties is a prolonged afterhyperpolarization (AHP). AH neurons are sensitive to stretch, and they project to the lamina propria, where they can receive the chemical signals released from enteroendocrine cells. As a result of their functions, these neurons have been called intrinsic primary afferent neurons (6); however, some researchers feel that they do not fit the formal definition of primary afferent/sensory neurons and prefer to call...
Inflammation-induced changes in colonic motor circuitry

**AH neurons.** Because AH neurons project to the lamina propria of the intestine, where inflammatory responses are centered, they are particularly susceptible to the effects of pro-inflammatory signaling molecules in the inflamed gut. Indeed, to date, all studies of inflammation-induced changes to enteric neurons have reported alterations in the electrical properties of AH neurons. The locations and forms of inflammation that have been studied include myenteric (8) and submucosal (9) neurons in the colon inflamed by trinitrobenzene sulfonic acid (TNBS), myenteric neurons in the TNBS-inflamed ileum (10), and ileal myenteric neurons in animals with *Trichinella spiralis*-induced enteritis (11, 12). Furthermore, submucosal AH neuron excitability is enhanced when tissue from β-lactoglobulin–sensitized guinea pigs is exposed to the milk protein (13, 14).

The mechanisms responsible for inflammation-induced AH neuron hyperexcitability have not been completely resolved, but they appear to vary depending on the location and type of inflammatory response. A consistent feature of inflammation-induced neuroplasticity in AH neurons is a reduction in the magnitude of the AHP, which is important in limiting the excitability of these neurons. In TNBS colitis, a major contributor to the hyperexcitability in AH neurons is an increase in a hyperpolarization-activated cation conductance (8). In this model, the AH neuron hyperexcitability involves activation of COX-2 because chronic exposure to prostaglandin E2 results in AH neuron hyperexcitability in normal preparations (15) and because inhibition of COX-2, but not COX-1, prevents AH neuron hyperexcitability in animals with TNBS colitis (16).

In *T. spiralis* enteritis, the AH neuron membrane potential is depolarized and the neurons exhibit a lower input resistance (11, 12). Both of these features, along with the decreased AH, indicate that a decrease in the activity of the intermediate conductance Ca^2+-activated K^+ channel (IK_{Ca}) in these cells is involved in the...
increase in excitability in this condition. In these neurons the cAMP/pCREB signaling pathway is facilitated and acute administration of histaminergic receptors, COX enzymes, and leukotriene inhibitors dampens hyperexcitability (11). Taken together, these reports indicate that hyperexcitability of AH neurons is a common feature of inflammatory conditions in the intestines.

**Interneurons and motor neurons.** Neurons with the electrical properties classified as S neurons include neurons that function as interneurons and motor neurons, as well as S neurons that likely have a role as sensory neurons because they are sensitive to mechanical stimulation (3). The changes in electrical properties of myenteric S neurons in TNBS colitis are more subtle than those of AH neurons, and no significant changes are detected when considered as a single population (8). However, when the S neurons are divided into two subpopulations — those with ascending projections and those with descending projections — the S neurons with ascending projections exhibit a slight increase in excitability. While the electrical properties of colonic S neurons are not dramatically changed in response to colitis, the strength of interneuronal synaptic potentials is significantly increased in both the submucosal and myenteric plexuses of the TNBS-inflamed colon (8, 9, 17). In submucosal ganglia, synaptic facilitation appears to involve a pharmacological transformation from synaptic potentials mediated by nicotinic cholinergic transmission to a combination of nicotinic, purinergic, and in some cases serotonergic transmission (9). In the myenteric plexus, colitis-induced synaptic facilitation does not involve altered pharmacology, but rather an increase in the readily releasable pool of neurotransmitters available in the nerve terminals and a sustained activation of protein kinase A, which likely increases Ca²⁺ influx into the nerve terminals (17).

**Neuromuscular transmission.** The status of neuromuscular transmission in the inflamed colon has been investigated in TNBS colitis. In an intracellular recording study of synaptic junction potentials from circular muscle cells, Strong and colleagues demonstrated that the ascending excitatory neuromuscular transmission was unchanged in the inflamed region, whereas the descending inhibitory neuromuscular transmission was dampened (18). Pharmacological isolation of the nitrergic and purinergic contributions to the inhibitory junction potential (IJP) demonstrated that nitrergic neuromuscular transmission is normal in colitis, whereas the purinergic component was significantly attenuated. Since nitrergic and purinergic neuromuscular signals likely arise from the same nerve terminals in the intestines, the finding that purinergic IJPs are disrupted but nitrergic IJPs are normal was a bit perplexing. A recent investigation has shed light on this dilemma by demonstrating that the disruption of the purinergic IJP likely involves an interruption of purine production in the mitochondria due to oxidative stress in the inflamed region (19). IJPs in non-inflamed preparations are attenuated when the tissue is exposed to free radical donors or ATP synthase inhibitors, and neurally mediated purine release is decreased in inflamed preparations. In both guinea pig TNBS colitis and murine DSS colitis, treatment of animals with a free radical scavenger protects purinergic IJP. Addition of tempol to the drinking water during the development of colitis prevents oxidative stress in the tissue and maintains the purinergic IJPs at a normal amplitude even though the inflammatory scores of the colons are comparable to those not treated with the free radical scavenger. This disruption in the descending, inhibitor limb of the peristaltic reflex could affect propulsive motility by decreasing the extent of relaxation that occurs in the receiving segment of the gut during peristalsis.

To summarize, findings described above indicate that a number of changes are occurring in the intrinsic motor circuitry of the colon in response to inflammation (Figure 1). These include the following: (a) an increase in serotonin availability due to increased numbers of EC cells and decreased serotonin-selective reuptake transporter (SERT) levels; (b) an increase in the excitability of AH neurons that can result from a decrease in IKca function and/or an increase in hyperpolarization-activated cation current (Ih); (c) a facilitation of interneuronal synaptic activity due to an increase in the readily releasable pool of neurotransmitters in myenteric nerve terminals; (d) a decrease in purinergic neuromuscular transmission due to an oxidative stress–induced decrease in purine release from the terminals of inhibitory motor neurons. In addition, colitis is associated with a loss of about 20% of the myenteric neurons (20, 21).

**Linking inflammation-induced neuroplasticity to disrupted motility**

The questions of whether and how these inflammation-induced changes in the neural circuitry affect motility has been studied using an ex vivo guinea pig colon propulsive motility assay that provides the investigator with an opportunity to run numerous trials and evaluate the effects of test compounds that interact with receptors and ion channels. Initial studies simply involved measuring the amount of time it took for a fecal pellet to progress along a segment of normal versus TNBS-inflamed distal colon and demonstrated that the velocity of propulsive motility was slower in the inflamed preparations (22). Subsequent studies that used a Gastrointestinal Motility Monitor (23) to evaluate the progress of the fecal pellet along the segment of colon demonstrated that propulsive motility was completely obstructed or temporarily halted in heavily inflamed regions (18, 24).

**Attention deficit disorder in the ENS.** Given that major neurophysiological changes induced by colitis are associated with increased neuronal activity and enhanced synaptic transmission, one might intuit that these changes would be associated with enhanced reflex activity and accelerated motility. However, another possibility is that the increased neuronal activity in inflamed regions results in a form of attention deficit disorder in the ENS. Under normal, healthy conditions, the background activity in the ENS motor circuitry is thought to be rather quiescent due to the relative inexcitability of AH neurons. Thus, in response to a luminal stimulus or stretch at one site, there is an activation of AH neurons at that level of the gut, leading to unambiguous signals passing proximally and distally to contract and relax the bowel, respectively. As the bolus is propelled in the distal direction, this response repeats itself, resulting in a propagating wave motility pattern (Figure 2A).

In inflamed regions of the colon, about 50% of the AH neurons exhibit spontaneous action potentials. This likely leads to the activation of ascending contractile and descending inhibitory signals arising from multiple regions along the inflamed segment of bowel. This aberrant neuronal behavior may then result in mixed...
excitatory and inhibitory messages converging on smooth muscle throughout the inflamed region — a form of intestinal fibrillation.

To test whether enhanced neuronal activity in the myenteric plexus would slow rather than accelerate propulsive motility, segments of guinea pig distal colon were treated with veratridine, which at a concentration of 1 μM causes enteric neurons to generate action potentials repetitively (25). Even though there was abundant neuronal and muscle activity in the tissue, it was not possible to generate an adequate pressure gradient to move the fecal pellet forward.

**Links between neuroplasticity and disrupted motility in colitis.** Strong support for the concept that neuroplastic changes in colitis that are associated with increased neuronal activity contribute to disrupted propulsive motility comes from experiments in which similar neuroplastic changes were induced pharmacologically in normal colons or were reversed in TNBS-inflamed preparations (24). In normal colons, the velocity of propulsive motility is significantly decreased when IKCa channels are inhibited, resulting in an increase in AH neuron excitability. The velocity is further reduced when interneuronal synaptic activity is enhanced with a 5-hydroxytryptamine type 4 (5-HT4) receptor agonist (24, 26). Thus, by mimicking the AH neuronal hyperexcitability and synaptic facilitation features of colitis, motility is inhibited rather than accelerated.

Propulsive motility is restored in inflamed preparations when major features of neuroplasticity are reversed or prevented. As described above, in TNBS colitis, AH neuron hyperexcitability involves activation of COX-2 and an increase in the Ih current. The rate of colonic motility is significantly higher in TNBS-inflamed colons from animals that are treated daily with a COX-2 inhibitor (16). Consistent with this, propulsive motility is dramatically improved in TNBS-inflamed colons following application of hyperpolarization-activated cyclic nucleotide–gated (HCN) channel blockers to inhibit the Ih in AH neurons (24). In these experiments, obstructed or halted motility patterns were converted to relatively linear motility patterns, with the fecal pellets traversing the inflamed region without pausing. Motility is also improved in TNBS-inflamed animals that had been treated with a free radical scavenger to protect the purinergic IJP from the effects of oxidative stress, and when HCN channel blockers are applied to these colons, motility is further improved to a healthy control level.

Taken together, these findings underscore the delicate and complicated nature of the peristaltic reflex circuitry, and they demonstrate that overstimulation of the system can actually lead to disruptions in propulsive motility. These studies also demonstrate that even though mucosal erosion, muscularis hypertrophy, and inflammation are still present, propulsive motility can be protected or reestablished when neuronal activity is restored to its normal level.

**Persistent postinflammatory changes in enteric neurons and motility**

Understanding the changes in neuronal function during active inflammatory responses is important in the context of linking these changes with altered gut functions as described above. It is also important to consider which aspects of inflammation-induced neuroplasticity persist following recovery from the inflammatory response. This is a relevant question because alterations in gut function are often sustained during remission of IBD (27–31), and there is mounting evidence to suggest that irritable bowel syndrome (IBS) and other functional gastrointestinal disorders involve an inflammatory component (27, 32–35). In fact, in a large contingent (~35%) of diarrhea-predominant IBS patients, a careful history reveals that the symptoms began during a bout
of infectious enteritis (36), and these individuals are described as having post-infectious IBS. It is important to note that the types of inflammation-induced neuronal changes that have been identified (i.e., changes in neuronal excitability, ion channel function, and synaptic strength) are invisible in any clinical test. Thus, inflammation-induced neuroplasticity that outlasts the inflammatory response offers a mechanism that could be responsible for hard-to-explain, persistent symptoms experienced during remission from inflammation in IBD or associated with IBS.

*T. spiralis.* Myenteric neurons have been studied with electrophysiological techniques at “post-inflammatory” time points in *T. spiralis* and TNBS-inflamed animals, and while AH neuronal hyperexcitability was detected in both of these models, the mechanisms appear to differ. Christofi and colleagues examined the excitability of jejunal AH neurons at a time point of 35 days post-infection, by which time the parasite would have been cleared from the gut (11). Strictly speaking, this model is probably best described as a post-infectious model rather than a post-inflammatory model because evidence of inflammation is still detectable in the wall of the gut, including elevated myeloperoxidase, mast cell tryptase, and prostaglandin levels. Regardless, AH neuron hyperexcitability in this model likely involves post-translational changes in ion channel function because it is reversed by acute exposure to blockers of adenylyl cyclase, histamine, cyclooxygenase, or leukotriene pathways. However, long-term transcriptional changes may also contribute to persistent neuroplasticity in these cells because elevations in adenylyl cyclase and CREB phosphorylation levels are still detectable 35 days after infection.

**TNBS colitis.** When considering the TNBS colitis model, it is important to recall that six days after administration, during the active phase of the inflammatory response, AH neurons are rendered hyperexcitable through a COX-2-dependent mechanism, interneuronal synaptic activity is augmented, and purinergic transmission is blunted. In this model, macroscopic damage scores return to normal 14 days after TNBS administration, and all signs of inflammation, including elevated myeloperoxidase and prostaglandin levels, have disappeared by 28 days (37). At 56 days (four weeks) after recovery from inflammation, AH neurons are still hyperexcitable and synaptic facilitation is still augmented. As detailed above, AH neuron hyperexcitability contributes to dysmotility in TNBS colitis, but both excitability and motility are relatively normal in animals that have been treated with a COX-2 inhibitor for four days prior to euthanasia. To determine whether the post-inflammatory persistent changes in the colon involved a sustained low-level activation of COX-2 to maintain the neuroplasticity and dysmotility, TNBS-inflamed animals were treated with a COX-2 inhibitor on days 52 to 56. In these preparations, the colons still exhibited significant dysmotility, which suggest that the changes are somehow imprinted into the AH neurons during the inflammatory response when it can be prevented or reversed by COX-2 inhibitors. It remains to be determined what transcriptional or epigenetic factors are responsible for the changes in AH neurons, and what might be done to reverse them. Regardless of the mechanism, however, changes such as these may contribute to post-inflammatory and post-infectious changes in gut function and sensitivity.

While AH neuron hyperexcitability and synaptic facilitation persist in the post-inflamed colon, purinergic transmission recovers with the resolution of inflammation (18). This is not too surprising because its disruption is due to an effect of oxidative stress on mitochondria and the oxidative stress subsides with the recovery from inflammation.

**Allergen sensitivity in the ENS.** While it is not necessarily a post-inflammatory condition, food sensitivity represents another circumstance in which memory in the system can lead to pathological responses involving immune cells and the ENS. To investigate this, guinea pigs have been sensitized to cow’s milk at weaning, and then responses to β-lactoglobulin are evaluated (13). When challenged with the allergen, mast cell degranulation leads to AH neuron depolarization and hyperexcitability and a suppression of sympathetic synaptic activity in enteric ganglia. In this paradigm, the memory is stored in mast cells, which in turn activate acute changes in local neuronal function. Together with increased secretion, these changes result in enhanced propulsive motility and a flushing of the allergens from the lumen.

**Colonic motility in the inflamed human colon.** It is difficult to make direct comparisons between the experimental results of animal studies and the pathophysiological results of human procedures. Such comparisons are difficult due to differences in the chronicity of animal models versus human diseases, the lack of animal models that perfectly reflect the human disorders, the diversity in human populations versus the homogeneity of animal strains, the ability to tightly control experimental conditions in animal studies, and the wider variety of ex vivo and invasive in vivo approaches that can be used in animal studies. In the case of IBD, very few investigations of motor activity have been performed and most were conducted decades ago. That said, several reports indicate that the intuitive presumption that diarrhea, a common feature of IBD, is associated with increased motility in inflamed colons is not supported by clinical investigations of colonic motility. In fact, diarrhea in IBD is more closely linked to decreased contractile activity (38–43).

Rao and colleagues detected decreases in whole gut and small intestinal transit rates and found that subjects with active colitis exhibited proximal colon stasis (38). More distally, they reported that rectosigmoidal transit was increased and that stool frequencies were elevated in active colitis relative to quiescent disease. They concluded that diarrhea in colitis is due to rectosigmoid irritability and exudation of mucus and blood in the sigmoid and rectum, rather than rapid transit, and that caution should be used when treating active colitis with anti-diarrheal agents that could further inhibit proximal colon motility. Consistent with this, in an earlier study, Kern and colleagues reported that the most striking abnormality of the motility pattern in ulcerative colitis was a reduction or complete absence of phasic activity (39). They reported that the reduced phasic activity was closely correlated with severe diarrhea and a more normal motility pattern appeared when the diarrhea terminated. Interestingly, they noted, “although it might have been expected that the absence of phasic activity would be due to rigidity and fibrosis of the colon in advanced disease, this phenomenon is clearly unrelated to the degree of anatomical change” (39). The gastrocolic reflex response is also disrupted in ulcerative colitis. Snape et al., using simultaneous myoelectric and pressure recordings, reported that while colonic spike activity increased rapidly following a meal, the maximal response was smaller and of shorter
duration than that observed in controls, and that no postprandial increase in contractility was observed in the inflamed patients (43). They concluded that the lack of contractility might contribute to the increase in diarrhea observed in these individuals after a meal.

While contractility appears to be dampened in IBD, the frequency of propagating contractions is increased. In ulcerative colitis, propagating contractions are always anterograde, resulting in rapid movement of contents to the sigmoid colon (41, 44). The lack of retrograde contractions could be another contributing factor in IBD-associated diarrhea.

Concluding remarks

Effective movement of luminal contents requires a coordinated orchestration of excitatory and inhibitory signals from enteric neurons to the muscularis in a given region of the intestine. Propulsive motility involves a series of overlapping reflex circuits along the intestine that are capable of first receiving and then propelling the luminal contents along. It is possible that alterations in enteric neural function contribute to the disrupted contractility that has been reported in the human colons of inflamed subjects. The data obtained from animal models indicate that inflammation-induced enteric neuroplasticity can, and likely does, contribute to disrupted motility in colitis. These findings indicate that the orchestrated pattern of neuronal activity in the peristaltic reflex circuitry is disrupted in colitis, largely through an increase in neuronal activity and interneuronal synaptic strength, as well as a decrease in inhibitory purinergic neuromuscular transmission. Furthermore, many of these changes persist long after the recovery of colitis. These changes may contribute to alterations in gut function that are maintained during remission from IBD, and to altered motility in IBS; however, these functional properties of gut neurons are currently undetectable in a clinical setting.

It is important to keep in mind that in addition to the neurophysiological changes that alter gut motility and that have been the focus of this Review, there are a number of other concurrent changes within and outside of the colon that also play a role. These include changes within the colon such as the loss of neurons (20, 21), muscle changes such as hypertrophy, altered ionic conductances and responsiveness to neurotransmitters (45, 46), changes in mucosal serotonin signaling that appear to result in increased serotonin availability in inflamed regions (47), and changes in the physical compliance of the bowel in inflamed regions due to fibrosis. Changes in the sensitivity of extrinsic afferent neurons and/or changes in sympathetic input could also contribute to altered colon function as well (33). In addition to alterations in sympathetic activity due to inflammation-induced changes in the electrical properties of prevertebral neurons, there is a loss of synaptic input to the prevertebral ganglia from the gut due to a loss of intestino-fugal neurons in colitis (48).

To more fully understand the role of neuroplasticity in IBD, it will be necessary to undertake the difficult task of studying colonic neurons from human subjects. In an intracellular study of human myenteric neurons, Brooks and colleagues determined that neurons with electrical properties similar to those in guinea pig were observed and, as in guinea pig, about 4% of the colonic myenteric neurons exhibited the features of AH neurons (49). A more recent investigation has described the use of combined intracellular recording, dye filling, and immunohistochemistry in human myenteric neurons (50), demonstrating that it may be possible to ultimately determine whether the functional properties of myenteric neurons are altered by inflammation. Another approach that may be useful in investigating the pathophysiology of human enteric neurons involves the use of voltage-sensitive dyes (51). However, an imaging approach is less useful than electrical recordings for understanding ionic conductance, and this electrophysiological information has been critical for understanding, mimicking, and reversing inflammation-induced changes in animal models. As addressed in this Review, determining the role of neuroplasticity in human colonic function may prove to be crucial in providing more accurate information in the pathogenesis of IBD and ultimately lead to improved therapeutic techniques.

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