Disruption of Circadian Rhythms and Gut Motility
An Overview of Underlying Mechanisms and Associated Pathologies

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Abstract: Circadian rhythms ensure that physiological processes occur at the most biologically meaningful time. The circadian timing in the gastrointestinal tract involves interlocking transcriptional and translational feedback loops that culminate in the rhythmic expression and activity of a set of clock genes and related hormones. The suprachiasmatic nucleus and peripheral core molecular clocks oscillate every 24 hours and are responsible for the periodic activity of various segments and transit along the gastrointestinal tract. Environmental cues may alter or reset these rhythms to align them with new circumstances. Colonic motility also follows a circadian rhythm with reduced nocturnal activity. Healthy humans have a normal bowel motility during the day, frequently following awakening or following a meal, with minimal activity during the night. Maladjusted circadian rhythms in the bowel have been linked to digestive pathologies, including constipation and irritable bowel syndrome. Our advanced knowledge of the link between the circadian clock and gastrointestinal physiology provides potential therapeutic approaches for the treatment of gastrointestinal diseases. This review seeks to establish evidence for the correlation between circadian rhythm, bowel movements and digestive health, and examine the implications of disrupted circadian rhythms on gut physiology.

Key Words: circadian rhythm, bowel motility, clock genes, gastrointestinal tract

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Many organisms have circadian clocks that anticipate daytime and establish endogenous 24-hour rhythms, which organize their physiology and behavior.1,2 The circadian clock is orchestrated by a central pacemaker in the brain that resets the clocks in peripheral organs to control expression of key genes throughout the day.3 Circadian rhythms, driven by cell-autonomous biological clocks, take cues from cycles of light and dark, hormone levels or from the metabolic status of the individual.3,4 The first recorded reports of circadian control and the gastrointestinal system date back to the second or third century BC, where hunger was used as a time signal.5 Circadian rhythms control several gastrointestinal functions ranging from gastric enzyme and fluid production to small intestine nutrient absorption, and gastric/gut motility. Environmental cues alter or reset these rhythms to align with new circumstances.5,6 The most documented external cue, exposure to light, does not drive the circadian rhythm of the gut but merely resets the 24-hour cycle.5,6

NORMAL GUT PHYSIOLOGY

The gastrointestinal system is governed by a circadian rhythm characterized by quiescence during the night, rapidly elevated activity at the time of awakening and increased activity throughout the day.7 This rhythm, and associated endogenous clocks, prepares the body for anticipated stimuli such as feeding.8,9

DISRUPTED GUT PHYSIOLOGY

Disrupted circadian rhythms in the bowel have been linked to gastrointestinal conditions including constipation and irritable bowel syndrome (IBS).10,11 For example, differences in colonic pressure activity12,13 and time of bowel movements13,14 are observed in constipated patients versus healthy persons depending on the time of day. Similarly, differences in the frequency of bowel contractions15 and the sensitivity of rectal mechanoreceptors16 have been reported in irritable bowel syndrome-constipation (IBS-C) patients relative to healthy controls with respect to morning versus evening hours. These observations and their implications for gastrointestinal health are discussed in more detail later. Figure 1 depicts normal and disrupted circadian control of the gastrointestinal tract.

The objectives of this review are to establish evidence for the correlation between circadian rhythm, bowel movements, and digestive health, and implications of disrupted circadian rhythms on gut physiology.

METHODOLOGY

A systematic literature review was conducted by searching the Medline database (PubMed; November 1, 2017) to collate relevant published data on circadian rhythms and the gastrointestinal system. Studies included trials (randomized controlled trials, quasi-experimental and preexperimental), observational studies or reviews. Only articles written in English and published in peer-reviewed international journals were selected.

In phase 1, a search string was constructed to identify relevant articles:

("gastrointestinal [All Fields] OR “gut” [All Fields] OR (“intestines” [MeSH Terms] OR “intestines” [All Fields]

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In phase 2, search results were manually filtered for relevance based on their title and the inclusion/exclusion criteria included in the abstract. This was carried out by 2 independent reviewers and discrepancies were resolved by a third reviewer. Relevance was defined as coverage of the circadian rhythms or underlying mechanisms influencing the digestive system and its health.

In phase 3, data extracted from the long list were reviewed by 2 independent reviewers and discrepancies were resolved by a third reviewer. The results are shown in Figure 2.

**PHYSIOLOGY OF CIRCADIAN RHYTHMS IN THE GASTROINTESTINAL TRACT**

The molecular machinery synchronizing circadian rhythms in the gut is believed to be controlled by 2
complementary systems: a central mechanism in the brain and organ-specific peripheral mechanisms.\(^5\)

**Central and Peripheral Mechanisms**

In humans, the suprachiasmatic nucleus (SCN) in the ventral hypothalamus acts as the central circadian pacemaker which resets itself using light signals transmitted via the retinohypothalamic tract.\(^5\) The SCN communicates with peripheral tissues via neural and humoral pathways, most prominently via corticosteroids and melatonin.\(^6\) The SCN also controls the circadian release of digestive peptides, including vasoactive intestinal polypeptide and gastrin-releasing peptide.\(^16\)

Peripheral circadian clocks generally operate to a phase that is 4 hours behind the central clock. Local cues, such as food intake, are used to reset or entrain the circadian rhythm of the gastrointestinal system, independently of signaling from the SCN.\(^5,6,13\) Even when the SCN is rendered inactive due to a lesion, circadian rhythms remain in the gastrointestinal system, suggesting the system is primed to anticipate alternating levels of demand between day and night.\(^5,6\)

**Clock Genes**

Clock genes are a group of genes that are controlled by molecular feedback mechanisms.\(^13\) They are expressed rhythmically in epithelial cells of the colon and in neurons in the myenteric plexus which have a significant role in coordinating colonic motility.\(^3,5,13,17\) Clock genes are self-regulating, facilitating their 24-hour rhythm, and modulate the activity of other genes at specific times during each day.\(^13\)

Hoogerwerf and colleagues investigated measures of colonic motility in wild-type mice, per1per2 double-knockout mice, and neuronal nitric oxide synthase knockout mice. Assessments consisted of a combination of in vivo and ex vivo methods.\(^6\) Persistence of rhythmicity in stool output, intracolonic pressure changes, and tissue contractility in wild-type mice under constant darkness and the absence of this rhythmicity in per1per2 double-knockout and neuronal knockout mice confirmed that these measures are circadian and controlled by endogenous clock-driven processes.\(^6\) Examples of clock genes include the helix-loop-helix transcription factors clock and bmal1, per1, per2, and per3 genes, and the cryptochromes cry1 and cry2.\(^13\) Approximately 8% to 10% of genes in peripheral organs are ultimately controlled by clock genes.\(^13\) In humans, single nucleotide polymorphisms in clock genes have been associated with changes in gastrointestinal motility.\(^18\) The CLOCK3111T/C single nucleotide polymorphism is associated with significantly lower dominant frequency on electrogastrography (EGG) recordings compared with subjects with wild-type (TT) clock genes, indicating slower gastric motility.\(^18\) The PER3 variable-number tandem-repeat polymorphism (either 4 or 5 repeats, 54 nucleotides in length), which is also associated with circadian rhythm, has been shown to exacerbate CLOCK3111T/C-related slowing of gastric motility.\(^19\)

**Entrainment**

Light is an essential external cue for entrainment of the central circadian clock located in the SCN.\(^20,21\) In addition, there are peripheral clocks in nearly all tissues and cells of the body that are directly entrained by food, independent of the SCN.\(^20,21\) Although the mechanisms involved in entrainment have not been definitively identified, potential candidates include hormonal signaling (eg, melatonin), neural signaling, food intake, and body temperature regulation.\(^20,21\)

### Influence of Light on Gastrointestinal Circadian Rhythms

When subjects are exposed to dim-light versus bright light conditions during the day, less carbohydrate from the evening meal is absorbed in the cecum, although the circadian phases in each group appear to be consistent.\(^22\) Furthermore, baseline running spectral total power (ie, total spectral power of each frequency region on an EGG for 30 min) is comparable between subjects exposed to dim or bright light conditions. Taken together, evidence suggests that exposure to dim-light conditions during the daytime suppresses the digestion of an evening meal, resulting in malabsorption of dietary components.\(^23\)

### Influence of Food on Gastrointestinal Circadian Rhythms

Peripheral clocks are influenced by timing cues related to food intake. Restricting feeding synchronizes clock gene expression in the gastrointestinal system, independently of central control via the SCN.\(^6\) Furthermore, feeding may send metabolic cues to the SCN, entraining the central circadian rhythm.\(^6\) Studies in rodents show that rhythmic presentation of food to rats with SCN lesions can entrain anticipatory wheel running suggesting the existence of a food-mediated entrainment oscillator, semi-independent of the SCN.\(^23\) Food in contact with the gastrointestinal epithelium has been ruled out as an entrainment mechanism, although humoral mechanisms or food-related entrainment mediated by a corticotrophin-releasing hormone is a possibility.\(^5,6\)

### Influence of Exercise on Gastrointestinal Circadian Rhythms

Graded aerobic exercise has been shown to decrease phasic colonic motility in healthy, untrained subjects\(^24\) but after exercise there was a resurgence of predominantly propagating pressure waves, suggesting that exercise may enhance stool transit.\(^24\) Similarly, acute physical exercise increases both high amplitude propagated contractions (HAPCs) and low amplitude propagated contractions (LAPCs) in healthy individuals.\(^25\) It has been demonstrated that voluntary exercise can shift the SCN clock or modulate the photic synchronization of circadian rhythms.\(^26\) A simulated night work-study indicated that moderate-intensity exercise during the night shift produced larger circadian phase shifts compared with a sedentary control condition.\(^27\) Other groups indicated that low-intensity and high-intensity exercise during the night influenced circadian rhythms by the following day.\(^28,29\) As discussed earlier, these changes in circadian parameters can have profound effects on gastrointestinal motility.

### Circadian Rhythms and Control in the Gastrointestinal System

Migrating motor complex (MMC) are waves of electrical activity that start in the stomach and move along the gut triggering peristaltic contractions.\(^7\) Hormones such as motilin and ghrelin are involved in the generation of MMCs, while others (eg, gastrin, cholecystokinin, serotonin) are involved in the generation of propagation sequence spikes.\(^7\) These processes result in peristaltic or segmental contractions in the small (duodenum, jejunum, ileum) and large intestines (colon).\(^1\) In the small bowel, the length of the interdigestive cycle is divided into 4 phases. The first phase of the MMC involves a prolonged period of quiescence (40% to 60% of total time); phase II is characterized by
increased frequency of action potentials and smooth muscle contractility (20% to 30% of total time); phase III consists of a few minutes of peak electrical and mechanical activity (5 to 10 min); phase IV includes declining activity which merges with the next phase. The cycle is consistent throughout a 24-hour period, but the relative proportion of each cycle attributed to each phase differs between day and night. A significantly higher number of cycles at night have phases of motor quiescence compared with daytime, while the duration of the phase II period of increased frequency of action potentials and smooth muscle contractility is longer during the day versus night ensuring bowel movements occur during waking hours. In the colon, there is no MMC but sleep is known to strongly inhibit both propagating and nonpropagating colonic motor activity.

**Gastric Activity**

Normal gastrointestinal motility occurs through coordinated contractions of smooth muscle which derives from 2 patterns of electrical activity across membranes of smooth muscle cells: slow waves and spike potentials. The gastric activity can be recorded by EGG, a noninvasive technique. The dominant EGG frequency reflects the frequency of stomach contractions. The dominant frequency observed in EGG recordings is believed to reflect gastric slow waves; rhythmic smooth muscle cell depolarisations that act as the gastric pacemaker to coordinate contraction. Mean dominant frequency is increased during daytime hours and decreases during sleep. Esophageal function and circadian rhythms are important due to their association with gastroesophageal reflux disease (GERD). Typically, GERD patients complain about frequent night reflux symptoms. Nighttime GERD can profoundly impair the quality of life by causing pain and disturbance of sleep, that interferes with mental and physical functioning the next day. The upper esophageal sphincter tone changes little during sleep compared with waking hours. However, the motility of the esophagus is reduced during sleep, with the frequency of both primary and secondary contractions progressively diminishing from stages N1 to N3. N1 stage sleep refers to the transition stage between wakefulness and the deeper stages of sleep, while the N3 stage refers to deep or slow-wave sleep. In contrast, secondary contractions increase during rapid eye movement (REM) sleep. The lower esophageal sphincter has transient reductions in tone during sleep, which are conducive to reflux episodes.

**Small Intestine Activity**

Motility follows a circadian rhythm with reduced nocturnal activity. Using twin intraluminal pressure-sensitive radiotelemetric capsules, Kumar et al demonstrated significant variations between daytime and nocturnal propagation velocities of the MMC in the small intestine. Nocturnal MMCs are less frequent and have slower velocities of the MMC in the small intestine.

**Colonic Activity**

Normal colonic motility follows a rhythm, with minimal activity during the night and increased activity during the day. Typical activity during the day consists of increased colonic propagation sequences compared to those occurring at night in healthy persons. For example, Narducci et al showed that HAPCs were more prevalent in the morning in a study that measured 24-hour manometric recording of colonic motor activity in healthy subjects. Furthermore, Bassotti et al showed that LAPC waves were constantly present with an average of about 61 events/subject/day and mean amplitude of about 20 mm Hg. More than 80% of LAPCs waves occurred during the day and were increased after meals and morning awakening ($P<0.05$). At night, Bampton et al demonstrated that there was nocturnal suppression of colonic motor activity. Comparing area under the curve data in a 2-hour period after awakening in the morning to a 2-hour period before waking showed almost a 2-fold increase in colonic activity (4874 to 8335 mm Hg; $P < 0.0001$). The increase in colonic activity remained elevated throughout the waking hours.

Retrograde pressure waves are less frequent than antegrade propagating pressure waves and demonstrate a lower frequency at night followed by a slightly increased frequency after awakening. Under normal physiological conditions, propagating sequences were seen to migrate in both antegrade and retrograde directions in healthy persons. Antegrade propagating sequences were more frequent and of greater amplitude, as measured by a prolonged multipoint recording of colonic manometry in the unprepared human colon in subjects with normal bowel habits in daytime hours. Likewise, Furukawa et al found retrograde propagation sequences were of significantly lower amplitude. Dinning et al showed that postprandial distal colonic activity consists primarily of a retrograde propagating sequences following a meal in healthy persons. These conditions may prevent premature rectal filling, while also allowing stools to remain for the appropriate amount of time for absorption of water and electrolytes. In patients with constipation, a meal failed to induce the normal increase in the distal colonic propagating sequences seen in healthy persons.

**Rectal Activity**

The rectum exhibits intermittent cyclical motor activity specifically at night which has been termed the rectal motor complex or periodic rectal motor activity (PRMA). In healthy individuals, PRMA is confined largely, but not exclusively, to the rectosigmoid region and follows a distinct circadian rhythm. Rao et al demonstrated that PRMA often follows a motor event in the more proximal colon and that there is a distinct circadian rhythm suggesting that PRMA may serve as an intrinsic braking mechanism that prevents untimely flow of colonic contents. The number, duration and peak amplitude of rectal motor complexes (regular pressure fluctuations with a frequency of either 3 or 6 cycles/min) are significantly reduced at night.

A summary of the associations between circadian rhythms and gut motility is shown in Table 1.

**HORMONAL CONTROL OF CIRCADIAN RHYTHM IN THE GASTROINTESTINAL SYSTEM**

**Melatonin**

Melatonin is the neuroendocrine “clock factor” generated by the pineal gland, a structure that dominates...
TABLE 1. Summary of the Associations Between Circadian Rhythm and Gut Motility

| Region          | Esophageal motility follows a natural circadian rhythm                                                                                     | Colonic motility follows a circadian rhythm                                                                 |
|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------|
| Esophagus       | Overall esophageal motility is reduced during sleep.                                                                                         | There are slight variations in colonic motor activity during different phases of the sleep cycle.           |
|                 | There are slight variations in esophageal motility during different phases of the sleep-cycle.                                                | Upper esophageal sphincter tone does not change during sleep vs. waking hours.                             |
|                 | Upper esophageal sphincter tone does not change during sleep vs. waking hours.                                                               | There are transient reductions in lower esophageal sphincter tone during sleep vs. waking hours.          |
| Small bowel     | Normal small bowel motility follows a circadian rhythm with reduced nocturnal activity.                                                      | Normal colonic motility follows a circadian rhythm, with minimal activity during the night and increased activity during the day. |
|                 | There are variations in MMCs between daytime vs. nocturnal propagation velocities.                                                          | HAPCs increase after meals and during the day.                                                           |
|                 | Nocturnal MMCs are less frequent and have slower velocities.                                                                                   | LAPCs increase after meals and during the day.                                                           |
|                 | There are conflicting reports on MMC duration, ie, shorter vs. no difference.                                                                | Antegrade propagating sequences are more frequent and of greater amplitude during daytime hours.            |
| Colon           | The number, duration and peak amplitude of rectal motor complexes are significantly reduced at night.                                         | Retrograde propagation sequences are less frequent and have lower amplitude during daytime hours.         |
| Rectum          | Normal rectal motility follows a distinct circadian rhythm consisting of intermittent cyclical motor activity specifically at night.            | Postprandial distal colonic activity consists primarily of retrograde propagating sequences following a meal in healthy persons. |
|                 | PRMA is confined largely, but not exclusively, to the rectosigmoid region.                                                                    | HAPC indicates high amplitude propagated contractions; LAPC, low amplitude propagated contractions; MMC, migrating motor complex; periodic rectal motor activity; REM, rapid-eye movement. |

HAPC indicates high amplitude propagated contractions; LAPC, low amplitude propagated contractions; MMC, migrating motor complex; periodic rectal motor activity; REM, rapid-eye movement.

Gastrointestinal Pathology and Circadian Rhythms

Circadian clock desynchronization often leads to functional abnormalities in the gastrointestinal system (abdominal pain, constipation, and diarrhea), and metabolic diseases (obesity, and nonalcoholic fatty liver disease), and to increased susceptibility to alcoholic liver disease due to increased intestinal permeability.5,6,13,17,20,51

Circadian disruption displaces the timing of eating and normal gastrointestinal functions, such as gastric, bile, and pancreatic secretions, enzyme activity, intestinal motility, and the rate of nutrient absorption.52,53 Surveys have documented gastrointestinal problems in shift-workers in which nighttime workers complained of constipation.54,55 Altered gastrointestinal physiology and/or gut microbiota may drive gastric pathologies by increasing gastrointestinal permeability, which in turn increases the risk of injury and inflammation.39 This is supported by reports that melatonin may have a role in maintaining the integrity of the gut wall.56

Constipation

As discussed earlier, several lines of evidence suggest that altered circadian rhythm contributes to the manifestation of constipation. For example, patients with constipation demonstrate a significantly lower colonic pressure activity following waking or food consumption compared to nonconstipated subjects.5,15 This is in contrast with the normal diurnal rhythm observed in nonconstipated patients where colonic pressure is higher in the morning.12,13 Likewise, bowel movements occur most frequently in the evening rather than in the morning in patients with irregular bowel habits of just 3 to 4 times per week. Mass movements (MMs) in healthy persons have been shown to be more frequent during the day than at night, and that the percentage of subjects who expressed MM between 6 AM and 2 PM was greater than those who did so between 4 PM and 4 AM.57 Although the number of MM did follow a diurnal rhythm in constipated patients, it was blunted and the percentage of patients with MM was less in the morning in constipated patients compared to healthy controls.7 Prolonged manometry studies demonstrate that adults with slow transit constipation have fewer spontaneous HAPCs than healthy subjects.12,58–60 This observation will be discussed in more detail later in relation to the potential mechanism of action of the laxative medication.

There is also some evidence to suggest that the circadian clock plays a role in “traveler’s constipation.” This concept was investigated in a study that included 70 people traveling from Europe to the United States for a short stay.61 In addition to the usual questionnaires, 65 subjects kept diaries on their bowel habits, had stool samples evaluated for consistency according to a standardized methodology, and had their colonic transit time (CTT) measured after ingesting radioactive tracers.61 Nearly 40% of the subjects complained of constipation and this was most pronounced during the first days of travel. The degree of constipation correlated with the degree of jetlag.61 A potential caveat of this study could be that factors other than travel (changes in diet and physical activity) may have played a role and further studies are needed to replicate these data.

Other evidence suggests that the incidence of constipation increases during periods of fasting during the daylight. During Ramadan, many adults Muslims fast during daylight hours eliciting a change in circadian rhythm. A questionnaire measured the rate of constipation in 900 individuals and found that those who fasted for >14 days reported more “severe” or “very severe” constipation than those who fasted for <14 days.62 However, alimentation is a confounding factor, affected by the increased volume of food intake and, in some cases, the introduction of traditional dishes.

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IBS

IBS is a prevalent functional gastrointestinal syndrome defined as a constellation of symptoms including abdominal pain, alternating constipation, and diarrhea.53-65 Sleep disturbances are a common comorbidity in IBS, affecting 26% to 55% of patients.48,49 Likewise, the severity of IBS may be exacerbated by poor sleep, leading to the hypothesis that IBS may be associated with a disrupted circadian rhythm.5,17,49 In a study of 399 nurses, IBS was more prevalent in rotating shift workers than in day-shift nurses. This association was found to be independent of sleep quality suggesting circadian disruption.63 The frequency of bowel contractions observed upon awakening in healthy subjects is significantly blunted in the descending colon of patients with constipation-predominant IBS (IBS-C).15 In addition, decreased sensitivity of rectal mechanoreceptors in patients with IBS-C relative to healthy controls has been reported in the morning, followed by significantly higher sensitivity during the day.14

Neurodegenerative Diseases

Circadian desynchronization (disrupted daily rhythms of physiological parameters such as sleep, activity, and hormone secretion) has been regarded as a symptom of several neurodegenerative diseases.66 Disruption in circadian rhythms has a negative impact on quality of life, cognitive performance, mental health, motor control, and metabolism. Many of these functions become impaired in neurodegenerative disorders such as Alzheimer disease, Parkinson disease, and Huntington disease, where chrono-degeneration is a common feature.67 These disorders are associated with constipation. For example, the pathophysiology of constipation in PD includes central mechanisms encompassing changes in dorsal vagal nucleus function and peripheral mechanisms with loss of dopaminergic neurons.68 It is not known whether disruptions of circadian rhythm are causal or symptomatic of neurodegenerative disorders.

A summary of the association between compromised circadian rhythms and gastrointestinal pathologies is shown in Table 2.

THERAPEUTIC BENEFIT OF MODULATING THE CIRCADIAN RHYTHM IN THE GASTROINTESTINAL SYSTEM

A concept has emerged that pharmacological control of the circadian clock may provide a novel therapeutic strategy for gastrointestinal disorders. Chronopharmacological techniques (chronotherapy) aim to optimize the efficiency of a drug via timed administration following a circadian cycle.69 The timing of treatment in coordination with the body-clock may increase the desired effect of drugs, lower the dose and decrease toxicity.69

Melatonin

Melatonin has been considered as a potential co-adjuvant treatment for some gastrointestinal diseases (e.g., IBS-C).49 Studies have shown that melatonin has regulatory effects on gastrointestinal tract motility and sensation that may improve bowel habits and alleviate abdominal pain or distension in IBS patients.49 For example, low-dose melatonin treatment accelerates intestinal transit time whereas high doses decrease gut motility.70,71 The finding that low dose melatonin accelerates intestinal transit time suggests that it could be beneficial in the treatment of IBS-C. For example, this action could also be of benefit for the treatment of constipation alone, although studies investigating this potential have yet to be performed.

Song et al72 investigated the effects of melatonin on abdominal pain in IBS patients who had concurrent sleep disturbances. Melatonin administered at a dose of 3 mg for 2 weeks attenuated abdominal pain and reduced rectal pain sensitivity. Therapeutic efficacy was associated with heightened pressure and volume thresholds for both urgency and pain sensations in melatonin-treated patients.72 Melatonin did not influence sleep parameters, including total sleep time, sleep latency, sleep efficiency, sleep onset latency, arousals, duration of stages 1 to 4, REM sleep, and REM onset latency.72 The authors concluded that the beneficial effects of melatonin in IBS may be related to its action on gut visceral hypersensitivity and independent of its action on sleep.72

A study by Lu et al73 showed that nightly treatment with melatonin (3 mg) for 8 weeks improved mean IBS scores compared with placebo in female patients with IBS. Response rates, defined as the percentage of subjects achieving mild-to-excellent improvement in IBS symptoms, were higher in melatonin-treated patients. Symptoms of sleep disturbance and anxiety/depression scores were comparable between groups.73

TABLE 2. Summary of the Association Between Compromised Circadian Rhythms and Gastrointestinal Pathologies

| Pathology | Description |
|-----------|-------------|
| Disruption of circadian rhythms and gastrointestinal pathologies | Circadian clock desynchronization often leads to functional abnormalities in the gastrointestinal system |
| Circadian disruption displaces timing of normal gastrointestinal functions, including intestinal motility |
| Constipation | Altered circadian rhythms contribute to manifestations of constipation |
| Constipation is more prevalent in night-shift workers |
| Patents with constipation demonstrate a significantly lower colonic pressure activity following waking or food consumption compared to the normal diurnal rhythm observed in nonconstipated subjects |
| Adults with slow transit constipation have fewer spontaneous HAPCs than healthy subjects |
| Mass movements occur less in the morning in constipated patients compared to healthy controls |
| IBS | Sleep disturbances are a common comorbidity in patients with IBS |
| Symptoms of IBS are exacerbated by poor sleeping patterns |
| Patients with IBS-C are more prevalent in night-shift workers |
| Frequency of bowel contractions observed upon awakening in healthy subjects is significantly blunted in the descending colon of patients with IBS-C |
| Patients with IBS-C have decreased sensitivity of rectal mechanoreceptors in the morning, followed by significantly higher sensitivity during the day relative to healthy controls |
| Neurodegenerative diseases | Neurodegenerative diseases including AD, PD, and HD are associated with constipation |
| Circadian dysynchronization has been regarded as a symptom of neurodegenerative diseases |
| It is not known whether disruptions of circadian rhythm are causal or symptomatic |

AD indicates Alzheimer disease; HAPC, high amplitude propagated contractions; HD, Huntington disease; PD, Parkinson disease; IBS, irritable bowel syndrome; IBS-C, irritable bowel syndrome-constipation.
Melatonin levels have also been shown to influence CTT in control subjects and patients with IBS. Participants were treated daily with melatonin (3 mg) or placebo for 8 weeks, followed by a 4-week washout, and then placebo or melatonin in the reverse order for a second 8-week period. Melatonin treatment induced an increase in CTT in control subjects. In contrast to baseline CTT in control subjects, CTT in constipated IBS patients was prolonged. The CTT did not change in IBS patients after melatonin treatment. This study highlights the complexity of mechanisms involved in the effects of melatonin and whether the beneficial effects are more related to rectal physiology. As discussed earlier, there is a fine balance between increasing or reducing CTT via melatonin treatment, and this is largely governed by dose and the specific methodology employed in each trial.

Even though placebo-controlled studies investigating melatonin suffer from considerable heterogeneity in methodology, an extensive literature review found that melatonin improved abdominal pain in many studies, with some studies showing improvements in quality of life.

**Probiotics**

Research has demonstrated that the bacteria in the gastrointestinal tract vary over the course of a day, with the relative abundances of bacterial taxa, the proximity of bacteria to the colonic epithelium, and microbial metabolism all exhibiting diurnal rhythms. Furthermore, the gut microbiome appears to have a reciprocal relationship with the circadian clock and eating habits in humans.

It has been proposed that disturbances in diurnal bowel function in chronic constipation may lead to changes in colonic flora. These alterations in the intestinal microflora could alter the metabolic homeostasis of the colon with resultant changes in the concentration of physiological substances that may influence the motor and secretory functions of the bowel, a theory that was first suggested over 30 years ago. This proposal has been confirmed by evidence showing the ability of probiotics to stimulate the motility of the large bowel and to normalize the intestinal microflora in patients with constipation. Probiotic supplementation was also shown to alleviate constipation-related symptoms: patients defecated with more frequency and ease, and normalized stool consistency in constipated patients. It is plausible that products released by the abnormal flora may contribute to the colonic motility changes that lead to constipation. The observation that the normalization of the intestinal microflora, associated with probiotic therapy, is accompanied by a stimulating effect on colonic motility further supports this concept.

**Oral Laxatives**

As therapeutic agents, modern laxatives are safe to use, and severe reactions are rarely reported. Current laxatives promote defecation by increasing bulk, decreasing stool consistency (softening) or stimulating colon motility through several mechanisms. Laxatives, in addition to interventions like dietary fiber, fluid, etc. are key treatments for constipation.

As discussed throughout this review, a potential player in the manifestation of constipation is a disruption of circadian rhythms in the gut. A strategy conducive to the reinstatement of these rhythms in constipated patients could be to encourage the natural gastrointestinal functionality that occurs in healthy subjects (eg, peristalsis, optimal osmotic pressure). Colonic HAPCs are essential for facilitating the transit of colonic contents over long distances, leading to defecation. Prolonged manometry studies demonstrate that adults with constipation have fewer spontaneous HAPCs than healthy subjects. Studies have shown that intraluminal infusion of bisacodyl, a stimulant laxative that acts directly on colonic musculature, can elicit HAPCs in humans. This effect of bisacodyl was first established over 50 years ago. It has been demonstrated that bisacodyl increased the number of propagated HAPCs as measured by colonic manometry in more recent studies in constipated patients. Bisacodyl-induced HAPCs are quantitatively and qualitatively similar to naturally accumulating HAPCs in healthy persons. Further evidence for a favorable action of bisacodyl linked to circadian control of gut functionality is found in a small study by Kamy et al where the onset of action of bisacodyl was demonstrated to be 12 hours after administration. When taken at night, this property mimics the naturally occurring circadian rhythm promoting peristalsis and the secretion of fluid in the gut leading to the urge to pass stools in the morning, as is the case in healthy subjects.

Collectively, these characteristics are not shared by some other commonly used laxatives include bulking agents, osmotic agents, stool softeners and prokinetics. Although these treatments are effective, there is little evidence to suggest that they influence the reduction of naturally occurring HAPCs seen in constipated patients, and they have variable offsets of action that do not necessarily follow the body’s natural rhythm.

As pointed out earlier, probiotics have been shown to stimulate motility of the large bowel, relieve constipation-related symptoms (eg, reduced frequency of bowel evacuation and compromised stool consistency) and concurrently normalize changes in the intestinal microflora seen in patients with constipation. Probiotics had returned to pretreatment levels by 3 months. These changes were most pronounced among those who were most severely constipated and demonstrated the slowest transit through the large bowel; concentrations of E. coli and Candida were increased by 10- to 100-fold in more than half of the patients in this subgroup. Normalization of evacuation function by bisacodyl treatment was accompanied by a relative normalization of the microflora. Three months after the end of treatment, the values for most microorganisms had returned to pretreatment “diseased state” levels. Bisacodyl did not influence potentially toxic bacteria (eg, Clostridium, E. coli or Enterobacteria). Although this study was performed in a small number of patients, it does provide a plausible link between the gut microbiome and constipation, particularly as the microbiota returned to an imbalanced state following cessation of treatment.

A summary of the therapeutic benefit of modulating the circadian rhythm in the gastrointestinal system is shown in Table 3.

**CONCLUSIONS**

There is convincing evidence that the gastrointestinal system is governed by a circadian rhythm with both central
TABLE 3. Summary of the Therapeutic Benefit of Modulating Circadian Rhythm in the Gastrointestinal System

| Treatment |
|----------------------------------|
| Melatonin | Melatonin has regulatory effects on gastrointestinal tract motility. Melatonin may improve bowel habits and alleviate abdominal pain or distension in IBS patients. Several studies have demonstrated potential benefits of melatonin in patients with IBS and shown improvements in intestinal transit time and abdominal and rectal pain. |
| Probiotics | Bacteria in the gastrointestinal tract vary over the course of a day. The relative abundances of bacterial taxa, proximity of bacteria to the colonic epithelium, and microbial metabolism all exhibiting diurnal rhythms. It has been proposed that disturbances in diurnal bowel function in chronic constipation leads to changes in colonic flora. Treatment with probiotics stimulates motility of the colon and normalizes changes in the intestinal microflora seen in patients with constipation. Probiotic supplementation alleviates constipation-related symptoms: patients defecated with more frequency and ease, and normalized stool consistency in constipated patients.
| Oral laxatives | Use of oral laxatives that act in harmony with circadian rhythms could encourage the natural circadian gastrointestinal functionality that occurs in healthy subjects (eg, peristalsis, optimal osmotic pressure). When taken at night, this onset of action mimics the naturally occurring circadian rhythm promoting peristalsis and the secretion of fluid in the gut leading to the urge to pass stools in the morning, as is the case in healthy subjects.

HAPC indicates high amplitude propagated contractions; IBS, irritable bowel syndrome.

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