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Pathophysiology and management of diabetic gastroenteropathy

Theresa Meldgaard, Jutta Keller, Anne Estrup Olesen, Søren Schou Olesen, Klaus Krogh, Mette Borre, Adam Farmer, Birgitte Brock, Christina Brock and Asbjørn Mohr Drewes

Abstract: Polyneuropathy is a common complication to diabetes. Neuropathies within the enteric nervous system are associated with gastroenteropathy and marked symptoms that severely reduce quality of life. Symptoms are pleomorphic but include nausea, vomiting, dysphagia, dyspepsia, pain, bloating, diarrhoea, constipation and faecal incontinence. The aims of this review are fourfold. First, to provide a summary of the pathophysiology underlying diabetic gastroenteropathy. Secondly, to give an overview of the diagnostic methods. Thirdly, to provide clinicians with a focussed overview of current and future methods for pharmacological and nonpharmacological treatment modalities. Pharmacological management is categorised according to symptoms arising from the upper or lower gut as well as sensory dysfunctions. Dietary management is central to improvement of symptoms and is discussed in detail, and neuromodulatory treatment modalities and other emerging management strategies for diabetic gastroenteropathy are discussed. Finally, we propose a diagnostic/investigation algorithm that can be used to support multidisciplinary management.

Keywords: diabetes mellitus, complications, diabetic neuropathies, enteric nervous system, gastrointestinal motility, gastrointestinal transit, pharmacology, enteropathy

Introduction
The development of diabetic polyneuropathy can potentially alter neuronal structure and function anywhere in the peripheral, central and enteric nervous systems. Where small and large fibres (unmyelinated C and myelinated Aβ and Aδ fibres) in the somatic sensory nervous system are affected, the typical manifestation is a distal symmetrical polyneuropathy. Importantly, these symptoms can also be found in people with prediabetes. The somatic polyneuropathy is characterised by alterations in the sensory system, and classically presents with marked changes in a number of sensations (including temperature and fine touch, balance, etc.) and can be painful or painless, although the former predominates. In community samples with diabetes, the prevalence of clinical symptoms is in the order of 30%, although this is likely to represent an underestimate as many people are not formally diagnosed.

Although often overlooked, effects in the autonomic nervous system most likely occur concurrently with those observed in the somatic nervous system. For example it is plausible that early damage to small fibres, preceding large fibre neuropathy, of the somatic nerves, which can be observed in skin biopsies also take place in other small fibres such as in the enteric nervous system. Autonomic neuropathy can be clinically silent and is often present when polyneuropathy of the somatic nerves is present. The diagnosis is typically based on cardiovascular abnormalities. In particular, studies of RR-complexes in electrocardiograms have been used to describe heart rate variability, leading to the diagnosis of cardiac autonomic neuropathy. However, autonomic neuropathy also leads to gastrointestinal (GI) complications such as diabetic gastroenteropathy, which may affect the entire length of the GI tract. The underlying pathophysiology, which is driven by a multitude of factors,
including microenvironmental factors such as hyperglycaemia, have a negative impact on the enteric motor and sensory functions and manifest as symptoms related to, e.g., motility and secretory dysfunctions.5–7 Hence, people with diabetic gastroenteropathy may experience a variety of burdensome symptoms, including dysphagia, dyspepsia, pain, bloating, diarrhoea, constipation and faecal incontinence, all of which adversely influences quality of life.

Although the focus of this review is diabetic gastroenteropathy, it should be emphasized that functional GI disorders are prevalent in the community. Hence, the presentation of diabetic gastroenteropathy and functional disorders such as functional dyspepsia and irritable bowel syndrome overlap and cannot be distinguished on the basis of symptoms or medical history (i.e. diabetes) alone. In fact, examinations of small bowel biopsies from patients with diabetes revealed neuropathy and myopathy only in a minority of patients.8 It is also clear that individuals with chronic disease have a higher prevalence of psychiatric disorders and are exposed to more stress than healthy subjects. These factors play an important role in the presentation and perceived severity of functional GI diseases.

The aims of this review are, however, to describe the pathophysiology of GI complications to diabetic neuropathy (gastroenteropathy) and to outline diagnostics and recent pharmacological and nonpharmacological treatment modalities for this burdensome condition.

Pathophysiology of diabetes-induced gastrointestinal complications
The term diabetic gastroenteropathy encompasses the cumulative impact that diabetes exerts on the GI tract. The pathophysiology is multifactorial and to date remains incompletely understood. However, changes in the neuronal and microenvironment are believed to be a major driver in the pathogenesis. Microvascular complications to diabetes lead to alterations of blood flow in the GI wall, and hence also to alterations in the microenvironment. Smooth muscle myopathy is also thought to be a contributing factor to diabetic gastroenteropathy. However, smooth muscle myopathy may not be a primary disturbance, but is more likely a result of smooth muscle cell atrophy induced through reduced trophic cues from interstitial cells of Cajal (which are also reduced in number).9 This link between reduced numbers of interstitial cells of Cajal and smooth muscle myopathy has been observed in animal models of diabetic gastroparesis and likely contributes to abnormal motility.

The autonomic nervous system comprises (a) the sympathetic nervous system, (b) the parasympathetic nervous system (whose main neural substrate is the vagus nerve) and, according to the early and recently re-established definition, (c) the enteric nervous system.10,11 Alterations in either of these systems are involved in the underlying mechanism of burdensome GI complications in diabetes. As both the enteric nervous system and central nervous system (CNS) are involved in the bidirectional regulation and control of the GI homeostasis, any changes in either of these interconnected systems may result in altered GI function.

Changes at the level of the enteric nervous system
In diabetes, the microenvironment within the enteric nervous system is significantly altered due to the effect of e.g. long-term hyperglycaemia; oxidative stress; inflammation; reduced levels of neurotransmitters, local hormones and nerve growth factors; and increased levels of fatty acids.12,13 Recently, altered gut luminal microbiota has also been proposed to exert an influence. For example, lowered numbers of bacteria involved in production of short chain fatty acids have been observed in diabetes. Short chain fatty acids have anti-inflammatory effects in the GI wall and promote the secretion of the incretin hormone glucagon-like peptide-1 (GLP-1). Lower levels of GLP-1 influence glucose metabolism and can increase low-grade inflammation.14–18

Various components of the enteric nervous system, (including enteric neurons, interstitial cells of Cajal, enteric glial cells) and smooth muscle cells are affected by the changes described above. Enteric neurons and interstitial cells of Cajal are particularly vulnerable to hyperglycaemia.19 When hyperglycaemic episodes are frequent, or when hyperglycaemia is persistent, shifts in the intracellular glucose metabolism of neurons occur, consequently leading to the formation of advanced glycation end-products, osmotic and oxidative stress as well as inflammation. Collectively, this
leads to cellular damage and ultimately to cell death, a process often referred to as glucose neurotoxicity. These mechanisms are primarily described in the peripheral nervous system, but similar mechanisms are present in the enteric nervous system.20

The pathophysiological changes described above may lead to various degrees of diabetes-induced damage to enteric neurones. Preferential loss of large fibre neurons in the dorsal root ganglion and inhibitory motor neurons in the gut wall have been observed. In particular, selective loss of nitric oxide synthase and neuropeptide Y expressing inhibitory motor neurons has been shown in the human diabetic colon.21 This obviously has consequences for the contractile activity of the smooth muscle layers in the GI tract and hence the motility pattern. Furthermore, loss of interstitial cells of Cajal throughout the GI tract has been reported in both animal models and in humans,22,23 causing reduced frequency of spontaneous muscular contractions. In addition, decreased activity of gastric enteric glial cells has been observed in animal models of diabetes.24 This may contribute to the development of gastrointestinal neuropathy in diabetes mainly because enteric glial cell functions such as neurotrophic support, immunosuppression and anti-inflammation is diminished with decreased activity.25,26

In contrast to the loss of motor neurons, it has recently been shown that levels of neurones containing substance P and calcitonin gene-related peptide are increased in the stomach of porcine models of diabetes.27 Both these molecules are primarily involved in the afferent transmission of GI sensory and nociceptive information. Hence, such alterations may be related to the pathophysiological mechanisms of sensory symptom generation. Like in the somatic system, when sensory nerves are affected by enteropathy, pain from the gut can likely be spontaneous or evoked by external or internal stimuli such as disturbances in motility and glandular functions. On the other hand, previous experimental studies where the gut was stimulated with electrical, mechanical and thermal stimuli, hypoalgesia to peripheral stimulation of both the upper and lower gut were shown, likely reflecting abnormal central processing of the afferent activity.28 Moreover, rectal pain thresholds are correlated to cutaneous and autonomic dysfunction.29,30 The autonomic components of visceral hyperalgesia have only been investigated in detail in healthy subjects, but similar mechanisms are likely involved in diabetes.31

Changes at the level of the autonomic nervous system

Autonomic afferent and efferent signalling through the vagus nerve is directly involved in the extensive communication with the brain, forming the so-called gut-brain-axis. In both people with diabetes and in animal models, the number of neurons in the sympathetic and parasympathetic ganglions of the vagus nerve connected to the GI tract is reduced32–34 as well as structural changes in the axons have been reported.35,36 In consequence, autonomic neuropathy, influencing the vagus nerve, contributes to reduced GI function.

Changes at the level of the CNS

Although the blood–brain barrier offers some protection to the brain against hyperglycaemia, it has been observed that the microstructure in specific brain regions (diabetic encephalopathy) involved in visceral sensory processing is changed in diabetes.37 Manifestation of sensory symptoms from the abdomen, such as nausea, vomiting and pain, as well as unspecific fullness, shooting sensations, etc., may relate to abnormal function of the sensory visceral nerves in combination with the encephalopathy (Figure 1).

We have previously shown that central reorganisation of brain responses to visceral stimuli is associated to the burden of GI symptoms as well as heart rate variability as a proxy for diabetic autonomic neuropathy.38,39 This was supported by another study where visceral hyposensitivity was correlated to an increase in somatic referred pain areas, indicating central neuropathic-like changes.28 Although controversies exist regarding the pathogenesis of these changes, the findings in the CNS may be secondary to the peripheral neuropathy, because the reduced afferent activity may cause adaptive shrinking.37,40 Finally, there is evidence from both neurophysiological and imaging studies that descending pathways from the brainstem that normally ‘gates’ the incoming nociceptive barrage is malfunctioning in people with diabetes,30,41 Hence, as in somatic peripheral
neuropathy, abnormal central sensory processing and hyper-excitability may explain the visceral symptoms despite the peripheral hypoalgesia (as described above). Such neuroplastic mechanisms were also seen e.g. in people with chronic pancreatitis, which is thought to have a strong neuropathic component, and this validated the findings.

In summary, long-term diabetes induces marked structural and functional changes of the GI wall, parasympathetic, sympathetic and CNS alterations. Especially, intrinsic and extrinsic neuronal communication of the GI tract is altered. Taken together, this leads to burdensome panenteric alteration of GI sensation and function, including the biomechanics that drive GI motility.

**Diagnosis**

As outlined above, diabetic gastroenteropathy can affect the entire GI tract, and consequently symptoms are not only very heterogeneous, they are also unspecific. The cardinal symptoms are nausea, vomiting, bloating and early satiety, however, symptoms range from dysphagia and heartburn to faecal incontinence. Thus, based on the patient reported symptoms, it is hardly possible to distinguish sequelae of diabetes-associated GI dysfunction from organic diseases. This diagnostic dilemma is further aggravated by the fact that people with type 2 diabetes have an increased risk for a multitude of organic GI diseases, including reflux oesophagitis, gallstones and GI malignancies. An increased cancer risk has also been observed in type 1 diabetes. Moreover, increased prevalence of other diseases with an autoimmune component affecting the GI tract such as coeliac disease are frequent and need to be taken into consideration.

Accordingly, it is important to first exclude organic disease using appropriate laboratory tests, endoscopy and imaging techniques in people presenting with symptoms that could be attributed to diabetic gastroenteropathy (Figure 2). If these tests are unrevealing and symptoms do not respond to simple therapeutic measures (e.g. laxatives in chronic constipation), GI function tests can be used to diagnose disturbances associated with diabetic gastroenteropathy. Again, most clinically available function tests identify and quantify dysfunction of various GI segments and organs without being able to specifically prove the causation by or the relative importance of diabetic gastroenteropathy. For instance, severe oesophageal hypomotility diagnosed by high resolution manometry can explain nonobstructive dysphagia in a person with diabetes, but could also be due to other aetiologies. Therefore, clinical plausibility and affection of other organ systems, e.g. cardiac autonomic neuropathy, should be taken into account. Impaired pancreatic polypeptide release has been suggested for specific diagnosis of gastroenteropathy, but measurements are not widely available. Likewise, antroduodenojunal manometry with increased frequency of phase III-motility and hypomotility during phase II as well as postprandially is typical of autonomic neuropathy, but available only at highly specialized centres and reserved for people with very severe symptoms.
Gastric emptying tests are recommended at an early stage in people with diabetes and dyspeptic symptoms. One reason for this is that gastric emptying is of paramount importance for blood glucose control. Another reason is that approximately 20% of people with diabetes and disturbed gastric emptying have functional dumping syndrome, i.e. accelerated gastric emptying without prior upper GI surgery, in contrast to the more frequently reported complication gastroparesis. Both disturbances cannot be differentiated reliably based on symptoms, but obviously require different treatment strategies. Recommendations regarding optimal use of gastric emptying tests are reviewed in detail elsewhere. Even in the absence of dyspeptic symptoms, gastric emptying testing can play a role in exclusion of diabetic gastropathy as a cause of impaired blood glucose control not responding to antidiabetic medication.

Once settled on the diagnosis diabetic gastropathypathy, treatment options are not overwhelming, but the cornerstone is tight glycaemic control.

**New diagnostic modalities**

The following modalities are selected because they are directly applicable in clinical practice...
and therefore can be used to assess diabetic neuropathy.

13C-breath test. This test reflects the conversion of 13C isotope to 13C-CO by hepatic metabolism after absorption from the small intestine, thereby serving as a proxy for gastric emptying. This modality may not be as sophisticated as, for instance, magnetic resonance imaging or scintigraphy. However, this very accessible technique correlates well with scintigraphic data, can be used repeatedly in the same person, and may be a marked improvement in assessing gastric emptying in clinical situations.

The wireless motility capsule. This system compromises an indigestible capsule that continuously measures pressure, temperature and pH as it passes through the GI tract. Although the modality has been available for some time, the optimal yield from this technology is still to be determined. The pH measurements are used mainly to establish the GI segment. However, with the growing evidence of the influence of gut microbiota on gastroenteropathy and metabolism in general, one could speculate that associations between the pH of the different segments and composition of the gut microbiome could be of interest. For example, more acidic caecal pH has been demonstrated in type 1 diabetes, which may represent increased caecal fermentation.

3D-transit system. Finally, a number of emerging modalities are currently being developed for research use. The ambulatory Motilis 3D-transit system tracks electromagnetic capsules as they traverse the GI tract and measures changes in position, velocity of movements and orientation of the capsules. This reflects gut contractile activity and progression dynamics. Anatomical information allows for detailed description of colonic motility, including regional transit times and motor patterns.

Management

Management of blood glucose fluctuations

There is no cure for diabetic gastroenteropathy. Hence, the treatment aims are to delay progression, ease symptoms, manage complications and if possible restore function. The primary strategy to achieve this remains tight glycaemic control. Glycaemic management should be guided by age, disease duration and overall health and, if successful, symptoms may improve. Dietary and lifestyle advice can provide persons with diabetes with tools for better long-term glycaemic control. In people with diabetes and gastroparesis, it can often be helpful to administer pre-prandial insulin after the meal or in reduced amount. Employment of an insulin pump may further contribute to tight glycaemic control in persons with insulin-dependent diabetes. Devices that allow for continuous glucose monitoring in interstitial fluids in real time have become available and enable monitoring of time spent in target glucose range (“time in range”) as well as warning trends toward hypoglycaemia or hyperglycaemia in real time. This modality has already proven its potential in modelling intestinal glucose absorption and thus it can be expected to be a future important tool in validating glucose metabolism of the enteric system in physiological and pathophysiological setups. Continuous glucose monitoring is recommended by national and international medical organisations and expert clinician consensus guidelines, both in combination with pumps and in persons on multiple daily insulin injections. As the number of hyper- and hypoglycaemic events are reduced, the concept is believed to have a neuroprotective effect. Besides optimisation of glycaemic control, no available treatments address the underlying polyneuropathy. Hence available nonpharmacological and pharmacological treatment options targets symptoms of gastroenteropathy, the latter of which may be complicated by altered drug absorption in the diabetic gut.

Pharmacological management: absorption of medications

Widespread disease of the GI tract will have consequences for the absorption of orally administered drugs. However, only few studies on small populations have addressed the effects of diabetes on net absorption of drugs. As diabetes-induced structural and functional alterations are observed throughout the GI tract, diabetic gastroenteropathy may alter drug absorption after oral administration.

Two important factors may be affected: (1) The release of drug substance from the controlled release formulations, which are designed to release drug with a predefined rate throughout the GI-tract. Changes in the intraluminal environment due to gastroenteropathy may alter drug release. (2) Drug absorption following oral administration, which is
possible throughout the GI-tract, with the upper small intestine as the main site for absorption. Several changes of GI physiology and function related to diabetic gastroenteropathy may therefore influence drug absorption (Table 1). This can ultimately result in therapeutic failure due to altered plasma levels.

One study in diabetes demonstrated that alterations in gut transit time impacts mainly the pharmacokinetics of drugs with poor intestinal absorption and controlled release formulations. Further studies on disease–drug interactions are needed as the study on the influence of GI dysfunction on drug absorption from oral formulations is still in its beginning.

In conclusion, potential unpredictable drug absorption and the likelihood of treatment failure should be considered in people where diabetic gastroenteropathy may be present.

**Pharmacological management of motility dysfunction in the upper GI tract**

**Prokinetics.** A number of prokinetic drugs have been studied for the management of motility disturbances in the upper GI tract in people with diabetes, and have generally proven to be effective for symptom improvement in placebo-controlled trials. Until now, no absolute association between symptom improvement and changes in upper GI motility before or after treatment have been shown. This may be due to heterogeneity of study groups and use of sub-optimal methods for measuring gastric emptying time. Hence, two new metaanalysis showed that there may in fact be an association between upper GI motility and symptoms when optimal methods are used. On the other hand it cannot be ruled out that central effects of prokinetics may explain the observed improvement of symptoms. The classical prokinetic drugs fall into two categories based on their primary molecular target. The D₂-receptor antagonists metoclopramide and domperidone have been used to treat gastroparesis for many years and have proven effective for this indication in randomised placebo-controlled trials.

A multicentre study comparing the effectiveness of metoclopramide and domperidone found the drugs to be equally effective against symptoms of gastroparesis, but with more adverse effects in the CNS reported in persons treated with metoclopramide. This finding can be explained by the ability of metoclopramide to cross the blood-brain barrier to a higher degree, thereby having an increased potential for mediating central adverse effects. In February 2009, the U.S. Food and Drug Administration (FDA) and European Medicines Agency appointed black box warnings for long-term use (more than 12 weeks) of metoclopramide due to the risk of irreversible tardive dyskinesia, which has limited its use. The FDA has approved only metoclopramide for gastroparesis, although the risks for cardiovascular side effects seems to be higher than for domperidone.

In Europe, domperidone is most often used, but caution should be taken due to risk of cardiac arrhythmias in the presence of prolonged QT-syndrome. Risk factors such as

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**Table 1.** Diabetic gastroenteropathy related factors that may influence drug absorption

<table>
<thead>
<tr>
<th>GI factors influencing drug absorption</th>
<th>Consequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dysmotility (gastroparesis, functional dumping syndrome, diarrhoea and constipation inclusive)</td>
<td>Altered transit time and/or luminal water content</td>
</tr>
<tr>
<td>Small intestinal bacterial overgrowth</td>
<td>Altered pH and transit time</td>
</tr>
<tr>
<td>Altered secretory function</td>
<td>Reduced or increased luminal water content</td>
</tr>
<tr>
<td>Altered enteric microbiota</td>
<td>Altered pH and luminal drug metabolism</td>
</tr>
<tr>
<td>Structural remodelling of the wall of the GI tract</td>
<td>Altered intestinal transporters and gut wall metabolism</td>
</tr>
<tr>
<td>Reduced microvascular blood flow</td>
<td>Altered absorption to systemic circulation</td>
</tr>
</tbody>
</table>

GI: Gastrointestinal.
female sex, age above 65 years, electrolyte disturbances, polypharmacy, known heart disease, etc. should be taken into consideration when D₂-receptor antagonists are used. Erythromycin is a motilin and cholinergic receptor agonist that has also been used for decades to treat upper motility dysfunction in people with diabetes. However, its clinical efficacy often diminishes after 2–4 weeks due to tachyphylaxis, and many patients experience adverse effect during long-term use. A survey among people with gastroparesis observed a correlation between willingness to accept potentially lethal side effects and symptom severity. This emphasizes how burdensome GI complications are.

The prokinetic action of erythromycin is likely a drug class effect, and other macrolides with less toxicity may be used, but evidence from controlled trials is not available. Novel molecular targets, including highly selective serotonin agonists, are currently under investigation for debilitating symptoms associated with diabetic gastroparesis, but have not yet been approved for this indication. Prucalopride is currently available for other GI dysmotility disorders (see next section). It has a safe cardiovascular profile and may be used off-label for treatment in selected cases. The synthetic ghrelin analogue, relamorelin, with prokinetic properties, also appears to be safe and has shown promising results for the treatment of gastroparesis in phase IIA studies. Hence, relamorelin may be a potential drug to use in the future.

Tricyclic antidepressants. Low-dose nortriptyline, amitriptyline, and desipramine can diminish symptoms in people with diabetes and chronic vomiting, who had an inadequate response to prokinetics. A multicentre trial has reported that amitriptyline relieve symptoms, although gastric emptying time was not lowered, in subgroups of patients with painful functional dyspepsia. This indicates a mechanism of action of tricyclic antidepressants on visceral hypersensitivity and not gastric emptying. However, in a large multicentre randomised controlled trial in adults with idiopathic gastroparesis, the use of nortriptyline (up to 75 mg per day) compared with placebo for 15 weeks did not improve the overall symptom score. Thus, further research is warranted in order to make any conclusive recommendations.

Taken together, the serotonin and ghrelin receptor agonists have generally been well tolerated and safe in humans without the cardiac or neurologic adverse effects associated with ‘classic prokinetics’. Therefore, these agents (and potentially tricyclic antidepressants) comprise promising therapeutics for future treatment of upper motility dysfunction in diabetes and other upper GI dysmotility disorders.

Pharmacological management of motility dysfunction in the lower GI tract

Natural bulking, osmotic and stimulant laxatives. Lower GI symptoms in people with diabetes type 1 are associated with poor glycaemic control and quality of life. In randomised, placebo controlled trials, the intake of the natural bulking psyllium (10 g bid) or flaxseed (10 g bid) reduced symptoms of constipation and improved glycaemic control in people with type 2 diabetes. No studies have specifically investigated the effects of laxatives in persons with constipation as a complication due to diabetic gastroenteropathy. It is commonly suggested, but not evidence based, to start with osmotic laxatives such as magnesium or polyethylene glycol. If this is insufficient, stimulant laxatives such as bisacodyl, senna or picosulfate can be added. Lubiprostone, a chloride channel activator, increases secretion from the colon, thereby reducing colonic transit time and increasing the number of spontaneous bowel movements in persons with diabetes-related constipation. Prucalopride, a 5-HT₄ agonist, reduces transit time throughout the GI tract, and especially through the colon. Though not evaluated in DM, the pharmacological profile of prucalopride indicates that it is useful against constipation as part of the panenteric disorder often found in diabetes. Linaclotide, although registered for irritated bowel syndrome, may also be used in selected cases. Finally, the enzyme transglucosidase (used as a dietary supplement) changes the gut microbiome and increases the weekly number of bowel movements in people with type 2 diabetes and constipation. Antibiotics, enzyme supplementation and dietary intervention. In addition, people with diabetes have an increased prevalence of diarrhoea as sequel to small intestinal bacterial overgrowth, bile acid diarrhoea, pancreatic insufficiency and coeliac disease. These should be treated specifically with antibiotics, bile acid sequestrants, enzyme supplement or gluten-free diet whenever appropriate.
**Antidiarrhoeal products.** Compared to the general population, people with diabetes have a two-fold risk of having diarrhoea (11 vs 6 %).\(^{88}\) Very often, diarrhoea is induced by glucose-stabilizing treatment such as metformin or other medications commonly taken by people with diabetes. If no underlying condition is found for development of diarrhoea, pharmacological treatment will usually include dietary assessment and intervention in combination with loperamide, an opioid receptor agonist. Uncontrolled observational studies have shown that the \(\alpha_2\) adrenergic agonist clonidine may reduce diarrhoea in people with diabetes. The use of clonidine is, however, restricted by its cardiovascular side effects.\(^{89}\)

Furthermore, diabetes significantly increases the risk of having faecal incontinence, which in frequently is aggravated by diarrhoea. If an underlying cause of diarrhoea can be identified, it must be addressed. If not, loperamide or dietary intervention may be indicated. In a number of persons, faecal incontinence is associated with neuropathy and reduced sensibility of the anal canal. In these cases, loperamide, suppositories or enemas should be considered. Treatment of refractory cases is very complex and may even require a stoma.

**Neuromodulation.** Neuromodulatory electrical stimulation of the sacral nerve, which is described in detail later, is an emerging technique for the treatment of faecal incontinence and potentially sensitivity in the anal canal.\(^{90}\) However, the role of sacral nerve stimulation in people with diabetic gastroenteropathy has not been investigated specifically.

**Pharmacological management of sensory symptoms**

In the management of abdominal pain in diabetes it is often impossible to distinguish between the different organ manifestations. This is due to the vague presentation of visceral symptoms with changing presentation of the pain and referral to somatic structures, and this should be taken into consideration.\(^{91}\) It can also be difficult to distinguish between symptoms such as nausea and pain evoked by dysmotility and those that are related to the sensory (and central) neuropathy *per se*. In such cases the primary reason for pain such as constipation should be treated primarily. If all such reasons for the pain can be excluded, management should be directed against the neuropathy. Although only few studies have addressed treatment of visceral sensory symptoms in people with diabetes, the pharmaceutical options most often applied are reviewed here. On the other hand, the individual variability in phenotypical presentation of pain in diseases is greater between people than between pain syndromes. This indicates that mechanistic aetiologies and subsequent successful treatment should be based at the level of the individual rather than the disease *per se*.\(^{92}\) Hence, the recommendation for pharmacological management of visceral neuropathic pain follows the guidelines used for somatic neuropathies.\(^{93}\)

**Tricyclic antidepressants.** As stated above tricyclic antidepressive medications can be used to manage symptoms in gastroparesis, likely because many of the clinical presentations are consequences to sensory neuropathy. A detailed description is outside the scope of this paper, but for disorders with peripheral and central neuropathy, these adjuvant analgesics are often used.

**Selective serotonin-noradrenaline reuptake inhibitors and gabapentinoids.** Depending on the clinical situation, other pharmacological compounds for the use of treating diabetes induced somatic and neuropathic pain also include selective serotonin-noradrenaline reuptake inhibitors as well as the gabapentinoids (gabapentin and pregabalin), and the drugs can be used in combination.

**Opioids.** In difficult and severe cases, more potent analgesics such as opioids may be needed, but, if possible, long-term treatment should be avoided due to adverse effects, e.g. high risk of masking hypoglycaemic events.\(^{94,95}\) Although most drugs have a certain potency, all have GI (or CNS) side effects and safety is often the major limitation in pain management, especially for opioids.\(^{75,96}\) Hence, the balance between effect and side-effects is more relevant than the potency of the analgesics. This balance is however highly individual, and as no valid predictors for individualised treatment is available, a period with ‘trial and error’ is often necessary. There should also be awareness on opioid-induced bowel dysfunction that by itself may lead to gastroparesis and constipation and lead to a vicious circle that can intensify the pain.\(^{97,98}\)

**Modern pain management.** There is always more to analgesia than analgesics, and pharmacological management can seldom stand alone. Therefore,
modern pain management of diabetic gastroenteropathy also includes e.g., invasive procedures, supportive care and nursing (multimodal analgesia). Finally, it should not be forgotten that individual experiences and manifestations of pain are influenced by complex interactions between sensory, pathophysiological, affective, socio-cultural, behavioural and cognitive elements. An active screening for psychiatric comorbidity, including anxiety and depression, should be done as up to 40% of chronic pain patients are depressed, and identification of mood disorders may select persons where adjuvant therapy with antidepressants are particular beneficial.99 It should be stressed that treatment of abdominal pain secondary to diabetic gastroenteropathy is complex and involves a multidisciplinary approach including diabetologists, gastroenterologists, pain specialists, dietitians and psychologists.

Dietary management
People with diabetic gastroenteropathy are at risk of developing dehydration and poor nutritional status.100 Most studies on dietary treatment in diabetes have included people with gastroparesis who may have insufficient dietary intake due to early satiety, postprandial fullness, nausea and vomiting. However, the literature regarding the evidence for the nutritional intervention is scarce. Among people with gastroparesis, 64% had insufficient daily intake of energy, vitamins and minerals.100 In contrast, another study demonstrated that people with diabetes were capable of maintaining a sufficient daily caloric intake.101 The normal dietetic recommendation for diabetics includes a high-fibre content, which is not appropriate for people with gastroparesis. A small particle diet cause less upper GI symptoms than a conventional diet and represents food items that are easily processed into small particle size (maximum 2 mm in diameter). In a study by Olausson et al., the fibre and fat content of the diet was normal, but it excluded husks/peels (e.g. corn), membranes (e.g. orange), stringy foods (e.g. rhubarb), seeds and grains (e.g. whole grain), and poorly digestible particles (e.g. salad).102

It has been observed that gastric emptying is significantly delayed in healthy volunteers after a high-fat meal compared with a low-fat meal.103 However, the majority of well-controlled studies indicate that gastric emptying upon high-fat meals is similar to other nutrient compositions matched for calorie and volume.104 One explanation may be that high-fat meals increase visceral sensitivity. This may explain, why correspondingly, a high-fat solid meal causes more symptoms than a low-fat liquid meal in people with diabetic or idiopathic gastroparesis.105,106 Also, high-fat liquid meals were better tolerated than high-fat solid meals.107 High-osmolality liquids with more than 350 mOsm.kg delay gastric emptying in healthy persons108 but the effect in people with diabetes is unknown.

Since fat, fibre, meal-size and consistency of foods all seem to influence gastric emptying and symptoms, the dietetic intervention in people with diabetes and gastroparesis should include several small meals (five to six) with a small particle size, a moderate-to-low content of fat and fibre, and a high content of liquids both in and alongside the meal. In case of weight loss, high-fat liquid meals might be an option. Oral intake is preferred, but in people with severe symptoms and weight loss, enteral (particular jejunal) or parenteral nutrition can be indicated.109

Neuromodulatory treatment
Although not neuromodulatory per se, a novel nonpharmacological vibrating capsule is assumed to induce a normal peristaltic wave in the large intestine to alleviate constipation. Although it has been shown to improve transit times in some people with functional constipation by a conveying by vibration approach,110 further evidence is needed.

Electrical stimulation of the GI tract was first used over 50 years ago to improve motility in postoperative ileus.111 Subsequent studies demonstrated that gastric electrical stimulation could improve gastric emptying and gastric dysrhythmias, both important pathophysiological features of gastroparesis.112 Gastric electrical stimulation is currently used for the treatment of people with gastroparesis whose symptoms are refractory to medical interventions, particularly in the context of weight loss, however, it has not affected the transit time. The method is invasive and involves laparoscopic surgical placement of electrodes on the externa muscular wall of the gastric antrum (Figure 3a). These electrodes are then subsequently attached to a programmable signal generator box, which is implanted in a subcutaneous pouch in the left flank. The complication rates are in the order of
10%, the most common being subcutaneous pocket infection. The mechanism of action of gastric electrical stimulation is incompletely understood, but it is thought that it may either modulate vagus nerve function to improve gastric accommodation or influences vagal afferent signalling to the CNS. Outcome and efficacy data on gastroparesis in people with diabetes are largely derived from open-label single-centre studies in highly selected persons. Meta-analytic evidence, however, suggests that gastric electrical stimulation improves the cardinal symptoms of gastroparesis, such as nausea and vomiting, improves generic quality of life and reduces the need for enteral or parenteral nutritional support. It has been shown that acute hyperglycaemia inhibits anal sphincter function, leading to a reduction in rectal compliance, which can lead to faecal incontinence. Although the mainstay of management remains medical, a number of neuromodulatory therapies are emerging, such as sacral nerve stimulation or percutaneous tibial nerve stimulation. Typically, a test or trial stimulation period is undertaken where the signal generator is external, and, if successful, this can be internalised in a subcutaneous pouch (Figure 3b). The majority of the studies to date have used cross-over designs, which have demonstrated a degree of efficacy in reducing the number of episodes of faecal incontinence, although these have not been specifically designed for people with diabetes. Albeit not tested in diabetes, percutaneous tibial nerve stimulation is a novel ambulatory therapy for faecal incontinence (Figure 3c). However, a large multi-centre randomised controlled trial comparing 12 weeks of percutaneous tibial nerve stimulation with sham stimulation did not demonstrate any benefit over 12 weeks of treatment.

Non-invasive electrical stimulation is a promising novel approach for the treatment of abdominal symptoms that appears to increase gastric emptying and colon transit time. If proven, this would constitute a very applicable treatment approach.

Conclusion
The understanding of diabetic neuropathy has improved rapidly over the last decade. Increased understanding of the different symptoms, and how they interact with motor and secretory functions of the gut may be a major breakthrough in the treatment of the, often very diffuse, symptoms and complaints that have a major impact on quality of life. The management of diabetic gastroenteropathy is still difficult and should involve a multidisciplinary team including pharmacologists, nurses, dieticians, diabetologists, gastroenterologists and surgeons as well as health professionals from other specialities. On the other hand, new techniques to unravel the gut function as well as better treatment modalities are emerging. Together with increased awareness on the symptoms and better glycaemic control, diabetic gastroenteropathy and its different manifestations will undoubtedly be less burdensome in the near future.

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Figure 3. Neuromodulatory modalities in gastroenteropathy.
(a) A laparoscopic view of gastric electrical stimulation electrodes sutured to the wall of the gastric antrum. Photograph courtesy of Sri Kardirkamanthan, Broomfield Hospital, Essex, UK. (b) A schematic representation of sacral nerve stimulation demonstrating the electrodes and signal generator. (c) Percutaneous tibial nerve stimulation. In this photograph, the tibial nerve is being stimulated using a 34-gauge needle inserted in/around the tibial nerve with a cutaneous electrode on the sole of the foot.
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**Conflict of interest statement**

The authors declare that there is no conflict of interest.

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