Over the past few decades, corticotropin-releasing factor (CRF) signaling pathways have been shown to be the main coordinators of the endocrine, behavioral, and immune responses to stress. Emerging evidence also links the activation of CRF receptors type 1 and type 2 with stress-related alterations of gut motor function. Here, we review the role of CRF receptors in both the brain and the gut as part of key mechanisms through which various stressors impact the gastrointestinal system. We also examine how these mechanisms translate into the development of new approaches for irritable bowel syndrome, a multifactorial disorder for which stress has been implicated in the pathophysiology.

**Stress and gut motor function**

A real or perceived threat to the homeostasis of mammalian organisms, originating internally or externally, triggers adaptive stress responses that affect behavior, as well as endocrine, immune, autonomic, and visceral functions (1, 2). Regarding effects in visceral functions in particular, early seminal investigations reported by Cannon and by Almy and colleagues provided clear evidence that emotional stress impacts gastric and colonic motor activity in healthy volunteers and cats (3, 4). Subsequent studies established that delayed emptying of the stomach is the most common response evoked by various acute stressors in both experimental animals and healthy human subjects (5). By contrast, in the colon, various stressors (including anxiety, dichotomous listening, fear, intermittent hand immersion in cold water, and stressful interviews) increase colonic motility in healthy subjects (5). Similarly, in rodents and dogs, stressors as diverse as open-field test, fear conditioning, loud noises, restraint, cold exposure, water avoidance, inescapable foot or tail shocks, and increased levels of cytokines induced by endotoxins stimulate propulsive colonic motor function (2, 5–7). The autonomic nervous system provides a peripheral neuronal network by which the effects of stress can be rapidly imposed on gut function. These are mediated through the sympathetic and vagal and pelvic parasympathetic innervation of the enteric nervous system (20, 21), particularly because of the high prevalence of coexisting psychiatric disorders, prominently anxiety and depression (16–18). Consequently, emphasis has been placed on elucidating the level of involvement of CRF signaling pathways in both the brain and the gut, in the alterations of gut motor function known to be associated with stress (5, 19).

In addition, the clinical relevance of overactive CRF signaling in the brain and periphery in functional bowel diseases is receiving increasing interest (20, 21), particularly because of the high prevalence of coexisting psychiatric disorders, prominently anxiety and depression, in IBS patients (22). In this Review, we first discuss recent advances in our knowledge of CRF signaling as it relates to CRF family members and their receptors, as well as the insight provided by the development of CRF receptor antagonists. Preclinical studies supporting a primary role for CRF receptor activation in the brain and gut in mediating the alterations of gastric and colonic motor function associated with exposure to various stressors are outlined. The potential relevance of CRF signaling pathways in the pathogenesis and treatment of IBS is also addressed (20, 23).

**CRF signaling pathways**

**CRF family members and receptors.** CRF was the first peptide isolated (15) of a family of mammalian CRF-related peptides that now includes urocortin 1, urocortin 2 (also known as stresscopin-related peptide), and urocortin 3 (also known as stresscopin).
In summary, CRF and urocortins have a common carboxy-terminal region and structurally distinct amino-terminal extracellular domains (the region involved in ligand binding) that contain 34 aas for CRFα1, 61 aas for CRFβ2, and 20 aas for CRFγ2 (24, 28). Recently, a novel soluble splice variant has been identified in the mouse brain and shown to encode only the first extracellular domain of CRFα1 and to function as a soluble binding protein for CRF and urocortin 1 (29).

Despite sharing 70% amino sequence similarity, CRF1 and CRF2 display distinct characteristic affinities for the CRF family of peptides (Figure 1) (24, 26). CRF has a higher affinity (10– to 40-fold higher) for CRF1 than for CRF2, whereas all the urocortins signal preferentially through CRF2. Urocortin 1 binds CRF2 with 100-fold greater affinity than does CRF, and CRF1 with 6-fold greater affinity than does CRF (30–32). Urocortin 2 and urocortin 3 exhibit high selectivity only for CRF2 (31, 32), with a slightly higher affinity for CRFβ2 compared with CRFα1, and a low affinity for the new soluble CRFα1 splice variant (29, 32) (Figure 1). CRF2 variants display tissue- and species-specific expression. In nonprimate mammals, CRFβ2 is expressed only by neurons and CRFα1 mainly in the periphery and by non-neuronal cells of the brain (33), whereas CRFγ2 is found only in the amygdala of the human brain (34). It is well documented that stimulation of CRFα1, CRFβ2, and CRFγ2 activates adenylyl cyclase/cAMP signaling pathways through coupling and activation of Gs proteins (26). However, several recent reports indicate that the nature of a trimeric complex, made of CRF agonist, CRF receptor, and G protein, influences the pattern of intracellular signaling in a tissue-specific manner (26).

CRF receptor antagonists. The development of competitive CRF receptor antagonists was an important early development in the endeavor to determine the functions of CRF receptors under basal and stress conditions (35). The first CRF antagonist was α-helical CRF9-41, a CRF analog generated by deletion of 8 aas from the amidated carboxyl terminus of CRF (27, 36). A subsequent approach to enhance the potency of CRF antagonists was to design analogs in which the secondary structure of CRF was constrained, leading to the generation of [α-Phe12,Nle21,38,αMeLeu37]CRF22-41 (α-Phe12CRF12-41) and cyclo[30–33][β-Phe12,Nle21,38,Glu50,Lys35]Ac-CRF9-41 (known as astressin) (37). Additional astressin-like analogs were later developed, of which cyclo[30–33][β-Phe12,Nle21,αMeLeu27,Glu50,Lys33,Nle38,αMeLeu40]Ac-CRF9-41 (known as astressin-B) is the most efficacious and long-acting (being still effective 24 hours after a single peripheral injection) (38). The use of these CRF antagonists has unraveled the many roles that CRF receptors have in orchestrating the behavioral (anxiety, decreased feeding, and drug seeking), cognitive (anxiety and anxiety), neuroendocrine (ACTH and β-endorphin release), autonomic (activation of the sym-
pathetic nervous system), immunological, and visceral (hypertension and alterations in gut motor function) responses to stress.

However, these CRF antagonists bind both CRF- and CRF2 receptors; therefore, do not provide selectivity to assess the involvement of the 2 CRF receptor subtypes (27, 42) (Figure 1). An important goal was reached recently when competitive and selective peptide antagonists for CRF2 were developed and shown to bind equally to the a, b, and c variants of CRF2 while having little to no affinity for CRF1 receptors (29, 43, 44) (Figure 1). Three of these peptides are [d-Phe<sup>11</sup>, His<sup>13</sup>]-sauvagine<sub>11-40</sub> (known as antisauvagine-30), [d-Phe<sup>11</sup>, His<sup>12</sup>, Nle<sup>17</sup>]-sauvagine<sub>11-40</sub> (known as K141998), and the long-acting analog with additional conformational constraints, cyclo[(31–34)[d-Phe<sup>11</sup>, His<sup>13</sup>, Nle<sup>17</sup>, Cz-MeLeu<sup>13,39</sup>, Nle<sup>27</sup>, Glu<sup>19</sup>, Lys<sup>38</sup>]] Ac-sauvagine<sub>28-40</sub> (known as astrassin-B) (Figure 1). So far, there are no peptide analogs that are selective CRF1 antagonists; however, a flurry of patents for orally bioavailable, nonpeptidic selective CRF1 antagonists, it has become clear that the constellation of physiological effects produced by endogenous CRF peptides might be attributed to actions on distinct CRF receptor subtypes. Compelling evidence indicates that activation of the brain CRF-CRF<sub>1</sub> signaling pathway has a leading role in coordinating many of the physiological responses to adaptive stress as it relates to the activation of the hypothalamic-pituitary-adrenal (HPA) axis, sympathetic nervous system, and changes in cardiovascular, colonic, and immune functions in rodents and primates (23, 39, 47–49). Preclinical and clinical studies also indicate that abnormally increased central CRF<sub>1</sub> signaling contributes to the pathogenesis of anxiety and depression and can have implications in the pathophysiology of IBS (18, 20, 49). With regard to CRF<sub>1</sub> receptors in the brain, emerging evidence supports a role for these receptors as mediators of ways to dampen and/or facilitate the proper recovery of the CRF-initiated behavioral, endocrine, and visceral responses to stress (50–52). However, in some systems — for instance, the suppression of feeding behavior — activation of CRF<sub>1</sub> has an additive effect with the CRF<sub>1</sub>-mediated orexigenic effect (49).

**Link between CRF receptors in the brain and stress-related alterations of gut motor function**

Convergent experimental reports have shown that central injection of CRF and urocortins reproduces stress-related alterations of gut motor function in naive rodents, whereas central injection of CRF antagonists prevents the effects of various stressors, supporting a crucial role for CRF receptors in the brain in the regulation of stress-induced alterations in gastrointestinal motility (53).

**Stress, CRF receptors in the brain, and gastric transit.** CRF, urocortin 1, urocortin 2, and the nonmammalian CRF-related peptides sauvagine and urotensin I inhibit gastric emptying of noncaloric liquid, caloric liquid, and solid food when injected into the cerebrospinal fluid (CSF) of several nonprimate mammals (53). These peptides also inhibit basal and cholinergic-stimulated gastric motility in rodents and dogs (54, 55). CRF<sub>2</sub> is the CRF receptor subtype in the brain through which CRF and urocortins injected into the CSF primarily initiate their inhibitory effect on gastric transit and motility (56–58). Sites of CRF action in the brain are specific and localized in the paraventricular nucleus of the hypothalamus (PVN) and the dorsal vagal complex nuclei, both of which contain neurons bearing CRF<sub>2</sub> and are known to influence autonomic nervous outflow to the stomach (Figure 2) (53, 59, 60). Consistent with this, blocking of the transmission of impulses by the autonomic nervous system prevents CRF injected into the CSF or PVN from inhibiting gastric transit (61–62). Removal of the pituitary gland or adrenal glands had no effect on CRF inhibition of gastric emptying; this indicates that the gastric effect is not secondary to the activation of the HPA axis (61, 62). All reports, except 2 (61, 63), have identified the vagus nerve as the main pathway mediating the delayed gastric transit and inhibition of gastric motility induced by central injection of either CRF or urocortin 1 in rats and dogs (54, 55, 57, 59, 62, 64, 65). By contrast, the delayed gastric emptying induced by injection of urocortin 2 into the CSF is not altered by gastric vagotomy and instead requires the integrity of the sympathetic nervous system and peripheral α<sub>2</sub>-adrenergic receptors (57). These data indicate that CRF and its related peptides differentially modulate vagal and sympathetic components of the autonomic nervous system to suppress gastric motor function.

The importance of CRF signaling in the brain was established using pharmacological blockade of CRF receptors. Injection of α<sub>2</sub>-helical CRF<sub>6–41</sub>, d-Phe<sup>12</sup>CRF<sub>12–41</sub>, astressin, or astressin-B into the CSF or PVN completely prevented the delayed gastric emptying induced by various psychological, physical, visceral, immunological, or chemical stressors, including swim stress, restraint, abdominal or cranial surgery, peritoneal irritation with 0.6% acetic acid, systemic or brain injection of IL-1β, and exposure to ether anesthesia (5, 63, 66). Consistent with this, various stressors, including abdominal surgery and peripheral administration of IL-1β, activate neurons that express CRF and upregulate levels of mRNA encoding CRF in the PVN (67–70). Likewise, urocortin 1, urocortin 2, and urocortin 3 are present in the PVN, and expression of the urocortins is upregulated during stress (49, 71, 72). The CRF receptor subtypes and endogenous CRF ligands primarily involved in delaying gastric motor function under stress conditions have been characterized in only a few studies so far. The CRF<sub>2</sub> antagonist astressin<sub>2</sub>-B injected into the CSF abolished the delayed gastric emptying in response to restraint in rats (63). By contrast, pharmacological blockade with a selective CRF<sub>1</sub> antagonist and the use of mice lacking CRF<sub>1</sub> suppressed the inhibition of gastric transit induced by surgical stress (caused by abdominal surgery or cecal palpation) (73). Collectively, these data provide new insights into the role of specific CRF receptors in the brain in the altered gastric digestive function that occurs in response to stress of surgical or immunological origin. As inhibition of gastric propulsive activity after surgical intervention represents a substantial medical problem for which effective treatments are still lacking (74), these experimental findings may open new therapeutic avenues.

**Stress, CRF receptors in the brain, and small-intestinal transit.** Alterations in small-intestinal motor function induced by stress and CRF ligands have been investigated in parallel with those occurring in the stomach (53). Acute psychological stress and central injection of CRF and urocortin 1 exert an inhibitory effect on duodenal and small-intestinal transit and propulsive motility, similar to their effects on gastric functions (6, 54, 65, 66, 75–77). Peptide-inhibitory action is mediated directly by vagal nerves and is independent of activation of the HPA axis (6, 61, 66, 78).
The inhibitory action of central CRF on small-intestinal transit is, however, less prominent than its inhibitory action on gastric transit; this most probably reflects the lesser density of vagal innervation of the small intestine as compared with the stomach (79). In addition, the role of CRF receptors in the brain in stress-related inhibition of small-intestinal transit is yet to be characterized. The reduction of small-intestinal transit induced by partial restraint was reported to be blocked in male, but not in female, rats injected with α-helical CRF9–41 into the CSF (6, 66). Whether these conflicting data relate to sex differences or other experimental components remains to be determined.

Stress, CRF, in the brain, and colonic transit. In contrast to the inhibitory effects of CRF and urocortin 1 injected into the CSF on gastric and small-intestinal motor function, these peptides stimulate colonic transit and defecation and induce diarrhea through increased sacral parasympathetic outflow to the large intestine in female and male rats, mice, and gerbils (6, 7, 56, 61, 80–85). Convergent studies have shown that the activation of CRF1 receptors in the brain contributes to the stimulatory effects of central injection of CRF and urocortin 1, as well as the stimulatory effects of various stressors, on colonic motor function (23, 53). First, the colonic propulsive motor activity induced by stress is mimicked by central administration of urocortin 1 and the preferential CRF1 agonist ovine CRF (27), whereas the selective CRF2 agonists urocortin 2 and urocortin 3, injected into the CSF at a dose similar to that of CRF, are inactive (56). In addition, central administration of the pan–CRF peptide antagonists α-helical CRF9–41, D-Phe12–CRF12–41, and astressin blocked the colonic motor stimulation (motility, transit, and defecation) induced by central injection of CRF and urocortin 1, as well as by various stressors (wrap or partial restraint, water avoidance, conditioned fear, IL-1β injected into the CSF, and morphine withdrawal) (6, 7, 56, 66, 80, 82, 83, 86–89). Similarly, the selective CRF1 antagonists CP-154,526, CRA 1000, NBI 27914, NBI 35965, and antalarmin, injected either into the CSF or i.p., prevented the acceleration of colonic transit induced by restraint, dampened defecation in response to water avoidance, restraint, and social stress, and inhibited the diarrhea elicited by morphine withdrawal (23). Likewise, CRF1-deficient mice have lower defecation scores than wild-type littermates in an open-field test (90). Lastly, the CRF2-deficient mice have lower defecation scores than wild-type littermates in an open-field test (90). Lastly, the CRF2-deficient mice have lower defecation scores than wild-type littermates in an open-field test (90). Lastly, the CRF2-deficient mice have lower defecation scores than wild-type littermates in an open-field test (90).

The PVN, the locus coeruleus (LC), and the Barrington’s nucleus, which lies just ventromedial to the LC, are areas of the brain where CRF and stress stimulate colonic motor function and anxious behavior (16, 91) (Figure 2). These sites are activated by water-
avoidance stress, as is shown by their increased expression of FOS, a neuronal marker of cell activation (80). Water avoidance also induces rapid transcription of the gene encoding CRF in the PVN (92). Furthermore, α-helical CRF9-41 injected into the PVN prevents the stimulation of colonic transit and defecation induced by partial restraint and water avoidance (64, 80, 87). Likewise, in inbred rats with a genetically impaired hypothalamic CRF response to stress (93), water avoidance results in a reduced activation of neurons in the PVN and sacral parasympathetic nucleus (as shown by decreased expression of FOS) and is associated with an attenuated colonic motor response (86). Moreover, expression of the gene encoding CRF; in the PVN (60) is markedly increased by different types of interoceptive or exteroceptive stressors (94). It has been shown that CRF-synthesizing neurons in the Barrington’s nucleus project to both the noradrenergic LC and the sacral parasympathetic nucleus of the spinal cord, which innervates the descending colon (Figure 2) (91, 95). In the LC, CRF increases the rate at which noradrenergic neurons fire and thereby increases the amount of noradrenaline released into the brain cortex, leading to arousal and anxiogenic behavior (Figure 2) (91, 96). Consequently, the activation of CRF-CRF signaling in the PVN and LC (94, 97) is well positioned to participate in stress circuits that coordinate behavioral anxiogenic and autonomic responses that impact colonic motility (81, 91). It might be speculated that overactivity of these neurocircuits has relevance to the high incidence of anxiety disorders in patients with IBS and that these effects might be efficiently targeted by CRF; antagonists (16, 23, 98).

Peripheral CRF signaling and stress-related alterations of gut motor function

As established for a number of neuropeptides (such as somatostatin, opiates, and calcitonin gene–related peptides) that act in the brain to influence gut motility (99), the CRF ligands and receptors that were initially characterized in the brain (where they function to influence gut motor function) have recently been shown to be widely expressed in peripheral tissues, including the gastrointestinal tract of experimental animals and humans (19, 32, 100–103). The coincident expression of CRF ligands, mostly urocortins, with cognate receptors provided strong support for the idea that their local action could influence gut motor function (100–102, 104, 105).

**CRF receptors in the periphery and gut transit.** Initial functional studies showed that injection of CRF peripherally alters gut motility and transit in several mammalian species, including rodents, dogs, and humans (6, 62, 106–109). In particular, injection of CRF either i.v. or i.p. inhibited gastric emptying, delayed small-intestinal transit, stimulated colonic transit and defeation, and induced diarrhea with a potency similar to that of CRF injected into the CSF (6, 61, 106). Although central administration and peripheral administration of CRF result in similar gut transit alterations, distinct sites and mechanisms of action are involved (54, 57, 61, 110, 111). For example, pharmacological blockade of autonomic outflow does not modify the inhibition of gastric emptying and acceleration of colonic transit induced by injection of CRF i.p., whereas it abrogates the gastric and colonic responses induced by injection of CRF into the CSF (61). In addition, injection of α-helical CRF9-41 into the CSF does not modify the stimulation of colonic transit induced by injection of CRF i.v.; this indicates that the peripherally injected CRF did not activate CRF receptors in the brain (66).

Further studies in rodents have established that peripheral injection of CRF, urocortin 1, urocortin 2, or urocortin 3 delays gastric emptying by activating CRF; whereas peripheral injection of CRF or urocortin 1 stimulates colonic motility through activation of CRF; expressed by colonic myenteric neurons (19). This was shown by the fact that injection of the selective CRF; agonists, urocortin 2, or, less potently, urocortin 3 either i.p. or i.v. inhibited gastric emptying of a solid or liquid meal but did not influence distal colonic transit in rodents (106, 112). By contrast, under the same conditions, CRF and urocortin 1 inhibited gastric motor function and stimulated colonic propulsion and defecation in rats and mice (54, 106, 112). Moreover, in rodents, peripheral injection of the CRF;–specific antagonists astressin-B and antiguavine-30 prevented the inhibition of gastric emptying that is induced by CRF and urocortin 1 given i.v. or i.p. but did not modify their stimulation of distal colonic transit (106, 112). Conversely, peripheral injection of the CRF;–specific antagonists CP-154,526 and NBI 27914 blocked the stimulation of colonic transit, defeation, and diarrhea induced by i.p. injection of CRF and urocortin 1 but did not prevent delayed gastric emptying (106, 112–114). The mechanisms by which peripherally administered CRF and urocortin 1 stimulate colonic motor activity might involve direct activation of colonic myenteric neurons that lie between the longitudinal and circular muscles. Indeed, in rats, high levels of FOS expression are induced in neurons of the colonic myenteric ganglia by CRF injected i.p., and this is blocked by peripheral injection of astressin and CP-154,526 (115). These data are consistent with the expression of CRF; by rat colonic myenteric neurons (101, 104).

**CRF receptors in the periphery and gut motility.** Analysis of the motility changes underlying gut transit alterations showed that i.v. injection of urocortin 1 or CRF reduces the amplitude of postprandial gastric contractions, inhibits jejunal motility induced by i.v. injection of motilin, and increases propulsive colonic motility in experimental animals (54, 113, 116, 117). The motor alterations induced by CRF and urocortin 1 were reproduced in vitro in isolated stomach tissue and distal segments of the colon, which both have functional enteric neurons; this supports the idea of local peripheral action (102, 105, 113, 118, 119). Studies in healthy humans revealed that i.v. injection of CRF increases nonpropulsive postprandial duodenal motor activity and stimulates propulsive motor contractions in the descending colon (108, 109). Of interest is the report that patients with IBS show enhanced colonic motility in response to i.v. injection of CRF as compared with healthy volunteers, which indicates that they are hyperresponsive to CRF, and this might be linked by upregulation of colonic CRF; (108).

**Stress, CRF receptors in the periphery, and gut transit.** Pharmacological studies support the notion that gut CRF signaling occurs under stress (6, 19, 112, 113). Several reports showed that the delayed gastric emptying induced by abdominal surgery can be blocked by peripheral injection of α-helical CRF9-41, β-Phε12CRF12-41, and astressin (111, 120, 121). CRF; antagonists injected i.v. also prevented the inhibition of gastric emptying induced by acute wrap-restraint stress, whereas the selective CRF; antagonist CP-154,526 did not (112). With respect to stress-related stimulation of colonic motor function, peripheral administration of α-helical CRF9-41, astressin, and CP-154,526, but not of astressin-B, prevented or blunted the stimulation of distal colonic transit and defeation induced by acute wrap-restraint and water-avoidance stress (6, 83, 112, 113, 122). In patients with IBS, compared with healthy subjects, the administration of α-helical CRF9-41 improves colonic motility, visceral perception, and the negative mood elicited by rectal transmural electrical stimulation, without affecting the HPA axis (20).
The cellulos of the CRF and CRF-related peptides that activate peripheral CRF receptors present in the gut remain to be elucidated. CRF ligands have been detected in the gut myenteric nervous system, as well as in enteroneuroendocrine cells and lamina propria macrophages in rodents and humans (103, 123, 124). Because both central and peripheral administration of CRF receptor antagonists is able to counteract the impact of stress on gut motility, this supports the concept that stress influences the release of CRF ligands in the gut through autonomic pathways, where they can then function as local effectors of altered gastrointestinal motility.

Conclusions and future perspectives

CRF signaling in the brain, established as a leading mediator of the biochemical effect on the endocrine and anxiogenic behavior responses to stress, is also part of the underlying mechanisms through which stress inhibits gastric transit and stimulates colonic transit in experimental animals. In addition to the brain, the gut through which stress inhibits gastric transit and stimulates colonic transit the concept that stress influences the release of CRF ligands in IBS diarrhea-predominant patients, such as stimulation of colonic motility, defecation/watery diarrhea, visceral hypersensitivity, and anxiogenic/hypervigilant behavior, that are alleviated by CRF1 receptor antagonists (23). These data support the involvement of the CRF1 system at central and/or peripheral sites as part of the mechanisms whereby stress triggers or enhances gut complaints in patients with IBS (9–11). The promising completion of the first open-label clinical trial with the CRF1 antagonist R121919 in severely depressed patients (125), along with the improvement of colonic motility and visceral perception by peripheral injection of the CRF antagonist α-helical CRF(9–41) in IBS diarrhea-predominant patients (20), provides a strong basis for therapeutic use of CRF1 antagonists to treat IBS, particularly given the high frequency of comorbid psychiatric disorders in IBS (22).

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