Gastroparesis (Gp) is a chronic disease that presents with clinical symptoms of early satiety, bloating, nausea, vomiting, and abdominal pain. Along with these symptoms, an objective finding of delayed gastric emptying, along with a documented absence of gastric outlet obstruction, are required for diagnosis. This article focuses on updates in the pathogenesis and management of Gp. Recent studies on full thickness biopsies of Gp patients have shed light on the complex interactions of the central, autonomic, and enteric nervous systems, which all play key roles in maintaining normal gut motility. The management of Gp has evolved beyond prokinetics and antiemetics with the use of gastric electrical stimulators (GES). In addition, this review aims to introduce the concept of gastroparesis-like syndrome (GLS). GLS helps groups of patients who have the cardinal symptoms of Gp but have a normal or rapid emptying test. Recent tests have shown that patients with Gp and GLS have similar pathophysiology, benefit greatly from GES placement, and likely should be treated in a similar manner. (Gut Liver 2017;11:579-589)

Key Words: Nausea; Vomiting; Gastroparesis; Enteric nervous system; Gastric emptying

INTRODUCTION

Gastroparesis (Gp) is a clinical syndrome consisting of the principal symptoms of nausea, vomiting, bloating, early satiety, postprandial fullness, and upper abdominal pain. In addition, objective evidence of delayed gastric emptying (GE) and exclusion of mechanical obstruction is required. A solid-phase gastric scintigraphy is considered the gold standard in objective evidence required for diagnosis. The symptoms of nausea and vomiting are two of the prevalent symptoms in Gp. Abdominal pain is becoming recognized as a widespread symptom as well, with one study reporting abdominal pain in 90% of their 68 patients. Research on epidemiology was conducted in Olmsted County, Minnesota and reported the incidence, prevalence, and outcomes of Gp patients from 1996 to 2000. The incidence per 100,000 was 2.5 in men and 9.8 in women. The prevalence per 100,000 was 9.6 in men and 37.8 in women. Furthermore, patients with Gp had a significantly higher mortality than the rest of the population in that county. The conclusion of that study was while Gp is uncommon, patients have a recognized increase in health burdens.

ETIOLOGY

The majority of Gp cases can be classified into three main groups: idiopathic gastroparesis (IG, 36%), diabetic gastroparesis (DG, 29%), and postsurgical gastroparesis (PSG, 13%) as seen in Table 1.1,5 Patients with IG, the most common etiology, are those who meet clinical and objective evidence of Gp without a primary identified cause. One important subset of IG is postsurgical; these patients have rapid onset of Gp symptoms after a viral prodromal phase. The classic history is a healthy patient...
with acute Gp symptoms such as nausea and vomiting without resolution of their symptoms. Typically these patients will improve over the course of a year. In comparison, DG is the most common identified cause of Gp. In a study examining 527 type I and type II diabetic patients, 5.2% of type I diabetic and 1.0% in type 2 diabetic patients developed Gp over the course of a 10-year period. PSG is becoming a well-recognized complication of bariatric surgery for bypass and fundoplication for acid reflux. One study followed 615 patients for a year after Nissen fundoplication and found all patients had symptoms the first 3 months postoperatively. Early satiety (88%), and bloating (64%) were the most common symptoms, however 94% of patients had resolution of their symptoms a year after their operation. The other rare causes of Gp include diseases such as Parkinsonism, amyloidosis, paraneoplastic disease, scleroderma and mesenteric ischemia. The clinical guidelines of the American Journal of Gastroenterology recommend for patients with Gp to be screened for diabetes, thyroid dysfunction, neurological disease, bariatric surgery, and autoimmune disorders as possible underlying causes for Gp.

**DIAGNOSIS**

The diagnosis of Gp requires an objective demonstration of delayed GE. The three methods are scintigraphy, wireless motility capsule (WMC), and carbon breath testing. Scintigraphy of a solid phase meal is still the most widely used and validated test for diagnosis. A consensus protocol was published in 2008 to standardize the test. It is recommended that diabetic patients should have their fasting glucose checked before the test and ideally should be <275 mg/dL. In addition, prokinetic agents (Metoclopramide, Erythromycin, and Domperidone), opiate medications, and anticholinergic agents (Bentyl, Levsin, and Robinul) should be stopped at least 2 days prior to the test to prevent interference. Patients should be fasting the previous night and have the test early morning and refrain from smoking during the test. A solid meal was emphasized as liquid studies correlated poorly with Gp symptoms. The recommended meal consists of a technetium (TC)-99m sulfur colloid radiolabeled meal with two large eggs, two slices of bread and jam, with water. An allergy to eggs is a contraindication. The entire meal or at least 50% should be consumed. Imaging is then performed at 0, 1, 2, and 4 hours after ingestion. Delayed GE was then defined at >90% at 1 hour, >60% at 2 hours, >10% at 4 hours. The test is most specific at the 4-hour mark, suggesting the need for studies to be continued for that length of time rather than relying on the 2-hour scan. In addition, patients with rapid emptying should be defined as <30% retention after 1 hour. The GE study should also include the fasting blood sugar of diabetic patients, how much of the meal was ingested, the time, vomiting if it occurred, and medications taken in the last 24 hours to see if they affected the results. All of these recommendations were made to help provide a consensus protocol to help with physician and patient interpretation of results and increase reliability of the test.

One proposed method to increase standardization is the use of a WMC also called SmartPill which is able to measure pH, temperature, and pressure. The SmartPill measures GE time based on the duration of time from ingestion to an abrupt rise in pH from the gastric baseline (>3 pH units) which signifies the change from acidic antrum to the more alkaline duodenum. One study examined 148 patients (87 healthy patients, 61 Gp patients) by having them ingest both the pill cam and doing a GE study. They found that at 4 hours the overall correlation between the two was 0.73.

Another noninvasive alternative is the use of $^{13}$C labeled breath test using octanoate, which is a medium chain triglyceride which is mixed into a solid meal such as eggs or a muffin. The carbon labeled material is ingested and rapidly absorbed in the small intestine, transported to the liver by serum albumin, and oxidized to carbon dioxide which through respiration is eliminated. The rate limiting step of this process is GE. Analysis is done before ingestion and then every 15 minutes until 4 hours; delayed GE is if it takes more than 120 minutes for half of the radiolabeled material to be metabolized and excreted. Samples are measured using mass or infrared spectrometry. The current recommendation is that WMC and breath testing requires further study before it is considered as an equal alternative to a GE study, but its benefits of an in office procedure and lack of radiation help warrant further investigation.

**GASTROPARESIS-LIKE SYNDROME**

An emerging concept proven by increasing evidence demonstrated by pathophysiology is gastroparesis-like syndrome (GLS). New groundbreaking research is revealing that GLS should be classified as a spectrum of Gp. Table 2 shows a quick reference comparing aspects of GLS and Gp. GLS helps to classify

| Table 2. Comparison of Gastroparesis-Like Syndrome and Gastroparesis |
|----------------|----------------|----------------|----------------|
| GLS            | No difference | Normal, rapid  | Decreased      |
| Gp             | No difference | Delayed        | Decreased further |
| GCSI, Gastric Cardinal Scoring Index; GE, gastric emptying; ICC, interstitial cells of Cajal; GES, gastric electrical stimulator; GLS, gastroparesis-like syndrome; Gp, gastroparesis. |
patients who exhibit the symptoms of Gp but have a normal or rapid emptying study. One study examined 425 patients with chronic unexplained nausea and vomiting (CUNV) for at least 12 weeks without evidence of obstruction. Of those patients, 25% percent had GE study that were normal or rapid (less than 30% retention after an hour). The validated Gastroparesis Cardinal Symptom Index (GCSI) scale of severity (0–5) was used for both groups and at 48 weeks follow-up there was no difference in symptoms between Gp and CUNV patients. See Table 3 for the GCSI index.\textsuperscript{15} This begs the further question of how to classify patients exhibiting all the symptoms of Gp without an objective delayed GE study. These patients have also been shown to have a clinical course comparable to that of Gp patients and are proven to have a very similar burden of health. Research has shown that loss of neuronal nitric oxide synthase (nNOS) and loss of interstitial cells of Cajal (ICC) are the two most likely mechanisms that explain Gp.\textsuperscript{14,16,17} Further explanations of the physiology of these cells are described below. New studies involving gastric biopsies and ICC show that patients with CUNV have less ICC compared with healthy individuals. Gp have even fewer ICC. It is because of this correlative that CUNV should be characterized instead in part of a spectrum of Gp and subsequently termed GLS.\textsuperscript{18}

**PATHOGENESIS**

The pathophysiology behind Gp has proven to be complicated, with multiple variables involved. Fundus and antral con-

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**Table 3. Gastroparesis Symptom Cardinal Index**

| Symptom                                      | None | Very mild | Mild  | Moderate | Severe | Very severe |
|----------------------------------------------|------|-----------|-------|----------|--------|-------------|
| 1. Nausea                                    | 0    | 1         | 2     | 3        | 4      | 5           |
| 2. Retching                                  | 0    | 1         | 2     | 3        | 4      | 5           |
| 3. Vomiting                                  | 0    | 1         | 2     | 3        | 4      | 5           |
| 4. Stomach fullness                         | 0    | 1         | 2     | 3        | 4      | 5           |
| 5. Not able to finish a normal-sized meal    | 0    | 1         | 2     | 3        | 4      | 5           |
| 6. Feeling excessively full after meals      | 0    | 1         | 2     | 3        | 4      | 5           |
| 7. Loss of appetite                          | 0    | 1         | 2     | 3        | 4      | 5           |
| 8. Bloating                                  | 0    | 1         | 2     | 3        | 4      | 5           |
| 9. Stomach or belly visibly larger           | 0    | 1         | 2     | 3        | 4      | 5           |

For each symptom, please circle the number that best describes how severe the symptom has been during the past 2 weeks. If you have not experienced this symptom, circle 0.

Adapted from Revicki DA, et al. Aliment Pharmacol Ther 2003;18:141-150, with permission from John Wiley & Sons.\textsuperscript{15}

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**Fig. 1.** Electron microscopy of the interstitial cells of Cajal (ICC) network in gastric smooth muscle in a chronic unexplained nausea and vomiting (CUNV) patient. (A) Compared with the control (B). Labeled structural components: thick basal lamina (small asterisks), nerve endings (NEs), smooth muscle cells (SMCs), lamellar body (LB), contact between ICC and SMCs (arrows), peg-and-socket junction (large asterisks). Adapted from Angeli TR, et al. Gastroenterology 2015;149:56-66.e5, with permission from Elsevier.\textsuperscript{18}
Contractions timed with relaxation of the pylorus in the stomach are coordinated between smooth muscle, autonomic nerves, and ICC. Fig. 1 demonstrates the histological changes that occur with loss of the connection between SMC and ICC.16 ICCs are the most recognized cells that function to regulate smooth muscle contractility through slow wave progression. They can be thought of as the cells that operate as the pacemakers in the gut; the slow waves initiated by ICC result in depolarization of L-type calcium channels resulting in contractile response. These cells are not just chronotropic but ionotropic as well, regulating the strength of the response. It is because of this relationship that research associating motility disorders with ICC are at the forefront of research. ICC cells are being studied across a multitude of motility pathology such as Hirschsprung’s disease, achalasia, and pyloric stenosis. It is becoming increasingly apparent just how important ICC are for the regulation of intestinal motility.19

Examination for histological changes in Gp patients specifically examining ICC have been increasingly studied. Full thickness gastric biopsies obtained from the Gp Clinical Research Consortium, found that ICC cells were decreased. This pattern was seen in both IG and DG. This study consisted of 20 patients in each group of IG, DG, and a control group. In DG they found overwhelming results pointing to ICC correlating with Gp compared to the control group. Patients with DG had only 2.8±0.4 ICC per high power field versus 5.3±0.2, with a p<0.0001. In addition, lower ICC correlated with slower emptying scans. There was a similar correlation in IG patients as well.20

The ICC cells interact with enteric nerves that produce inhibitory and excitatory signals to help regulate smooth muscle and contractility. The study of neuronal changes from these signals has been of great interest as well. Stimuli that promote contraction, such as acetylcholine and substance P, are important for motility. Inhibition of these contractions have been studied as well, specifically nNOS which produces nitric oxide. Fig. 2 shows the interaction between ICC and nNOS for normal GE. Studies done on mice with knockouts for Nos1, which codes for nNOS, demonstrated that the mice developed gastric stasis. Pharmacologic inhibition of nNOS was consistent for delayed for nNOS, demonstrated that the mice developed gastric stasis. Pharmacologic inhibition of nNOS, demonstrated that the mice developed gastric stasis. Pharmacologic inhibition of nNOS was consistent for delayed GE in mice as well. The results of that study showed that the mechanism behind nitric oxide signaling is the inhibition of duodenal and pyloric contractions.21–23 Furthermore, in diabetic and streptozotocin-induced diabetic rats, nNOS expression was decreased along with delayed GE.24,25

Further examining DG, there are new studies looking at the role of systemic inflammation. One study examined 143 patients with DM and symptoms of Gp and found that C-reactive protein, an inflammatory marker, was associated with increased symptoms.26 Another study strongly supporting the role of inflammation in DG examined full thickness gastric biopsies in 28 patients. Six of 14 patients with DG had evidence of a lymphocytic infiltrate in the myenteric plexus as opposed to controls who had no lymphocytic infiltrate. Once again a decrease in ICC was seen as well.27

One pilot study examined the idea of immunotherapy to target the inflammatory role of Gp. This study examined 11 patients with refractory Gp that was resistant to drug and stimulants, and treated the patients with either intravenous immunoglobulin (IVIG), mycophenolate mofetil (MMF), or a combination of methylprednisolone and MMF for 8 to 12 weeks. Symptoms were scored and the study found that patients responded well to IVIG with the most improvement in vomiting symptoms.28 Another similar study was done with 23 patients suspected of having an autoimmune component to their gastrointestinal (GI) dysmotility. Seventeen out of 23 patients in this study improved and involved IVIG as the main therapy along with a few patients receiving methylprednisolone or combination.29 These two studies illustrate the importance of considering an autoimmune etiology and subsequent inflammatory response, especially in patients who have been refractory to medications and stimulants.

Disruptions of the autonomic nervous system (ANS) have also been well-recognized, with previously mentioned gastric biopsies showing a decrease in ICC and enteric neuronal stimulation. One study proposed the effect of the ANS on gastric stimulation by both a cholinergic and adrenergic response with evidence utilized from heart rate monitoring and electrogastrogram in five patients with DG.30,31 Studies are even looking at correlating gastric electrical stimulation to the central nervous system (CNS) in rats.32 These studies help to further emphasize prospective studies on ANS, CNS, and enteric pathways should be monitored in their role of pyloric, antrum and GE.

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**Fig. 2.** Interaction of interstitial cells of Cajal (ICC) cells and neuronal nitric oxide synthase (nNOS) signaling for normal gastric emptying (GE). NO, nitric oxide; Ach, acetylcholine; SMC, smooth muscle cell.
TREATMENT

The treatment of Gp requires wide consideration of multiple factors to best optimize care for the patient. Each patient should have a thorough evaluation of the past remedies and approaches to help tailor their future management. Optimizing oral nutrition should be the main goal. As discussed, the complex pathogenesis involving the CNS, ANS, inflammatory and neuromuscular dysfunction all contribute to management. See Fig. 3 for this complex interaction. The severity of Gp should always be assessed and one commonly used questionnaire is the validated GCSI. A severity index has also been created classifying patients as either grade 1-mild Gp, grade 2-compensated Gp, and grade 3-Gp with gastric failure, in order of severity. See Table 4 for this outcome score. Several studies have also supported use of patient-reported outcomes as correlating with severity of the disease. This tool has been shown to be useful as well in helping providers assess the current state of treatment.

**Table 4.** Patient-Reported Gastrointestinal Outcomes and Total Symptom Score

| Symptom                | Frequency | Severity | Average |
|------------------------|-----------|----------|---------|
| Vomiting               | 0 1 2 3 4 | 0 1 2 3 4| 0 1 2 3 4|
| Nausea                 | 0 1 2 3 4 | 0 1 2 3 4| 0 1 2 3 4|
| Anorexia/early satiety| 0 1 2 3 4 | 0 1 2 3 4| 0 1 2 3 4|
| Bloating/distension    | 0 1 2 3 4 | 0 1 2 3 4| 0 1 2 3 4|
| Abdominal pain         | 0 1 2 3 4 | 0 1 2 3 4| 0 1 2 3 4|
| Total GI symptom score | Sum of above | Sum of above | Sum of above |

The total symptom score instrument is used as a patient-reported outcomes tool. Rated as 0 to 4, none to worse and total (0–20). GL, gastrointestinal.

Adapted from Cutts T, et al. BMC Gastroenterol 2016;16:107.ât

**Fig. 3.** Conceptual framework for pathophysiology of gastroparesis syndromes.

CNS, central nervous system; ANS, autonomic nervous system; ENS, enteric nervous system; PNS, parasympathetic nervous system; SNS, sympathetic nervous system.
and whether adjustments should be made. The hallmarks of treatment are to correct nutritional deficiencies, decrease symptoms, and pursue the etiology in the hopes of rectifying the underlying cause of Gp. Combination strategies that focus on these three principles are the key strategies for management. More than 10% unintentional weight loss over a 6 month period signifies severe malnutrition. Please refer to Fig. 4 for a quick algorithm for the diagnosis and management of gastroparesis syndromes.

1. Diet

Diet modification is essential to help improve symptoms and should be emphasized as part of the treatment plan. Diets should consist of multiple small meals and should be limited in their fat and fiber content since those tend to delay emptying. Meal frequency should have a goal of at least 4 to 6 meals, and supplementation with high caloric liquids can help with nutrition without causing symptoms in mild Gp. Food should be chewed well and consumed in small-portioned bites. Supplementation with a multivitamin is helpful as well if tolerated. A comprehensive diet history should also be performed and foods that seem to aggravate Gp in the specific patient should be avoided, such as dairy or certain meats. Patients with a high intolerance of liquid supplementation may signal that tolerance of solid food will be even more difficult. A few studies have even shown alcoholic beverages, carbonated drinks, and cigarette smoking should be avoided as modifiable factors that affect emptying. In addition, heavy lipid containing foods should also be avoided since they potentially also slow down emptying.

2. Enteral nutrition

For patients who are having difficulties with oral nutrition, enteral nutrition using a jejunostomy feeding tube may be the preferred solution. A naso-jejunal (NJ) tube should be placed first to see if enteral feeding past the stomach helps with the symptoms. Nasogastric tubes and gastrostomy tubes are not recommended since symptoms of Gp still persist and the risk of aspiration for pneumonia is higher. NJ tube feeding should start slow at 25 to 50 mL an hour using feeds consisting of 1.5 calories per mL and advanced as tolerated by 10 to 25 mL an hour. Feeding should be done continuously 24 hours a day until nutrition improves where the patient can tolerate an oral diet better with the eventual goal to have nocturnal feeding to allow for normal daily activities. Enteral feeding should be the preferred method over total parenteral nutrition (TPN), which should be avoided if possible. TPN can cause complications with infection, access problems, and thrombosis, and may only be necessary if dysmotility involves the small bowel to help provide supplemental nutritional support. Parenteral support should be the last resort.

3. Hyperglycemia

Besides focusing on nutrition, regulation of blood sugar should be extremely important as well to help reduce the symptoms of Gp. Poor glycemic control inhibits GE and even interferes with emptying tests. Numerous studies have shown GI symptoms manifesting in patients who already have other diabetic complications such as neuropathy and retinopathy. A few studies have even shown especially in type 1 diabetics, blood glucose levels between 288 and 360 mg/dL have been shown to inhibit both solid and liquid emptying. Even in healthy individuals without Gp symptoms, acute hyperglycemia induces reduction of antral motor activity. Furthermore, correlation between increases in glucose alongside increases in symptoms of post-prandial fullness. All of these studies illustrate that glycemic control is important in optimization of Gp care in the reduction of symptoms and motility.

4. Prokinetic medications

Prokinetic medications have long been a mainstay of treatment. Dopamine receptor antagonists work by inhibiting the effects of dopamine on GE and also as a serotonin antagonist as
well to help with nausea. Metoclopramide is the only U.S. Food and Drug Administration (FDA) approved dopamine receptor antagonist specifically approved for the treatment of Gp. Studies comparing metoclopramide versus placebo improvements in Gp symptoms showed it helped decrease delayed emptying on scintigraphy.46-50 Metoclopramide is available in intravenous, intramuscular, oral, and liquid form and carries a black box warning of tardive dyskinesia. The intravenous form is more often used in hospitalized patients suffering from Gp. Metoclopramide has also been shown to increase symptoms of drug-induced parkinsonism, and akathisia compared to controls. Other side effects such as fatigue and drowsiness may limit the tolerance for the medication in up to 30% of patients. Increases in prolactin stimulation from dopamine antagonism can cause galactorrhea and menstrual irregularities in women. Increasing doses affect the incidence of these symptoms so starting at a low dose of 5 mg 3 times a day (tid) before meals and titrating up as tolerated is recommended.5,35,51-53 Maximum dose is 40 mg/day and the drug should not be used for more than 12 weeks without careful consideration of the risks and benefits due to the extra pyramidal side effects. Reduction of the doses and skipping days as tolerated should be encouraged to help limit symptoms.

Domperidone is also a dopamine receptor antagonist with the same efficacy but known for less extrapyramidal side effects compared to Metoclopramide since it does not cross the blood-brain barrier with starting doses of 10 mg tid with titration up to 20 mg tid and at bedtime. Max doses of 80 mg to 120 mg a day may be needed for at least 1 month to determine the efficacy of the medication. One study examined the medication’s effects in Parkinson’s patients and found that it reduced GI symptoms from levodopa therapy and accelerated GE of solid meals. Therefore, the medication is great for Parkinson’s patients with Gp symptoms. The main concern for the medication is QT prolongation and should not be given if the corrected QT is longer than 470 ms in males and 450 in females so an electrocardiogram should be performed.1,35,54 Unfortunately the medication is not FDA approved in the United States and obtaining the medication from other countries is discouraged by the FDA. An approved method would be through a FDA investigational new drug application with signature of informed consent and then subsequent purchase through an FDA-approved pharmacy.55 Another drug like Metoclopramide and Domperidone is Itopride, which has been studied for use in patients with dyspepsia and menstrual irregularities in women. Increases in prolactin have shown increased antral contractility with medication and a longer half-life than Erythromycin.62-65 There is a need for further investigation of chronic use of Azithromycin for use in Gp. Other motilin receptor agonists that are not antibiotics are currently being investigated for use as well. One agent called Mitemicinal appears to increase antral contractility. A few randomized, double blind studies with placebo and multiple different doses of Mitemicinal showed increased emptying in both DG and IG.66-69 The potential for benefit compared to the macrolide antibiotics is a decrease in side effects such as QT prolongation. One earlier motilin agonist, ABT-229, unfortunately was not proven to help with Gp.69

5. 5HT₄ receptor agonists

Cisapride is a 5HT₄ receptor agonist that leads to acetylcholine release and subsequent antral contraction and increased GE.70-73 It was initially FDA approved but was subsequently removed after cardiac arrhythmias developed due to QT prolongation and was removed in 2000. Tegaserod is also a 5HT₄ agonist and has already been in use in the treatment of constipation in irritable bowel syndrome and has been shown to help with GE. The advantage of Tegaserod was that it did not have the QT prolonging effects that Cisapride has. One study published the use of the Tegaserod in critically ill patients with impaired motility with resolution within 24 hours of administration.74,75 Further studies with larger sample sizes need to be completed to determine if Tegaserod could have a greater impact in Gp patients; however, Tegaserod is also not easily obtained.

6. Antiemetic therapy

Antiemetic agents have widely varying mechanisms and their use is recommended as second line behind the previously mentioned prokinetic agents such as Metoclopramide. The most common used drugs are phenothiazines (Prochlorperazine), antihistamines (Promethazine), and 5-HT₄ receptor antagonists (Ondansetron). These medications all carry side effects such as QT prolongation. Studies examining Ondansetron have shown every 8 hours by IV was shown to be helpful. In double blind studies, administration by IV was shown to normalize GE for solids and liquids.58 Studies examining both IG and DG patients on chronic oral Erythromycin showed improved emptying 4 weeks from the beginning of treatment.61 The drawbacks to motilin medications are the down regulation of motilin receptor with the effect of tachyphylaxis. This effect can happen within days of administration, so close monitoring and titration of oral Erythromycin is needed for therapeutic purposes. Lowest effective doses are recommended. Problems with cytochrome P450 interactions can also limit the use of the medication and cause more symptoms and carries a risk of sudden cardiac death as well.79 Another macrolide antibiotic, Azithromycin, has been shown to be equally effective as Erythromycin but without the cardiac risk and cytochrome interactions. Studies with Azithromycin have shown increased antral contractility with medication and a longer half-life than Erythromycin.62-65 There is a need for further investigation of chronic use of Azithromycin for use in Gp. Other motilin receptor agonists that are not antibiotics are currently being investigated for use as well. One agent called Mitemicinal appears to increase antral contractility. A few randomized, double blind studies with placebo and multiple different doses of Mitemicinal showed increased emptying in both DG and IG.66-69 The potential for benefit compared to the macrolide antibiotics is a decrease in side effects such as QT prolongation. One earlier motilin agonist, ABT-229, unfortunately was not proven to help with Gp.69
that it is not superior to Metoclopramide or Promethazine in patients when presenting to the emergency department and also are associated with QT prolongation.\textsuperscript{75,76} Studies also examined tricyclic antidepressants (TCA) in Gp. Due to the anticholinergics effects of TCA’s they could potentially worsen symptoms and cause further decreased GE.\textsuperscript{77,78} These anti emetic agents, while commonly dispensed to hospitalized patients, should be considered only after failure of prokinetic agents in Gp.

7. Gastric electrical stimulators

Gastric electrical stimulators (GES), have been approved for use by the FDA for drug-refractory IG and DG. The device is directly implanted into the stomach. For patients who are refractory to medications, GES has been proposed as a therapeutic option. The device consists of electrodes surgically placed into the anterior wall of the stomach, which are connected to a pulse generator. An early study in 2002 examined 38 patients with significant symptoms of Gp and found that 33 of them responded very well to temporary GES stimulation and underwent permanent stimulators. Patients were followed for a year and 97% experienced a reduction in symptoms of nausea and vomiting and the average weight gain was 5.5%.\textsuperscript{79} The most important aspect is that long-term data also showed a decrease in morbidity from Gp even 10 years after placement of the stimulator. Remarkably, hospitalizations, medications required, and symptoms were all decreased (p<0.05). The study involved DG, IG, and postsurgical Gp patients and all three groups saw weight increases after stimulation implantation. Even more astounding was that 89% of these patients were able to tolerate nutrition well enough to have their J-tube removed.\textsuperscript{80} With such great results a comparison of GES versus medical management was completed. Eighteen patients were enrolled, nine in the GES arm and nine in the medical management arm and parameters, symptoms, and hospital costs were compared after these patients were followed for 3 years.\textsuperscript{81}

The role of GES has even been investigated in the treatment of patients with GLS, which as mentioned previously have non-delayed gastric scintigraphy. One large study of 551 patients examined temporary GES placement after an emptying scan was performed. Patients were then classified into three groups: delayed emptying, normal emptying, and rapid emptying. Patients in all three subgroups had improvements in symptom scores; this demonstrates that patients with GLS should also be considered for stimulators and further supports the belief that GLS is part of the spectrum of Gp.\textsuperscript{82}

8. Surgical treatments

Besides GES, other modes of surgeries are being explored that aim to anatomically reduce resistance to GE. Surgical procedures range from pyloroplasty to partial or total gastrectomy to help bypass the stomach. One article reviewed surgical procedures for Gp and came to the conclusion that gastrectomy and long-term outcomes of Gp need to be further studied since only 26 subjects in two trials had been reviewed.\textsuperscript{83} Clinical guidelines for surgery are for patients who are being severely affected by Gp where risk of renal failure and shortened life expectancy is severely limited, with the intention to relieve effects of nausea and vomiting.\textsuperscript{1} One alternative, less invasive surgery than gastrectomy is laparoscopic pyloroplasty. One study involved 28 patients who underwent minimally invasive pyloroplasty, and saw improvements in GES times, symptom scores, and medication use; 83% of the patients improved.\textsuperscript{84} Overall, more research studies on surgical treatments for Gp need to be conducted to determine the long-term outcome and possibly to help identify those patients most at risk with mortality from Gp who have already tried traditional prokinetic and antiemetic therapies.

CONCLUSIONS

Progress has been made in the last few decades in understanding the pathophysiology of Gp and in the various treatment plans. The focus at the cellular level and function of cells that cause gastric dysfunction are key to understanding the pathogenesis of this disease. The term gastroparesis-like syndrome was created to help classify those patients who have all the symptoms of Gp without the objective decreased emptying scan. Ultimately, some of these patients may progress to Gp considering the progression of disease relating to the number of ICC cells seen. Some drug-refractory Gp patients have even been shown to benefit from GES. Other drug and device-resistant patients may be helped with immunotherapy. Progress towards understanding the pathophysiology of patients with Gp syndromes may help us treat patients with such profound symptoms that affect their quality of life.

CONFLICTS OF INTEREST

Thomas Abell is a speaker and investigator for Medtronic, is an investigator for Theravance and has IP through the University of Mississippi, now licensed to ADEPT-GI, of which he is the founder. He is also an editor for MedStudy and a reviewer for UpToDate. Nanlong Liu has no conflicts of interest to disclose.

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