The management of adult patients with severe chronic small intestinal dysmotility

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ABSTRACT
Adult patients with severe chronic small intestinal dysmotility are not uncommon and can be difficult to manage. This guideline gives an outline of how to make the diagnosis. It discusses factors which contribute to or cause a picture of severe chronic intestinal dysmotility (eg, obstruction, functional gastrointestinal disorders, drugs, psychosocial issues and malnutrition). It gives management guidelines for patients with an enteric myopathy or neuropathy including the use of enteral and parenteral nutrition.

1.0 FORMULATION OF GUIDELINES
1.1 Aim
This guideline aims to help clinicians manage patients who have or are thought to have severe small intestinal dysmotility that is causing malnutrition. It gives a logical guidance to determining the underlying diagnosis and shows how, knowing this, treatments may be tailored for each individual patient. It also aims to present some of the other factors contributing to the presentation and progress of the illness.

1.2 Development
The preliminary guidelines were compiled from the literature and a first document was drafted by Dr Nightingale and modified by members of the Neuro-gastroenterology and Motility Committee of the British Society of Gastroenterology (BSG) and by the Small Bowel and Nutrition Committee of the BSG. The article was reviewed by the patient organisation PINNT (Support and Advocacy group for people on home artificial nutrition) and modifications made to result in the current document.

The guidelines have applied the GRADE system. Strength of recommendation can be strong, moderate or weak. The evidence was graded according to the Oxford Centre for Evidence-based Medicine. Level 1 includes systematic reviews with homogeneity, individual randomised controlled trials (RCTs); level 2 includes systematic reviews of cohort studies, low quality RCTs and outcomes research; level 3 includes systematic reviews with heterogeneity and individual case-control studies; level 4 includes poor quality cohort or case series; and level 5 includes expert opinion without critical appraisal.

1.3 Scheduled review
The content and evidence base for these guidelines should be reviewed within 5 years of publication. We recommend that these guidelines are audited and request feedback from all users.

1.4 Service delivery
► Patients with severe chronic small intestinal dysmotility are not common but should be managed by a multidisciplinary team headed by a clinician with expertise in managing these patients. If managed appropriately, particularly if receiving parenteral nutrition (PN), there may be an improved quality of safe care and also considerable cost savings. (Grade of evidence level 4, Strength of recommendation strong)
► Any surgical procedure should be performed after extensive local/regional multidisciplinary team (MDT) discussions at hospitals with surgeons and histopathologists experienced in managing these patients. (Grade of evidence level 5, Strength of recommendation strong)
► Patients being considered for PN or a surgical bypass/resection should be seen at a dedicated PN centre or by an integrated intestinal failure (IF) unit respectively. (Grade of evidence level 5, Strength of recommendation strong)
► Regional networks may be established to ensure expertise is available to all and a national research forum may be established. (Grade of evidence level 5, Strength of recommendation strong)

1.5 Patients’ experience
► Patients with a severe chronic small intestinal dysmotility causing malnutrition should be managed as individuals; other factors including opioid use and psychosocial circumstances must be taken into account. (Grade of evidence level 5, Strength of recommendation strong)
► Patients will become experts in coping with their condition and management. All decisions should be made in conjunction with them. They are often more knowledgeable about their condition and treatments than the clinicians, nurses and dietitians and this should be respected. (Grade of evidence level 5, Strength of recommendation strong)
► Facilities for looking after these patients should be able to deal with physical,

emotional, psychological, social and quality of life issues. (Grade of evidence level 5, Strength of recommendation strong)

- Patients should be referred rapidly to places of more advanced expertise if management is difficult or unsuccessful. There should be a clinician, specialist nurse and/or a dietician available and a psychologist/psychiatrist to discuss the patients. The healthcare professionals should be familiar to the patient and know their history, thus eliminating the need for lengthy, time-consuming explanations. (Grade of evidence level 5, Strength of recommendation strong)

- The patients should be managed on wards or IF/home parenteral nutrition (HPN) units where the healthcare professionals are familiar with their specialist needs, especially if receiving PN. (Grade of evidence level 5, Strength of recommendation strong)

- Patients, especially if receiving HPN, should have access to a helpline (ideally 24 hours) so that emergencies are dealt with appropriately and quickly. (Grade of evidence level 5, Strength of recommendation strong)

- Written and audiovisual materials that explain and support the training techniques in enteral and PN should be available. (Grade of evidence level 5, Strength of recommendation strong)

- Meeting other patients with similar problems may improve the way in which a patient copes with their condition. (Grade of evidence level 5, Strength of recommendation strong)

- Where appropriate, patients should be offered contact numbers for the relevant support group that represents their specific disease. (Grade of evidence level 5, Strength of recommendation strong)

- Follow-up appointments should be as deemed appropriate to the MDT and patient, and ideally the patient should not have to travel long distances for appointments. The appointment should be with experienced and familiar staff, thus enabling continuity of care. (Grade of evidence level 5, Strength of recommendation strong)

- Staff should be aware of the latest research and developments and should make patients aware of any which may apply to them. (Grade of evidence level 5, Strength of recommendation strong)

For patients needing home parenteral nutrition (HPN)

- The techniques needed for HPN should be taught by competent and enthusiastic staff who can convey the confidence required to undertake the therapy successfully and safely. The procedures taught should follow the British Intestinal Failure Alliance (BIFA) standardised parenteral support catheter guidelines. (Grade of evidence level 5, Strength of recommendation strong)

- Patients need to know that an aseptic technique, in which the key parts are not touched, must be used whenever their feeding line is accessed. This is vital for safety and peace of mind. (Grade of evidence level 4, Strength of recommendation strong)

- All patients who require HPN whether short or long term should receive information about the patient support group ‘PINNT’ (Support and Advocacy group for people on home artificial nutrition) and should be registered with the British Artificial Nutrition Survey (BANS). (Grade of evidence level 5, Strength of recommendation strong)

2.0 SUMMARY OF RECOMMENDATIONS

2.1. Enteric myopathies are often a primary condition and have multi-viscer al involvement (especially of urinary tract) and these other manifestations should be sought. Enteric myopathies may be secondary to a muscular disease (eg, muscular dystrophy) and an awareness of this is needed when managing the primary illness. (Grade of evidence level 5, Strength of recommendation strong)

2.2. Patients with an enteric neuropathy often have a serious underlying condition (often neurological or metabolic) which should be sought. (Grade of evidence level 5, Strength of recommendation strong)

2.3. A definite diagnosis should only be given when a detailed history, symptoms and investigations (including histology in a unit that has expertise in this area of pathology) confirm the diagnosis. If, as is commonly the case, a definitive diagnosis is not possible, an empirical working diagnosis of probable severe dysmotility may be necessary. (Grade of evidence level 5, Strength of recommendation strong)

2.4. Mechanical obstruction must be excluded and occasionally (if radiology is inconclusive) this involves a trial of a low fibre (residue) diet or even a liquid diet. (Grade of evidence level 5, Strength of recommendation strong)

2.5. The contributing effects of drugs on gut motility (especially opioids and anticholinergics) must be taken into account. (Grade of evidence level 4, Strength of recommendation strong)

2.6. Psychosocial/behavioural issues often play a part in how the symptoms manifest and specialist psychosocial support must be available expediently. (Grade of evidence level 5, Strength of recommendation strong)

2.7. The effects of abrupt weight loss on the gut function must be taken into account. (Grade of evidence level 5, Strength of recommendation strong)

2.8. Investigations which include radiology, radioisotope studies, manometry, autoimmune screens and histology should be performed judiciously and in keeping with current guidelines, but may not give a definitive diagnosis. Care must be taken in interpreting them in the presence of some drugs (eg, opioids, cyclizine and anticholinergic drugs), severe malnutrition or an eating disorder. (Grade of evidence level 5, Strength of recommendation strong)

2.9. These patients are complex with multiple issues contributing to their presentation and so they need MDT management. The team is likely to include a gastroenterologist, gastrointestinal physiologist, gastrointestinal surgeon, pain team, psychiatrist/psychologist, rheumatologist (including fatigue management), urologist, gynaecologist, radiologist, dietitian, specialist nurses, clinical biochemist, histopathologist and pharmacist. Regional networking (can be via a virtual MDT) is encouraged both for support/guidance and to collect data. (Grade of evidence level 5, Strength of recommendation strong)

2.10. The aims of treatment for patients with severe chronic small intestinal dysmotility are where possible to:

- Reduce symptoms (eg, pain, vomiting, distension, constipation/diarrhoea, bloating/distension)

- Reduce morbidity and mortality

- Achieve a body mass index (BMI) within the normal range

- Achieve an improved/good quality of life. (Grade of evidence level 5, Strength of recommendation strong)

2.11. Treatment should be directed at the main symptom, using as few drugs as possible, avoiding high doses of opioids and cyclizine and avoiding unnecessary surgery. Try to avoid medicalisation (eg, enteral access, suprapubic catheters, etc) early in
the course of the illness. (Grade of evidence level 5, Strength of recommendation strong)

2.12. If the patient has taken long-term opioids, the narcotic bowel syndrome may have occurred and a gradual supervised opioid withdrawal should be considered. A pain specialist, if available, should be involved. (Grade of evidence level 4, Strength of recommendation strong)

2.13. If a patient is malnourished or at risk of becoming so, oral supplements/dietary adjustments should be tried. (Grade of evidence level 5, Strength of recommendation strong)

2.14. If feeding by the oral route is unsuccessful and if the patient is not vomiting, gastric feeding may be tried. (Grade of evidence level 5, Strength of recommendation strong)

2.15. If gastric feeding is unsuccessful, jejunal feeding initially via a nasojejunal tube may be tried and, if successful, a tube can be inserted endoscopically (through a gastrostomy (PEG)) or as a direct jejunostomy) or surgically. A jejunostomy also can be useful for drug administration. (Grade of evidence level 5, Strength of recommendation strong)

2.16. If jejunal feeding fails (often due to abdominal distension or pain as the feed is infused) and if the patient is malnourished, then parenteral support may be needed. (Grade of evidence level 5, Strength of recommendation strong)

2.17. A venting gastrostomy may reduce vomiting but can have problems (leakage, not draining, or poor body image). (Grade of evidence level 5, Strength of recommendation strong)

2.18. Nutritional status, when possible, should be optimised before any surgical procedure. A percutaneous endoscopic gastrostomy (PEG) or stoma is generally delayed in severely malnourished or physiologically unfit patients. (Grade of evidence level 5, Strength of recommendation strong)

### 3.0 KEY TERMS AND DEFINITIONS USED IN THE GUIDELINE

Many terms are used in the literature. Only chronic intestinal pseudo-obstruction (CIFO) has been consistently used in the past to imply a dilated gut that does not function. Other terms such as ‘enteric dysmotility’ have been used to define the ‘grey area’ sitting between the functional gastrointestinal disorders and more severe CIFO. Enteric dysmotility has been primarily defined by manometric abnormalities, although histological abnormalities can also be found. It lacks the small bowel dilation seen in CIFO and its current classification status remains uncertain.

In the past, different diagnostic labels have been given to patients based on the diagnostic test used. Unfortunately they are given to patients according to which diagnostic test is used. Currently they have not been merged to point to a specific diagnosis, which can be best given by histology. The histology may show abnormalities, but the extent they are due to malnutrition, previous surgery (including defunctioning bowel) or drug therapy is not completely clear.

**Phrases that may be found on investigations**

- Radiology
  - slow transit
  - dilated bowel
  - no transitional zone
- Scintigraphy
  - slow transit (gastric emptying, small bowel or colon) of liquid/solid
- Manometry
  - propulsive failure
  - giant contractions

- no migrating motor complexes (MMC)
  - Pathological
  - neuropathy
  - plexitis, leiomysitis
  - myopathy

In this document the term ‘intestinal dysmotility’ is used to cover all conditions in which there is a failure to propel the luminal contents without there being an organic obstructing lesion. The intestinal dysmotility is considered severe when nutritional or fluid support is needed due to objective evidence of malnutrition, dehydration or electrolyte disturbance.

This guideline relates to adults and does not cover acute pseudo-obstruction (eg, ileus).

### 4.0 INTRODUCTION

Chronic small intestinal dysmotility occurs when there is a failure of coordinated intestinal propulsion, giving rise to the symptoms and signs of intestinal obstruction (colicky abdominal pain, nausea, vomiting usually with abdominal distension, and often a dilated bowel) in the absence of a mechanical cause. A frank obstructive picture with distended bowel is not always the case, especially if there is a neurological aetiology. For the purpose of this article, chronic will be taken as more than 6 months of symptoms. Chronic intestinal dysmotility may be defined as severe when there is associated malnutrition (BMI of less than 18.5 kg/m² or more than 10% unintentional weight loss in last 3 months) and thus specific treatments which may include clinically assisted nutrition and hydration (CANH, formerly referred to as artificial nutritional support) may be needed.

Normal gastrointestinal motility is determined by intestinal smooth muscle function which in turn is influenced by neural and humoral factors. A disorder of one or more of these systems can result in intestinal pseudo-obstruction. There are many causes of acute or reversible/temporary intestinal dysmotility which include abdominal surgery, trauma, sepsis, metabolic (eg, hypokalaemia) or endocrine problems (eg, hypothyroidism), but they will not be discussed in this document as they do not usually result in long-term malnutrition. Dysmotility may occur in the small bowel and also in other areas of the gastrointestinal tract (eg, oesophagus, stomach and colon) and their involvement may complicate diagnostic tests and treatments. The primary problems with isolated oesophageal, gastric or colonic motility are not specifically discussed. Other medical causes of abdominal pain (eg, familial Mediterranean fever, angio-oedema, abdominal migraine and lead poisoning) are not specifically discussed but the clinician must be aware of them.

The diagnosis of these patients can be very difficult and may be empirical, especially when, as is most common, there is no definite histological confirmation of a disease process. The clinical features, results of investigations (eg, manometry) and histology may not all combine to indicate one specific diagnosis. Other factors that can give rise to the clinical picture or aggravate the condition are: unrecoégnised organic small bowel obstruction/the effects of previous abdominal surgery (including adhesions and neuropathic pain), drug usage (particularly opioids and drugs with anticholinergic effects), psychosocial problems including abnormal illness behaviour and malnutrition. In practice, all of these often play a part and contribute to the patient’s presentation. Untangling which of these factors gives rise to the dominant symptom can be challenging and needs the help of a MDT consisting of a gastroenterologist, gastrointestinal physiologist, gastrointestinal surgeon, pain team, psychiatrist/psychologist, rheumatologist (including a specialist in fatigue management),...
urologist, gynaecologist, neurologist, clinical biochemist, histo-
pathologist, radiologist and nutritional support team.

If there is uncertainty about the diagnosis, this should be clearly
documented and the diagnosis only described as working (prob-
ably or possible) and the contributing factors to this should be
stated on the patient’s problem list (eg, previous surgery, opioid
or cyclizine use, psychosocial problems or undernutrition). A
definitive diagnosis should only be given if there is a clear cause
identified. It is very difficult to remove a diagnostic label once it
has been given, and a premature or erroneous organic diagnosis
in those with predominantly psychosocial issues or abnormal
illness behaviour may make the management of contributing
issues very difficult. A definite diagnosis, although satisfying
to have, rarely affects the patient’s clinical management from a
medical perspective.

This document discusses the differential diagnoses, the medical
and nutritional treatment of chronic small intestinal dysmotility
(myopathy and neuropathy) which result in malnutrition. In its
most severe form, patients with small bowel dysmotility may
need long-term PN or even a small intestinal transplant, while in
a milder form dietary adjustments may suffice.

5.0 CONDITIONS THAT MIMIC OR CAN CONTRIBUTE TO THE
PRESENTATION

A patient with suspected small bowel dysmotility will have
had basal investigations to exclude other causes; these include
inflammatory markers (CRP, albumin, platelets and faecal
calprotectin) which, if normal, make active inflammatory bowel
disease unlikely. These may be followed, as appropriate, with the
use of endoscopies and cross-sectional imaging including with
intravenous contrast to diagnose structural/mucosal diseases.
Several other conditions may appear as severe chronic intestinal
dysmotility but with no primary bowel pathology (figure 1). In
one series the most frequent misdiagnoses for dysmotility were
volvulus, megacolon and chronic constipation.6

5.1 Organic obstruction

A major problem, that is often not diagnosed, is a localised
bowel obstruction as a result of adhesion formation. This may be
suspected clinically when a patient has had a number of abdomi-
nal operations (with or without extensive adhesion division).7
A history of intermittent colicky abdominal pain with abdomi-
nal distension, loud bowel sounds, no bowel or stoma action
and vomiting suggest this. A distal obstruction is suggested if the
vomit is faeculent, while a more proximal one by green/yellow
vomit. During an obstructive episode the bowel secretes more
fluid, and when the obstruction resolves, diarrhoea follows (or
a high stomal output). If a patient sticks to a low residue diet
fluid, and when the obstruction resolves, diarrhoea follows (or
vomit. During an obstructive episode the bowel secretes more
vomit is faeculent, while a more proximal one by green/yellow
and vomiting suggest this. A distal obstruction is suggested if the
A history of intermittent colicky abdominal pain with abdom-
inal operations (with or without extensive adhesion division).7
suspected clinically when a patient has had a number of abdom-
inal CT scan when the patient has an episode of severe pain.
Contrast follow through studies or MRI scans, although useful
when positive, may not be tolerated in the acute setting and do
not always demonstrate the obstruction. Unsuspected diagnoses
may be revealed (eg, small bowel volvulus from a band adhesion
or an intussusception). Further clues to an organic obstruction
are visible small bowel peristalsis, worse pain after prokinetic
drugs or giant jejunal contractions on manometry.9 9

Multiple laparotomies themselves may result in secondary
dysmotility, especially if the bowel becomes encased in fibrous
tissue as can occur with sclerosing peritonitis. In addition,
upper gut surgery (eg, a vagotomy, Whipple’s resection, gastro-
enterostomy, bariatric procedures or any bowel anastomosis) can
result in secondary small bowel dysmotility.10

Radiation damage can cause strictures and a localised obstruc-
tion and/or a generalised secondary dysmotility. Problems caused
by radiation damage tend to be progressive over many years.

5.2 Opioid and other drug effects on the bowel

Opioid-induced bowel dysfunction can result from both opioid
withdrawal and chronic opioid usage and manifests with features
of dysmotility (especially constipation) when pain is not the
dominant issue. The narcotic bowel syndrome may result from
chronic usage and is defined as chronic, worsening or frequently
occurring abdominal pain despite continued or escalating doses
of narcotics in addition to dysmotility.11 The opioid usage
induces a hyperalgesic effect. It may be becoming more preva-
 lent but it is often not recognised by clinicians.12 13 In addition
to being acknowledged to occur in patients with gastrointestinal
disease (functional or organic), it also occurs in patients with
no pre-existing gastrointestinal problems who take high doses
of the opioids for other painful conditions (eg, joint problems or
following surgery).

The components of therapy for narcotic bowel syndrome are
recognition of the disorder, a trusting therapeutic relationship
with the patient, replacement using neuropathic type pain drugs
and controlled reduction in the opioid dose.14 Specific drug
treatments have been tried for opioid-induced bowel dysfunc-
tion and narcotic bowel syndrome and include clonidine (to
reduce withdrawal symptoms) and peripheral mu opioid antag-
onists (naloxone, methylaltrexone, alvimopan).15–17

Opioids inhibit intestinal motility and so invalidate the tests
of small bowel motility. They may also increase the risk of line
infections in patients on long-term PN.

Survivors of cancer treatment may have bowel dysmotility
which may be due to chemotherapy or opioid medication. Their
management may require a wider MDT input.

Cyclizine is both an antihistamine and anticholinergic drug
which acts as a centrally acting anti-emetic. There are many
reports of it being taken for its euphoric effect, which is most
marked when taken intravenously. In addition to causing addic-
tive behaviour, it is of a low pH and so damages veins. It is not
recommended for long-term use, especially in patients receiving
PN.18

Other drugs such as anticholinergics, antidepressants,
calcium channel blockers, chronic laxative abuse or some

Figure 1 Progression of chronic small intestinal dysmotility.
chemotherapeutic drugs (eg, vincristine) may also cause reduced gut propulsion.

5.3 Functional gastrointestinal disorders
Many of the symptoms of small intestinal dysmotility are the same as for patients with other functional abdominal gastrointestinal disorders (eg, irritable bowel syndrome, functional dyspepsia, cyclical vomiting, functional bloating/distension, functional constipation/diarrhoea and centrally mediated disorders of gastrointestinal pain).19

The differentiation and overlap with these functional gastrointestinal disorders is difficult. They all may have genetic and psychosocial influences (early life trauma, life stresses, coping mechanisms, lack of social support, etc). In addition, bacterial flora, inflammation, visceral sensation and motility may all contribute to the symptoms. In irritable bowel syndrome there may be an overlap with enteric neuropathy as increased lymphocytes have been observed in the jejunal myenteric plexus.20 However, significant malnutrition is rarely a consequence of these disorders.

The treatment—as for all dysmotility problems—is to identify and treat the main symptom. If weight loss has occurred, then the same therapies as for intestinal dysmotility may be tried at the same time as nutritional support is given. Significant caution should be exercised, however, to avoid escalating to more invasive forms of nutrition support in patients with functional symptoms, especially in pain predominant presentations, in the absence of objective features of biochemical disturbance or those who have a high or normal BMI. Such escalation of invasive forms of nutrition support in patients with functional gut.29–31 The effects of undernutrition on gut motility and histological appearance are uncertain.

5.4 Psychological/psychiatric problems

5.4.1 Anorexia nervosa
The American Psychiatric Association (APA) Diagnostic and Statistical Manual of Mental Disorders (DSM- 5) stated in 2013 that to diagnose a person with anorexia nervosa they must display: (1) persistent restriction of energy intake leading to significantly low body weight (in context of what is minimally expected for age, sex, developmental trajectory and physical health); (2) either an intense fear of gaining weight or of becoming fat, or persistent behaviour that interferes with weight gain (even though significantly low weight); (3) disturbance in the way one’s body weight or shape is experienced, undue influence of body shape and weight on self-evaluation, or persistent lack of recognition of the seriousness of the current low body weight.21

However, the patients that present to gastroenterologists often do not have this typical presentation. Delayed gastric emptying, especially of solid, and delayed small and large bowel transit have been described in patients with anorexia nervosa.22–25 There is a report of a patient having a mega-duodenum and no propagating MMCs which both improved with an increased nutritional intake.26

5.4.2 Avoidant/restrictive eating
Some patients who have had psychosocial problems in the past may have disordered gut motility partly due to a disordered eating pattern, undernutrition and the drug treatments which they received.27

5.4.3 Psychiatric disorders and psychological distress
Other major psychiatric disorders can be encountered, sometimes masquerading as or confounding a dysmotility diagnosis.28

5.5 Effect of undernutrition on gut function
Malnutrition itself can impair gut function and cause malabsorption with mucosal atrophy, reduced gastric acid and pancreatic enzyme secretion and more bacterial colonisation of the upper gut.29–30 The effects of undernutrition on gut motility and histological appearance are uncertain.

5.6 Hypermobile Ehlers–Danlos syndrome (EDS) (joint hypermobility syndrome or Ehlers–Danlos syndrome hypermobility type)
Hypermobile Ehlers–Danlos syndrome (EDS) with its gastrointestinal associations is difficult to classify as most patients with EDS do not have a dysmotility of the small bowel, merely visceral hypersensitivity. Its symptoms may mimic dysmotility (neuropathy) and the presence of EDS can contribute to dysmotility, usually if postural tachycardia syndrome (PoTS) or other associated factors such as opiates are present. According to the 2017 classification, patients previously diagnosed with joint hypermobility syndrome and Ehlers–Danlos syndrome hypermobility type were reclassified as hypermobile Ehlers–Danlos syndrome (hEDS) if they met the strict criteria or Hypermobile Spectrum Disorders (HSD) if they had many but not all of the characteristics of hEDS.36 Patients with hEDS and HSD represent a third of patients referred to a tertiary neurogastroenterology clinic and these patients tend to be young, female with a poorer quality of life.37 HEDS/HSD is associated with a range of gut disorders (acid reflux, abdominal pain (especially is pain after eating or when any food arrives in the gut even from an enteral feed) and constipation).38 There is often autonomic dysregulation, particularly PoTS,39 chronic urinary retention due to a failure of the urethral sphincter to relax (Fowler’s
syndrome) and hypoglycaemia. Mast cell activation disorder is being increasingly reported (most commonly in those having PoTs). An increasing number of patients with joint hypermobility and gut dysmotility are being seen by nutrition support teams because of weight loss and malnutrition. This group seems especially sensitive to opioids and cyclizine, both of which can exacerbate all of their symptoms. It is currently unclear the degree to which the association of hEDS with gut symptoms encompasses specifically any greater degree of chronic small intestinal dysmotility, or whether there are any different treatment approaches to patients without hEDS who have the same functional symptoms. The same cautions therefore should apply when considering escalating invasiveness of nutrition support in this group as to that of functional gastrointestinal disorders in general, especially if there is a pain predominant presentation.

6.0 CLASSIFICATION OF CHRONIC SMALL INTESTINAL DYSMOTILITY

Three major histopathological entities are recognised: myopathies, neuropathies and mesenchymopathies, depending respectively on the predominant involvement of smooth muscle cells, enteric neurons or the interstitial cells of Cajal (ICC). Mesenchymopathies, which involve ICC, are the gut pacemakers, and are being recognised. The abnormalities described include a decreased ICC density, loss of processes and damaged intracellular cytoskeleton and organelles as revealed by immunohistochemical analysis and electron microscopy. It may be that abnormalities with the ICC are the primary event or may result from a neuropathy. In babies, immaturity may result in a delay in maturation of ICCs that can lead to the appearance of a reduction in these cells, so care needs to be taken with the histological diagnosis. As conditions specifically falling into this category are few, this document will only discuss conditions traditionally classified as a myopathy or neuropathy (figure 2, box 2).

It can be hard to determine if a condition is primarily a myopathy or neuropathy as some secondary conditions (eg, systemic sclerosis, vasculitis or amyloid) may appear as both. The end result of a neuropathy is often dysfunctional enteric muscle which occasionally can dilate, as in a myopathy. Overall, a neuropathy is more common than a myopathy in causing small bowel dysmotility but the reverse may be true in CIPO. The diagnosis can be difficult to define partly because the radiological, isotopic, manometric and histological diagnoses may be different.

6.1 Myopathies

Primary myopathies most commonly occur in children and young adults and are often familial (genetic). Myopathies often have multivisceral involvement and thus a relatively high mortality. They are characteristically associated with massive gut dilatation. Many predominantly affect the circular muscle (except hollow visceral myopathy). Primary myopathies are more common in children whereas secondary myopathies are more common in adults. Some causes will remain idiopathic.

6.1.1 Primary myopathies

Primary myopathies are due to abnormalities in enteric muscle (eg, hollow visceral myopathy or autoimmune myopathy). There has been interest in reduced immunostaining of alpha-isoactin in jejunal circular muscle, observed in one then 28 more patients (overall 24% of patients having full thickness jejunal biopsies). However, it is not clear if this is the primary pathology or secondary to other factors (eg, drugs, undernutrition or previous surgery) or a normal anatomical variant, and therefore not a specific finding. Precise information as to the location of the biopsy (jejunum or ileum) is essential for interpretation in this context.

Familial

Hollow visceral myopathy is the best known example of a familial visceral myopathy. It is a rare congenital disorder that usually presents in the first or second decade of life and, in addition to gross dilatation of the gastrointestinal tract (that often starts with a megaduodenum), there may also be associated dilatation of the urinary tract and associated frequent urinary tract infections. The disease has been reported to follow an autosomal dominant mode of inheritance. In infants other features may include malrotation, pyloric stenosis and bladder atony. There is a loss of enteric smooth muscle with vacuolar degeneration and fibrosis. The longitudinal muscle is predominantly affected. This may be due to the transformation of smooth muscle cells to collagen synthesising myofibroblasts. In one study of adult patients, four of six patients had urinary tract involvement with dilatation of the ureters and/or incomplete bladder emptying.

Acquired: autoimmune myopathy

A few cases only of a lymphocytic enteric leiomyositis involving the smooth muscle cells have been reported. Eosinophilic leiomyositis has also been reported, and in this context must prompt investigation for parasites, including dog hookworm.

Jejunal diverticulosis

Diverticula in the jejunum usually result from congenital abnormalities or dysmotility (eg, disordered high pressure bowel contractions) and may be associated with subclinical systemic sclerosis. Careful examination of any resected specimen should be carried out to examine the muscle and nerve layers of the bowel wall, both in the region of the diverticula and adjacent bowel.

6.1.2 Secondary myopathies

Secondary myopathies occur as part of a multi-system disease (eg, systemic sclerosis, amyloid, chronic irradiation damage or muscular diseases). The problems of pseudo-obstruction are often only clinically apparent towards the end of the illness when nutritional support may be needed.
**Classification of intestinal dysmotility**

**Myopathies**

*Primary*
- Familial
  - Hereditary myopathy (e.g., hollow visceral myopathy)
- Acquired
  - Autoimmune
  - Jejunal diverticulosis

*Secondary*
- Systemic sclerosis (and other connective tissue disorders [e.g., SLE])
- Amyloidosis
- Chronic irradiation damage
- Muscular diseases
  - Muscular dystrophies
  - Myofibrillar myopathies (e.g., desmin myopathy)
  - Hereditary inclusion body myopathies
  - Metabolic storage disorder

**Neuropathies**

*Primary (intrinsic)*
- Familial/congenital/developmental
  - Hirschsprung’s disease
  - Neuromyelomalacia
  - Mitochondrial disorders (MNGIE, DNA depletion, Alpers or Pearson’s syndromes)
  - Neuronal dysplasia
  - Infant colic (developmental)
- Autoimmune
  - Antineuronal antibodies
  - Ganglionosis
- Infective
  - Chagas’ disease
  - Herpes viruses (e.g., EBV, CMV, VZV) or
  - Polyoma viruses (JC virus)

*Secondary (extrinsic)*
- Generalised neurological disorders
  - Brainstem lesions
  - Spinal cord injury
  - Multiple sclerosis
  - Parkinson’s disease
  - Neurological effects of diabetes mellitus
  - Autonomic system degeneration
- Paraneoplastic syndromes (often with antineuronal antibodies especially anti Hu)
  - Small cell lung cancer
  - Carcinoid
  - Neuroblastoma
  - Thymoma
- Drugs/toxins
  - Vincristine, Adriamycin
  - Antidepressants, Ca channel blockers, anticholinergic drugs
  - Opioids
  - Clonidine
  - Isoniazid
- Other
  - MEN IIb
  - Porphyria (acute intermittent)
  - Fabry’s disease

In most cases an empirical working diagnosis of idiopathic dysmotility will be applied.

*Histology is generally inflammatory or degenerative.

**Systemic sclerosis (scleroderma) and other connective tissue disorders**

Most patients with systemic sclerosis do get gastrointestinal involvement particularly of the oesophagus. While the end result and main pathology is smooth muscle atrophy and gut wall fibrosis, it may start with microvasculature damage due to collagen deposits and inflammation which cause neural damage that progresses to muscle dysfunction and fibrosis. While systemic sclerosis patients with gross gastrointestinal involvement present in the terminal phase of the illness, this is not always the case and some present with gastrointestinal involvement early in the disease without cutaneous involvement and the disease may not progress for many years. The clinical outcome in elderly scleroderma patients is the poorest of all adult onset dysmotility patients. However, where indicated, long-term PN can offer a safe and effective means of nutritional support in patients with severe small bowel involvement. Other connective tissue and rheumatological disorders have been associated with dysmotility including systemic lupus erythematosus, rheumatoid arthritis and Still’s disease.

**Amyloidosis**

The primary type distribution may be associated with gut involvement. The most common underlying diagnosis is myeloma (often producing lambda chains). While classified as a myopathy, it can also cause neurological damage and, like systemic sclerosis, may cause both a myopathy and a neuropathy. The rectum can be spared so duodenal sampling should also be considered. Any full thickness biopsy for motility investigation should also be examined for amyloidosis. Genetic testing is now readily available for the hereditary type of amyloidosis, which may present more commonly in men and with peripheral neuropathy and cardiac as well as gut involvement.

**Chronic irradiation damage**

This usually occurs after pelvic irradiation for gynaecological cancers or genitourinary cancers. The sigmoid and terminal ileal areas are often most involved. However, the whole small bowel can be involved giving rise to a pseudo-obstruction picture or as discrete areas of strictures. Surgery is very difficult and has a high risk of an enterocutaneous fistula(s) being created. Gut permeability, secretion, motility and blood supply can all be affected, giving rise to any or all of the following: malabsorption, protein losing enteropathy, diarrhoea, perforation/fistulas, bleeding and obstruction. These problems can all occur months or years after more than 45 Gy irradiation has been given. The irradiation is more likely to give problems if a patient is already malnourished, has diabetes mellitus, hypertension or a vasculitis.

**Muscular diseases**

Myopathies may occur associated with congenital muscular disorders (muscular dystrophies, myofibrillar myopathies (e.g., desmin myopathy), hereditary inclusion body myopathies), but the muscle weakness (and often cardiac problems) dominates the clinical picture although gastrointestinal problems if sought are common. Metabolic storage disorders can have a myopathic process and occasionally can be treated with specific enzyme replacement therapy.

**6.2 Neuropathies**

An enteric neuropathy may occur as a primary pathology (congenital, autoimmune or infective), secondary to a generalised neurological disorder, paraneoplastic process, metabolic disorder (e.g,
diabetes) or drugs/toxins or as a developmental abnormality. Visceral neuropathy is less well reported in the literature than visceral myopathy. Visceral neuropathy is commonly acquired in adulthood or in old age and is associated with a high morbidity usually due to factors other than the neuropathy. Luminal dilatation is rarely seen except at the end stage of the disease.

Two forms of pathology are found—either enteric neural degeneration (in the absence of inflammation) or inflammatory. Degenerative neuropathies can result from mitochondrial dysfunction and the pathological findings include neuronal swelling, intranuclear inclusions, axonal degeneration and hypoganglionosis. The aetiology of many degenerative enteric neuropathies will remain idiopathic. Inflammatory neuropathies may include both plexitis and neuritis and can be lymphocytic or less commonly eosinophilic, the former promoting autoimmunity screening, the latter investigation for parasites.

6.2.1 Primary neuropathy
The enteric neuropathies can affect both the submucosal and myenteric plexuses but the myenteric plexus is predominantly affected. The term ‘visceral neuropathy’ is used for primary intrinsic enteric nervous damage.

Congenital/familial/developmental
Familial visceral neuropathies include Hirschsprung’s disease, mitochondrial cytopathies and Von Recklinghausen’s disease. Hirschprung’s disease can affect any part of the gut.

Mitochondrial disorders are relatively common if specifically sought. One study showed 19% of 80 adult patients labelled as CIPO to have this. Mitochondrial cytopathies such as mitochondrial neurogastrointestinal encephalomyopathy (MNGIE) (the most common) (also referred to as thymidine phosphorylase deficiency) is an autosomal recessive disorder characterised by severe gastrointestinal dysmotility (including bacterial overgrowth and lactic acidosis), cachexia and neurological problems including leukoencephalopathy (96%), polyneuropathy (96%), ophthalmoplegia (91%) (with ptosis) and hearing loss (35%).

The disease is caused by mutations in the thymidine phosphorylase (TP) gene. Gastrointestinal dysmotility is the most prominent manifestation, with recurrent diarrhoea and symptoms of obstruction. Patients with MNGIE present between the first and third decade (mean age 18 years) and usually have a very reduced life expectancy and tend to die in early adulthood (mean 37.6 years; range 26–58 years). It, like all mitochondrial defects, can be tested for by plasma and urine thymidine deoxyuridine, white cell thymine phosphorylase, the Tymp gene, MR brain scanning and muscle biopsy.

Autoimmune
Auto-antibodies directed at enteric neurons, usually neuronal ion channels (eg, voltage-gated potassium channels) can occur as a paraneoplastic phenomenon, when the anti-neuronal nuclear antibody (ANNA-1 or anti-Hu) is most commonly found, and antineuronal antibodies can occur in non-paraneoplastic motility disorders. Other auto-antibodies associated with dysmotility include acetyl cholinesterase receptor antibody (AChR) (ganglionnic, nicotinic and M3 type), antibodies against the voltage-gated potassium channel-complex (VGKC-complex), voltage-gated calcium channel antibodies (VGCC), smooth muscle and gonadotropin-releasing hormone (GnRH). The pathogenetic role of auto-antibodies is, however, currently unclear. Coeliac disease has also been implicated in some cases of dysmotility.

Infective
Both herpes (Epstein–Barr virus and cytomegalovirus) and polyoma viruses (John Cunningham (JC) virus) have had their DNA isolated in the myenteric plexuses of some patients with visceral neuropathy. They may be causative agents rather than innocent bystanders but this has yet to be proven. Chagas’ disease (South American trypanosomiasis) typically causes a megaesophagus and megacolon. Chagas’ enteropathy is common and gives rise to dyspepsia, intestinal pseudo-obstruction with bacterial overgrowth.

Lyme disease and botulism have also been reported as reversible causes of dysmotility.

6.2.2 Secondary neuropathy
The neuropathic process may affect the nerves external (extrinsic neuropathy) to the gut and so indirectly affect gut motility or may be part of a generalised illness. Indeed, most cases of visceral neuropathy are part of a generalised neurological disorder rather than a primary neuronal disorder of the gastrointestinal tract.

Generalised neurological disorders
Disorders of the parasympathetic or sympathetic nerves that innervate the gut (including autonomic system degeneration and the neurological effects of diabetes mellitus (most common) and other endocrine or metabolic disorders) can indirectly cause gut dysmotility. Brainstem lesions, spinal cord injury, multiple sclerosis, Parkinson’s disease (basal ganglia calcification) and leukencephalopathy can all affect gut motility. A lymphocytic leiomysitis and myenteric ganglionitis have been described in the ileum of children with cystic fibrosis and distal ileal obstruction. Myotonic dystrophy, multiple sclerosis, Parkinson’s disease and porphyria may all be associated with an enteric neuropathy.

Paraneoplastic
Small cell lung cancer, carcinoid tumours, neuroblastoma and thymoma with anti-neuronal nuclear antibodies have all been described to cause an enteral neuropathy. There is often a dense inflammatory infiltrate of lymphocytes and plasma cells affecting both plexuses but mainly the myenteric (myenteric ganglionitis). Anti-Hu neuronal antibody is characteristic. Removal of a thymoma may result in resolution of the dysmotility and the patient gaining weight. CRMP 5 (CV2) and AchR auto-antibodies have also been associated with paraneoplastic dysmotility and should prompt a careful search for occult malignancy.

Drugs/toxins
Vincristine is directly neurotoxic and causes a visceral neuropathy. Anticholinergics (eg, phentolamines and tricyclic antidepressants) have been associated with severe dysmotility. A case series of 102 life-threatening episodes of clozapine-induced gastrointestinal dysmotility episodes were collated with some evidence for dose dependence. A number of other drugs have been associated with severe dysmotility which, in most cases, improves with stopping the drug or reducing the dose; these include baclofen, buserelin, clonidine, fludarabine, phenytoin and verapamil. Lead poisoning can be a rare reversible cause.

6.3 Idiopathic
In the majority of patients and in most centres, the precise aetiology of chronic severe dysmotility is not characterised histopathologically and remains based on the clinical presentation, physiological testing and exclusion of obstructive and mucosal disease. This in part reflects a low uptake of full thickness
biopsies outside of the context of stoma formation or other surgical intervention. In selected populations of PN dependent patients with dysmotility, high rates of full thickness biopsies were associated with high rates of neuromuscular abnormalities of which two-thirds were primary and one-third were secondary causes, although not all biopsies yielded a diagnosis. The threshold and acceptability of full thickness biopsy testing, especially as most will not lead to a change in management, has not currently achieved consensus. There also remain some unresolved issues for gastrointestinal neuromuscular pathology standardisation and interpretation.59 For the near future, therefore, it is likely that the aetiology in the majority of dysmotility patients will remain idiopathic.

7.0 PHYSIOLOGICAL CONSEQUENCES OF SEVERE SMALL INTESTINAL DYSMOTILITY

7.1 Impairment of coordinated gut contractions and the migrating myoelectric complex (MMC)
If the MMC is impaired, then the small bowel will not be cleared of debris predisposing to gut stasis and bacterial overgrowth. With enteric neuropathies, gut coordination is disrupted and the presence of chyme in the small bowel can cause severe painful non-propulsive large contractions. This is one of the causes of abdominal pain shortly after eating.

7.2 Gut stasis
The failure of forward propulsion may also cause constipation and this is often the first symptom. Gut stasis results in abdominal distension and, if much fluid accumulates (oral intake and normal gut secretions), it may produce a large volume vomit. The vomit may be faeculent and contain food debris from several days previously.

7.3 Bacterial overgrowth and malabsorption
The combination of a dilated gut with reduced propulsion and ineffective MMC allows anaerobic bacteria to proliferate in stagnant loops of bowel. This bacterial overgrowth results in bile salts being deconjugated, less effective secondary bile acids (eg, lithocolic acid) being made and pancreatic enzyme degradation occurs so that steatorrhoea and malnutrition may occur. Associated with steatorrhoea is malabsorption of the fat soluble vitamins (A, D, E and K) with deficiency symptoms (night blindness, poor colour vision, dry flaky skin and ataxia). Vitamin K1 may be malabsorbed but both folic acid and vitamin K can be manufactured by the bacteria and so may give rise to high serum levels.

Occasionally the bacteria can manufacture D-lactic acid (normally L-isomer) giving rise to D-lactic acidosis (high anion gap acidosis) and other bacteria can manufacture ammonia which may appear in high levels in the blood.

Small intestinal bacterial overgrowth (SIBO) is when excess micro-organisms are present in the small intestine and lead to a malabsorption syndrome with occasionally a protein losing enteropathy. Subtotal villous atrophy may be found on histology. There are several endogenous mechanisms for preventing bacterial overgrowth: gastric acid secretion, intestinal motility, intact ileo-caecal valve, intestinal immunoglobulin secretion and bacteriostatic properties of pancreatic and biliary secretions. The aetiology of SIBO is usually complex, associated with disorders of these mechanisms. In some patients more than one factor may be involved.

There is currently no gold standard for the diagnosis of SIBO and the commonly available methodologies (the culture of jejunal aspirates and a variety of breath tests) are limited by large variations in their performance and interpretation.78

7.4 Problems of undernutrition
Patients who rapidly lose more than 10% of their body weight frequently have demonstrable physiological changes which include skeletal and cardiac muscle weakness, poor concentration and mental function including memory, prolonged sleeping, reduced sexual function, a low body temperature and a propensity to develop infections which are potentially severe.79 In the gut, malabsorption with mucosal atrophy, reduced gastric acid and pancreatic enzyme secretion and more bacterial colonisation of the upper gut can occur although the experimental backing for this is inferred from studies in patients with anorexia nervosa.22–26

8.0 CLINICAL FEATURES OF CHRONIC SMALL INTESTINAL DYSMOTILITY AND MANAGEMENT PLAN

8.1 History, examination and blood tests
The features of patients with a myopathy are summarised in Table 1. The clinical history and examination should determine if there are associated systemic neuromuscular, connective tissue or endocrine diseases (muscular diseases, neurological disease, storage diseases, systemic sclerosis, diabetes mellitus, irradiation, etc) and thus if the myopathy or neuropathy is a primary or secondary disorder. Exploring the family history will detect some congenital diseases, as will asking about foreign travel (Chagas’ disease). Examination especially includes the neuromuscular system and testing for joint hypermobility. Autonomic neuropathy should be considered if orthostatic, pupillary or sudomotor (sweating) dysfunction accompanies dysmotility. Simple clinical bedside assessment of orthostatic pulse rate change (lying to standing) may identify PoTs (box 3).52

Symptoms need to be listed in order of importance to the patient, and a record made of all the drugs currently taken or that have been taken for long periods (especially opioids and cyclizine).

A basic nutritional assessment will include measuring the patient’s height and weight and asking their usual weight in health and their weight change over the last 2 weeks, 3 and 6 months. From these their BMI and percentage weight loss can

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Table 1 Symptoms, medical history and medication of 28 patients with an enteric myopathy41

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>Duration, median (range)</th>
<th>Abdominal pain</th>
<th>Distension</th>
<th>Nausea/vomiting</th>
<th>Constipation</th>
<th>Diarrhoea</th>
<th>Weight loss</th>
<th>Medical history</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdominal pain</td>
<td>14 (4–33)</td>
<td>28 (100%)</td>
<td>23 (82%)</td>
<td>22 (79%)</td>
<td>17 (61%)</td>
<td>6 (21%)</td>
<td>10 (36%)</td>
<td>9 (32%)</td>
</tr>
<tr>
<td>Distension</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Previous laparotomies 8 (29%) (median 2 (range 1–3))</td>
</tr>
<tr>
<td>Nausea/vomiting</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>8 (29%)</td>
</tr>
<tr>
<td>Constipation</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6 (21%)</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6 (21%)</td>
</tr>
<tr>
<td>Weight loss</td>
<td>10 (36%) (5 given PN)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Table 1 Symptons, medical history and medication of 28 patients with an enteric myopathy.
be calculated. In addition they are asked about recent changes in their food intake.

Tests to help make the diagnosis are summarised in box 4. Blood tests will include routine blood count, renal (including potassium and magnesium), liver, bone chemistry, thyroid function, glucose, myeloma screen, anti-potassium and magnesium), liver, bone chemistry, thyroid function, glucose, myeloma screen, anti-

8. Try to establish a clinical diagnosis (or probable one). Perform physiologic assessment of the parts of the gastrointestinal (GI) tract that may be involved. These are done when nutritional status is near normal and the patient is off drugs likely to affect GI motility. Consider full thickness jejunal biopsy

9. Consider surgical options

10. Regular review and reconsider diagnosis as the clinical situation changes. Treat the predominant symptom/problem

Specific investigations are performed once organic obstruction, drug effects and eating disorders have been excluded. Physiological/histological tests are done after severe malnutrition has been treated.

1. Screen for hypothyroidism, coeliac disease and diabetes

2. Chest X-ray (or CT/PE)* for thymoma or other neoplastic conditions (eg, small cell carcinoma of lung)

3. Antibodies for scleroderma (anti-centromere, anti Sc170, anti M3R) and other connective tissue disorders (ANA, ANCA, anti DNA, anti SMA)

4. Antibodies that may be associated with paraneoplastic diseases (mainly small cell carcinoma and thymoma). These may include type 1 anti-neuronal nuclear antibody (ANNA-1 'anti Hu'), anti-collapsin response mediator protein 5 (anti CRMP-5 also known as anti CV2), ganglion acetyl cholinesterase receptor antibody (AChR antibody) especially if autonomic dysfunction, and anti-voltage gated potassium channel (VGKC)-complex antibodies.

5. Test for mitochondrial disorders with plasma and urine thymidine and deoxyuridine, WBC thymidine phosphorylase. If there is a high suspicion then test for the TYMP gene and also screen for related diseases (eg, 'MELAS' (mitochondrial encephalopathy with lactic acidosis and stroke-like episodes) with the m.3243A>G mutation). Muscle biopsy and sequencing of mitochondrial genome may be considered.

If none of these are positive consider full thickness jejunal biopsy.

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8.2 Radiological tests

The diagnosis is usually first suspected after plain abdominal radiographs have shown a dilated small and large bowel. Once suspected, investigations aim to confirm that there is impaired transit of luminal contents, to identify the region of the bowel affected, ideally to identify the propulsive abnormality and to show a specific pathology. Investigations help establish the presence of intestinal pseudo-obstruction and may delineate an underlying cause. In practice the diagnosis is often presumed after several laparotomies have excluded a physical obstruction, although CT/barium follow-through/MR enterography

Guidelines

### Box 3 Management plan for small intestinal dysmotility

1. Determine and order the primary symptoms
2. Exclude mechanical obstruction (CT abdomen with oral contrast)
3. Evaluate other contributing factors: drug therapy (eg, opioids, cyclizine and anticholinergics), psychosocial (may need formal psychological/psychiatric assessment) and quality of life issues
4. Nutritional assessment (BMI, percentage weight loss and other anthropometric tests)
5. Start nutritional treatment (consider/treat refeeding risks)
6. Perform tests to help establish aetiology (box 4) and consider tests of autonomic function
7. Therapeutic plan/objectives of care to address patient’s symptoms, nutritional status, psychosocial issues and quality of life
8. Try to establish a clinical diagnosis (or probable one). Perform physiologic assessment of the parts of the gastrointestinal (GI) tract that may be involved. These are done when nutritional status is near normal and the patient is off drugs likely to affect GI motility. Consider full thickness jejunal biopsy
9. Consider surgical options
10. Regular review and reconsider diagnosis as the clinical situation changes. Treat the predominant symptom/problem

### Box 4 Non-physiological tests to determine the aetiology of chronic small intestinal dysmotility

- **Screen for hypothyroidism, coeliac disease and diabetes**
- **Chest X-ray (or CT/PE) for thymoma or other neoplastic conditions (eg, small cell carcinoma of lung)**
- **Antibodies for scleroderma (anti-centromere, anti Sc170, anti M3R) and other connective tissue disorders (ANA, ANCA, anti DNA, anti SMA)**
- **Antibodies that may be associated with paraneoplastic diseases (mainly small cell carcinoma and thymoma). These may include type 1 anti-neuronal nuclear antibody (ANNA-1 'anti Hu'), anti-collapsin response mediator protein 5 (anti CRMP-5 also known as anti CV2), ganglion acetyl cholinesterase receptor antibody (AChR antibody) especially if autonomic dysfunction,** and anti-voltage gated potassium channel (VGKC)-complex antibodies.
- **Test for mitochondrial disorders with plasma and urine thymidine and deoxyuridine, WBC thymidine phosphorylase. If there is a high suspicion then test for the TYMP gene and also screen for related diseases (eg, 'MELAS' (mitochondrial encephalopathy with lactic acidosis and stroke-like episodes) with the m.3243A>G mutation). Muscle biopsy and sequencing of mitochondrial genome may be considered.**

If none of these are positive consider full thickness jejunal biopsy.

8.1.2 Neuropathy

Many of the features are the same as for a myopathy, particularly with severe abdominal pain after food; however, abdominal distension is often absent and the plain abdominal radiograph may appear normal.

8.2 Radiological tests

The diagnosis is usually first suspected after plain abdominal radiographs have shown a dilated small and large bowel. Once suspected, investigations aim to confirm that there is impaired transit of luminal contents, to identify the region of the bowel affected, ideally to identify the propulsive abnormality and to show a specific pathology. Investigations help establish the presence of intestinal pseudo-obstruction and may delineate an underlying cause. In practice the diagnosis is often presumed after several laparotomies have excluded a physical obstruction, although CT/barium follow-through/MR enterography


2083

Guidelines

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If none of these are positive consider full thickness jejunal biopsy.
excluding a transition point in a diffusely distended small bowel suggests CIP0 and may prevent unnecessary laparotomy. CT may also help distinguish severe dysmotility from functional bloating due to abdomino-phrenic dysinergia. Dynamic MRI of the small bowel is becoming increasingly helpful, though is less established. MRI brain can be helpful in the diagnosis of MNGIE.39

The measurement of whole gut time can be measured by serial X-rays of ingested radio-opaque markers (small lengths of barium-impregnated polyvinyl tubing).

Small bowel transit using a barium follow-through examination will usually give some indication of accelerated or delayed transit and a diluted duodenal loop (megaduodenum) may be one of the earliest signs of visceral myopathy. In addition, in HVM there may be oesophageal aperistalsis and variable dilatation of the small and large bowel.

8.3 Radioisotopic investigations

Gastric emptying can be measured using gamma scintigraphy to obtain serial images of labelled solid (scrambled eggs, liver or pancake), semi-solid (thick soup) or liquid (orange juice) meals in the stomach. Gastric emptying measurements may be helpful in determining whether the stomach is involved in a generalised disorder of propulsion or a localised one (eg, Chagas’ disease). These isotopic meals can be extended to measuring the transit of the meal through the small bowel and, if the isotope has a long half-life, oro-caecal and colonic transit may be determined. Liquid meals may not clearly demonstrate an abnormality.

8.4 Endoscopic tests

Jejunal aspirate for bacterial overgrowth is infrequently performed but usually by endoscopy (or fluoroscopy with jejunal intubation). A clinically significant overgrowth is when counts exceed 10^9/mL (usual is less than 10^9/mL). Common species include Bacteroides, Enterococcus and Lactobacillus. However, most of the bacteria likely to be relevant in causing symptoms cannot be cultured. Endoscopy also has a role in mucosal sampling and palliative venting. Capsule endoscopy examination can sometimes give transit information but is rarely used due to the risk of the capsule being retained and some regard it as contraindicated.

8.5 Non-invasive investigations of gut transit

Oro-caecal transit can be measured using the rise in breath hydrogen due to the degradation of ingested polysaccharides (eg, lactulose). In health this is caused by caecal bacteria fermenting the ingested agent, but this is unhelpful if there is propulsive failure as small bowel bacterial overgrowth is common and is not a recommended test for this application. It is also unhelpful following a significant small bowel resection or if there is an enteric fistula.

Breath tests to diagnose bacterial overgrowth may be misleading and produce false negative results compared with culture of small bowel aspirate. This has in part been due to broad variations in how these tests are performed and interpreted. Recent work in the UK and USA has taken place to develop evidence-based consensus guidelines for breath testing in terms of dose of substrate (75 g glucose, 10 g lactulose) and cut-off values. In addition, while hydrogen only breath testing was previously used, modern testing protocols have incorporated the measurement of methane. Increased intestinal methane levels have been associated with delayed small bowel transit as measured by scintigraphy and therefore should be measured in patients with suspected small bowel dysmotility to improve the test’s utility. Other tests that may indicate bacterial overgrowth include raised urinary indicans, blood D-lactate or alcohol levels.

8.6 Manometry

Intraluminal pressure sensors incorporated into a catheter can detect the patterns of contractile events. For the diagnosis of pseudo-obstruction, the logical investigation is small bowel manometry. Small bowel motor activity was initially studied using multi-lumen perfused tube systems, with a pump and strain gauge transducer external to the patient. This gave information on motor activity in the antrum and proximal duodenum, and detected abnormal motility and some types of pseudo-obstruction. This technique required the patient to remain in a laboratory attached to a machine for more than 6 hours and was not good at recording fasting, and postprandial activity.

An alternative is the wireless motility capsule (WMC) which is an ambulatory, minimally invasive diagnostic modality that allows continuous assessment of intraluminal pH, temperature and pressure during its transit through the gastrointestinal tract. The technology allows for both measurement of transit times in multiple regions of the upper and lower gastrointestinal tract, as well as pressure profiles in the antro-duodenum. The standardised equipment and procedures in WMC allow the comparisons of data across multicentres. The role of the technology has been best established in the evaluation of a large number of healthy volunteers and in patients with suspected gastroparesis and suspected chronic constipation. The worry with this technique in these patients is that the capsule may remain in the bowel and not be passed. However, a study in patients with Crohn’s disease has shown that the same precautions used when considering patients for capsule endoscopy (ie, clinical and radiological assessment and use of a patency capsule) can mitigate much of the risk in potential WMC patients.

Twenty-four hour ambulatory jejunal manometry uses a catheter with built-in miniature strain gauge transducers and records data to a solid-state digital recorder. The digital encoding of pressure data simplifies the analysis of continuous 24-hour recordings by computer software. This technique has proved useful in several conditions including pseudo-obstruction. During nocturnal sleep, normal stereotypic MMC activity is clearly evident and in some patients with pseudo-obstruction the abnormal contractile activity of the small bowel results in distortion of the fasting MMC pattern. Manometry of patients with pseudo-obstruction can be difficult in the later stages of disease because the peristaltic activity required to propel a manometry catheter into position in the proximal jejunum is lacking, and endoscopic assistance may be needed. A patient with a neuropathy may have a normal diameter gut but it may be hyperactive with many uncoordinated and often strong contractions (bursts).

The effect of any drugs the patient may be taking must be taken into account in interpreting any results (eg, opioids, anti-cholinergics and cyclizine). Manometry does not always produce results that are clinically helpful.

8.7 Pathology and histology

The biochemical examinations recommended are listed in box 4. Adequate biopsy material is not often available and few laboratories have an experienced gastrointestinal neuropathologist. There also remain some additional pitfalls for collecting and analysing samples including sampling error, effects of bowel handling, sparsity of normal data and specificity. Good
Box 5 Drug therapies for intestinal dysmotility

Laxatives (after adequate fluid in diet)

- Osmotic
  - macrogols (PEG), lactulose, magnesium salts
  - Stimulant
  - anthraquinone group (senna and dantron)
  - bisacodyl, sodium picosulfate, docusate sodium
  - phosphate enema, glycerol suppository
  - parasympathomimetics: bethanechol, neostigmine, pyridostigmine
  - 5HT, receptor agonists: prucalopride
  - Bulk forming
  - unprocessed wheat bran, methylcellulose, ispaghula and sterculia
  - Faecal softeners/lubricants
  - liquid paraffin, arachis oil (ground nut oil, peanut oil) enemas
  - Peripheral opioid-receptor antagonists
  - methylaltrexone
  - Secretagogues
  - Linaclotide, Tenapanor*

Antispasmodics

- Antimuscarinics
  - tertiary amine: dicycloverine hydrochloride
  - quaternary ammonium compounds: propantheline bromide, hyoscyne butylbromide
  - Direct smooth muscle relaxant
  - alverine, mebeverine and peppermint oil

Prokinetics

- Dopamine receptor 2 antagonists
  - metoclopramide, domperidone
  - Macrolides
  - erythromycin

Antidiarrhoeal drugs

- Opioid agonists
  - loperamide, codeine phosphate, diphenoxylate
  - 5-HT3 antagonist
  - Ondansetron

Anti-emetics

- D2 receptor antagonists (see above)
  - cyclizine
  - ondansetron
  - ramosetron*

Analgesics

- tricyclic antidepressant (low dose) amitriptyline
  - selective serotonin reuptake inhibitor
  - serotonin and norepinephrine reuptake inhibitor (duloxetine)
  - gabapentin
  - pregabalin
  - antispasmodic drugs
  - opioids (low dose)

Antibiotics for bacterial overgrowth

- amoxycillin-claevulinc acid
  - ciprofloxacin
  - metronidazole, tinidazole
  - cephalosporin
  - tetracycline, doxycycline
  - non-absorbable antibiotics: rifaxamin, neomycin

*Not licensed in the UK.

histological samples are needed to make a firm diagnosis. Close liaison between the surgeon and pathology laboratory is crucial so that a full thickness specimen of bowel is immediately processed. The samples should be divided. In an ideal situation some is snap-frozen in liquid nitrogen and the main portion fixed for routine histology and electron microscopy. In practice the latter two may be the best option. The immediate processing of samples is important if a detailed examination of the nerves, ganglia and muscle tissue is to be carried out. Diagnosis of a neuropathy may be difficult in conventionally orientated and stained sections of gut, and whole mount plexus assessment is a research tool. The most important element is to ensure that enough sections and material is examined in a centre with experience of dysmotility.

A full thickness jejunal biopsy is usually taken laparoscopically and is often helpful diagnostically, but the procedure in centres without much experience can be unhelpful and have a significant risk. Published data from centres with expertise suggest a median operating time of 30 min, conversion rate to open operation 2% and length of stay 1 day with an 8% readmission rate for obstructive symptoms.102 In myopathies the diagnosis may be established. A neuropathy in general shows either degenerative changes or inflammation.

All full thickness biopsies should be stained with Congo red stains to look for the presence of amyloid. As full thickness jejunal biopsies may not change the clinical management and are associated with risks, they are often performed when a laparotomy does not find an organic cause of obstruction or when the patient happens to be undergoing surgery for another reason (eg, a jejunal tube placement).103 MNGIE can be diagnosed with a skeletal muscle biopsy in addition to the blood and genetic testing.

9.0 TREATMENTS

The drug treatments for intestinal dysmotility are shown in box 5. Most of the drugs are commonly used to treat milder forms of the symptoms. Treatment is occasionally directed at the underlying condition but more often is targeted at a specific symptom.

9.1 Underlying condition

An underlying disease may need to be treated (eg, connective tissue disorder, enteric myositis, neoplastic disease or myotonic dystrophy). Diabetic control should be very good and may necessitate an insulin pump. Electrolyte, mineral or endocrine abnormalities should be prevented and treated when detected.

Immunosuppressive treatment has a small evidence base restricted to case series or reports. Prednisolone and ciclosporin have been reported to be of particular benefit in autoimmune myopathy.104 There is a case report of an improvement with initially prednisolone 1 mg/kg and azathioprine 2 mg/kg/day, then subsequently the prednisolone was replaced with budesonide 9 mg/day.52

There must always be awareness that organic obstruction can be missed as a diagnosis and, if a prokinetic drug105 makes pain worse, then an organic obstruction must be considered. Similarly, a successful trial of a low fibre or liquid diet suggests an organic obstruction.

Some metabolic storage disorders can be treated with specific enzyme replacement therapy.51

9.2 Specific drug treatments of symptoms

No treatment is ideal, and even though some help to correct physiological abnormalities, they may not affect the patient’s symptoms (boxes 3 and 5).

Drug therapy106 can be difficult and often drugs with conflicting actions are used (prokinetic for constipation and...
anticholinergic for colicky pain). Essentially, the drug therapy is targeted at the symptom perceived as most important by the patient.

Prokinetic treatments are used to try and improve the dysmotility and can return some of the measured abnormalities towards normal. They may especially help with vomiting and constipation. Prokinetic drugs are generally not used after a bowel anastomosis. Some of the previously used prokinetic drugs have been withdrawn or can only be used with extreme caution. They include domperidone and metoclopramide (D₂ dopamine receptor antagonists) which stimulate gastric emptying and small intestinal transit, and enhance the strength of oesophageal sphincter contraction. Metoclopramide also increases the release of acetylcholine from some enteric nerves. Domperidone is a selective antagonist of peripheral D₂ dopamine receptors, which does not have the acetylcholine-like effect of metoclopramide. National Patient Safety Agency (NPSA) alerts have been issued for domperidone highlighting problems with prolonged QTc, therefore long-term use should be subject to QTc monitoring. The extrapyramidal side effects of metoclopramide (especially in children) and the potentially irreversible tardive dyskinesia in elderly subjects, together with no evidence of consistent benefit in gastroparesis, caused the European Medicines Agency’s Committee to recommend that metoclopramide is not used in the long term. Cisapride, a 5-HT₄ agonist, enhances acetylcholine release in the myenteric plexus without having anti-dopaminergic effects and may have been of particular benefit if MMCs were present on small intestinal manometry. In a 6-week double-blind, placebo-controlled trial in 26 patients, cisapride helped abdominal pain, improved solid gastric emptying and the MMC. Unfortunately, due to an increased risk of fatal cardiac arrhythmias (probably relating to a prolonged QT interval) in patients taking other medications or suffering from underlying conditions known to increase the risk of cardiac arrhythmias, cisapride was withdrawn. Tegaserod, a 5-HT₄ receptor partial agonist, increased stool frequency in irritable bowel syndrome and improved the symptoms in functional dyspepsia, but was withdrawn due to an increased risk of heart attacks or strokes.

Prucalopride, a high affinity selective 5HT₄ receptor agonist, has been used for constipation and appears not to have the cardiac risks of cisapride or tegaserod as it does not affect the QT interval. This is by having no significant action on the 5-HT₁D and on the cardiac human ether-a-go-go K⁺ channels. Erythromycin, a motilin agonist, is potentially useful if there are absent or impaired antroduodenal migrating complexes but is subject to tachyphylaxis. Doses of 900 mg/day have been recommended. Azithromycin may be more effective for small bowel dysmotility. A somatostatin analogue (octreotide), given by a relatively painful subcutaneous injection, may be dramatically beneficial, especially in systemic sclerosis when other treatments have failed. It can improve vomiting and pain, partly because octreotide (in normal subjects) reduces the perception of volume distension due to inhibition of sensory afferent pathways. Octreotide may cause low amplitude MMCs to return. Octreotide may have a beneficial effect when erythromycin has been unsuccessful; its effect (50–100 μg once or twice a day) is apparent within 48 hours and is maintained for more than 2 years. It may be more effective when combined with erythromycin.

The parasympathomimetics betahexol, distigmine, neostigmine and pyridostigmine enhance parasympathetic activity in the gut and increase intestinal motility. They are rarely used because of both their gastrointestinal and cardiovascular side effects (diarrhoea and severe bradycardia). Pyridostigmine has, however, been shown to help refractory constipation (including in diabetes) and was well tolerated using a stepped dosing regimen.

Naloxone 1.6 mg given subcutaneously each day or methylaltrexone given subcutaneously on alternate days may be beneficial in blocking dysmotility effects of opioids or in improving motility through blocking endogenous opioids.

### 9.2.1 Constipation

Constipation may be a problem in early stages, but is rarely present when IF occurs. In the early stages of these diseases, constipation may be managed by diet ensuring that it includes an adequate intake of fibre and fluid. Bulk forming laxatives such as unprocessed wheat bran (or oat bran) taken with food or fruit juice are effective and methylcellulose (which is also a faecal softener), ispaghula, and sterculia are useful in patients who cannot tolerate bran.

**Osmotic laxatives** (macrogols (polyethylene glycol), lactulose or magnesium salts) increase the amount of water in the large bowel, either by drawing fluid from the body into the bowel or by retaining the fluid that was administered. Macrogols are inert polymers of ethylene glycol (PEG) which sequester fluid in the bowel. Lactulose is a semi-synthetic disaccharide which is not absorbed from the gastrointestinal tract. It produces osmotic diarrhoea of low pH and prevents the proliferation of ammonia-producing organisms. Magnesium salts are useful where rapid bowel evacuation is required. Sodium salts should be avoided as they may give rise to sodium and water retention.

If there is an inadequate response to an osmotic laxative, a **stimulant laxative** can be added. Stimulant laxatives increase intestinal motility and often cause abdominal cramp; they should be avoided in intestinal obstruction. Excessive use of stimulant laxatives can cause diarrhoea and hypokalaemia. The anthraquinone laxatives (senna, dantron, cascaria) are converted in the intestine to active sennosides, which may function by stimulating the myenteric plexus in the colon and also by inhibiting colonic water absorption. Their principal effect is in the descending and sigmoid colon. Their effect is largely local and depends on sufficient intestinal motility to present them to the colon for bacterial degradation to their active form. Sennosides, with prolonged use, had been thought to damage the intestine muscle and/or myenteric neurons but there is no clinical or animal evidence to support this. Poorly absorbed diphenylmethane derivatives (bisacodyl, phenolphthalein, sodium picosulfate) stimulate sensory nerves in the proximal colon and increase sodium and water movement into the colonic lumen. Castor oil can have a place with its principal effect on small bowel fluid secretion. Docusate sodium probably acts both as a stimulant and also as a softening agent.

Dantron, cascaria and castor oil are rarely used, dantron because of potential carcinogenicity.

5HT₄ receptor agonists (prucalopride) are selective serotonin 5HT₄ receptor agonists with prokinetic properties. Prucalopride is licensed for the treatment of chronic constipation in women when other laxatives have failed to provide an adequate response. Headache and gastrointestinal symptoms (including abdominal pain, nausea and diarrhoea) are the most frequent but rare side effects. The side effects generally occur at the start of treatment and are usually transient. It has the potential to be a useful prokinetic drug now that cisapride and tegaserod have largely been withdrawn. Linaclotide (a 14-amino acid peptide) which acts in the intestinal lumen on guanylate cyclase-C (GC-C)
so generating cyclic guanosine monophosphate (cGMP), which stimulates chloride secretion, resulting in increased luminal fluid secretion and an acceleration of intestinal transit. It also may have some visceral analgesic activity.

Methylnaltrexone is a peripherally acting mu-opioid-receptor antagonist that is licensed for the treatment of opioid-induced constipation in patients receiving palliative care when response to other laxatives is inadequate; it should be used as an adjunct to existing laxative therapy. Methylnaltrexone does not alter the central analgesic effect of opioids. Naloxegol is an oral agent and has the same properties.

**Faecal softeners** (liquid paraffin), the traditional lubricant, have potential disadvantages of minimal efficacy (hence usually used in combination with other agents) and safety issues (aspiration of paraffin, perianal burning). Bulk laxatives and non-ionic surfactant ‘wetting’ agents (docusate sodium) also have softening properties. Enemas containing arachis oil (ground-nut oil, peanut oil) lubricate and soften impacted faeces and promote a bowel movement. Diocyl sulfosuccinate, an anionic detergent, can be used to break down hard stools.

**Stimulant suppositories** (glycerol) or enemas (phosphate) may also be effective although they are often less acceptable to the patient. Glycerol suppositories act as a rectal stimulant by virtue of the mildly irritant action of glycerol. Constipation may need regular enemas initially using low volume phosphate preparations progressing to high volume saline washouts or transanal irrigation systems.

Treatment of faecal impaction may need a manual evacuation under anaesthetic if disimpaction does not occur after oral and rectal treatment, or if there is a megarectum. The outcome of colectomy±ileorectal anastomosis is poor for these patients and best avoided. Sometimes a defunctioning loop ileostomy, which is reversible, may be performed before considering a total colectomy.

### 9.2.2 Pain

Pain is often poorly correlated with motor events. A simple measure such as reducing fibre in the diet can reduce abdominal distension by reducing bacterial fermentation and the production of gases. Low FODMAP diets may also have a role, but are restrictive in nature and should not be used in an already malnourished individual. Peppermint oil may also help.

Antimuscarinics that are used for gastrointestinal smooth muscle spasm include the tertiary amine dicyclomycin hydrochloride and the quaternary ammonium compounds propantheline bromide and hyoscine butylbromide. The quaternary ammonium compounds are less lipid soluble than atropine and are less likely to cross the blood–brain barrier; they are also less well absorbed from the gastrointestinal tract. Dicyclomycin hydrochloride has a much less marked antimuscarinic action than atropine and may also have some direct action on smooth muscle. Hyoscine butylbromide is advocated as a gastrointestinal antispasmodic and is commonly tried, but it is poorly absorbed so intramuscular preparations may be more effective and can be used in the long term at home.

Direct relaxants of intestinal smooth muscle (alverine citrate, mebeverine and peppermint oil) which may relieve pain in irritable bowel syndrome are commonly tried and have no serious adverse effects.

Persistent abdominal pain may be a major problem and its mechanism may include central nervous system sensitisation making it very difficult to treat. Features of neuropathic pain should be sought and managed with neuropathic agents. Opioid-induced hyperalgesia as part of the narcotic bowel syndrome can also develop and needs appropriate management (see section 5.2). If other analgesics prove ineffective, opioids and their derivatives may be tried though they themselves have pro-absorptive/antisecretory effects and cause slowing of intestinal transit. Some opioids, such as tapentadol, may have a better dysmotility side effect profile. 'Targnact (oxycodone and naloxone combined) is marketed as a way of giving analgesia without causing constipation. A small amount of naloxone crosses the blood–brain barrier to block the dependence action of oxycodone, although when first used it may precipitate some withdrawal symptoms from the previous opioid-like drug. Naloxegol is a PEGylated naloxone formulation not combined with opioid and therefore should not cross the blood–brain barrier, owing to the PEGylation, to minimise withdrawal side effects.

Oral liquid preparations may be used but sublingual or transdermal buprenorphine or fentanyl have the advantage of bypassing the abnormal gut function. It may be of value to give a patient a ‘pain holiday’ in hospital during which sedation and continuous subcutaneous opiates, or even epidural anaesthesia, may reduce the pain threshold so allowing a reduction in maintenance analgesic dosage. Escalation beyond a low dose of opioid is likely to be ineffective in managing chronic pain and is associated with unacceptable risks, including catheter-related bloodstream infections, and should be de-escalated or discontinued if the chronic pain persists, even if no other effective pain medication is available.

There is a growing appreciation for the role in abdominal pain management of the gut-brain neuromodulators, frequently used in neuropathic pain management. These include a tricyclic agent that can be used at sub-antidepressant doses for abdominal pain or discomfort in patients who have not responded to laxatives, loperamide or antispasmodics. Low doses of a tricyclic antidepressant are used (eg, amitriptyline, initially 5–10 mg each night, increased if necessary in steps of 10 mg at intervals of at least 2 weeks to maximum 30–50 mg each night). A selective serotonin reuptake inhibitor (SSRI) may be considered in those who do not respond to a tricyclic antidepressant, but they are considered to be less effective analgesics than the serotonin-norepinephrine reuptake inhibitors (SNRIs), of which duloxetine is the first choice. The role of gabapentin and pregabalin is well established for chronic neuropathic pain, and there is emerging evidence that the use of combination gut-brain neuromodulators from more than one class is more effective than monotherapy. SNRI and SSRI classes, however, should not be combined due to risks of serotonin syndrome.

### 9.2.3 Vomiting

Now that domperidone and metoclopramide are no longer used in the long term and as cyclizine can cause psychological dependence and addictive behaviour, the 5-HT3 antagonists like ondansetron are most commonly used but can result in constipation.

If a nasogastric draining tube helps symptoms, then a venting gastrostomy (ideally over 20 French gauge (FG)) usually inserted endoscopically (though it may also be done radiologically or surgically) may reduce vomiting by decompressing the stomach. The difficulty is having a large enough gastrostomy tube to allow all debris from the stomach to drain. Sometimes a tube can be inserted into the small bowel if very dilated to decompress it. These venting ostomies are often successful but are associated with many complications such as leakage and infection (often

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with candida) and generally need to be changed more regularly than a feeding ostomy.

9.2.4 Diarrhoea: bacterial overgrowth and bile salt malabsorption

Antidiarreheal drugs such as loperamide, diphenoxylate or, rarely, codeine phosphate are very occasionally used for symptomatic benefit. Opioids with a central action, such as codeine, are not the first choice because of the risk of dependence and sedation.

Steatorrhoea may be secondary to an overgrowth of anaerobic bacteria in the motionless dilated loops of bowel.

As the disease progresses, bacterial overgrowth can result in diarrhoea. This can be reduced with oral amoxicillin-clavulanic acid combination, metronidazole/trimethoprim, cephalosporin, tetracycline (doxycycline), ciprofloxacin, cotrimoxazole or non-absorbable antibiotics such as rifaximin or neomycin. Rifaximin is often the first choice if it is on the local drug formulary. These may be used as necessary or in repeated courses every 2–6 weeks, often rotating (sometimes with a 1–2-week period of no antibiotic) to another antibiotic for a similar period of time before repeating. If metronidazole is used in the long term, the patient must be warned to stop if they develop numbness or tingling in their feet as an early sign of reversible peripheral neuropathy and be used at the lowest dose possible. Ciprofloxacin use longer term can cause tendonitis and rupture and again low dose and vigilance are required. The risk of resistant organisms, including *Clostridioides difficile*, should also be considered. There are no data currently about the use of probiotics.

Bacterial overgrowth is virtually inevitable and can cause cachexia without necessarily causing diarrhoea, so antibiotics (as suggested above) may be needed.

Bile salt malabsorption may occur and respond, if tolerated, to bile salt sequestrants (cholestyramine and colesvelam). It is most likely if terminal ileum has been resected or if there are large areas of fluid-filled dilated bowel.

Octreotide, occasionally used for its effects in reducing secretions and slowing gastrointestinal motility, has also been used in refractory SIBO.

9.2.5 Bloating/distension

Bloating and distension are common symptoms and not easy to treat. Reducing gas-forming microbes (eg, those producing CO₂, methane and hydrogen sulphide) with a low fibre diet or an antibiotic (rifaximin), giving simethicone or peppermint oil or reducing visceral hypersensitivity (antidepressants) or constipation may help along with increasing physical exercise.128

9.3 Neur modulation (pacemakers)

There is some limited evidence for gastric electrical stimulation (gastric pacing) to improve vomiting symptoms where gastroparesis is prominent and small bowel function relatively intact.129–131 Patients with diabetic gastroparesis respond best and, in general, responders tend to have more severe vomiting. Patients with idiopathic gastroparesis have a potentially higher rate of poor response to gastric electrical stimulation.122 Non-invasive vagal nerve stimulation has shown some promise both for improving gastroduodenal motility and reducing pain sensitivity.113

The dorsal column pathways are involved in the transmission of visceral pelvic pain. Spinal cord stimulation suppresses the visceral response to colon distension in an animal model and therefore may be an effective therapy for chronic pelvic pain of visceral origin. There has been success reported in one study of 35 patients in whom the catheter tip was situated at the T5 position for a median of 9 days (range 4–14).134 The Cochrane database concludes that more studies are needed.135

Sacral nerve stimulation uses electrical stimulation applied to the sacral nerves, eliciting a physiological effect on the lower bowel, anal sphincter and pelvic floor, and has shown some success in treating faecal incontinence and constipation.136–138 A Cochrane review concluded that from three studies there was “very limited evidence that sacral nerve stimulation can improve continence in selected people with faecal incontinence, and reduce symptoms in selected people with constipation” and larger “good quality randomised crossover trials are needed”.139 There is no evidence that sacral nerve stimulation helps any of the symptoms in patients with CIPO.

Small intestinal electrical stimulation is at an experimental stage.

9.4 Nutritional support

Nutritional assessment and support is an important aspect of management. With appropriate therapy, many patients with CIPO manage to maintain their nutritional status through the oral/enteral route without the need for PS.

9.4.1 Dietary adjustments/fluid management

Gastric motility may be far less deranged for liquids than for solids with the result that many patients tolerate liquid feeds better than solid meals. Sometimes frequent small meals with low-fat, low-fibre and liquid nutritional supplements may be helpful.

If the patient has a stoma and a short bowel, fluid restriction, a sipped glucose saline solution, use of loperamide sometimes in high dose (occasionally with the addition of codeine phosphate) will reduce the risks of dehydration, sodium and magnesium depletion. If a high net secretory output occurs, a proton pump inhibitor (or occasionally octreotide) may be needed.140

Post-feeding orthostatic symptoms in partial autonomic failure may respond to dietary adjustments and drugs such as fludrocortisone, midodrine and octreotide.

9.4.2 Vitamin/mineral deficiencies

Care is needed to ensure that micronutrient deficiencies particularly of iron, vitamin B₁₂, and the fat-soluble vitamins, especially vitamin A, D and E, do not occur. Magnesium deficiency is common especially if a high output stoma. Magnesium oxide may cause fewer osmotic effects than other preparations but is expensive.

Bone mineral density is important to address and should be assessed with dual energy x-ray absorptiometry (DEXA) scanning in those with malnutrition. For patients who cannot take oral measures to improve bone density, parenteral bisphosphonates such as zoledronate should be considered.

9.4.3 Enteral nutrition

Enteral nutrition is preferred if the gut is accessible and absorbing. In carefully selected patients, feeding jejunostomy with or without decompression (venting) gastrostomy may be tried. A percutaneous endoscopic or radiological gastrojejunostomy is preferred to a direct jejunostomy where possible as direct jejunostomy tubes are more subject to leakage, retention, pain and skin problems, and gastro-jejunostomies can achieve both post-pyloric feeding and venting with generally easier endoscopic placement whereas direct jejunal tubes will often need to be placed surgically. Invasive enteral tube insertion should be preceded where possible by a trial of naso-enteral tube
feeding to ensure absorption and tolerance prior to running the risks of mortality and morbidity associated with invasive tube placement.

If liquid enteral feeds are given, any excess can be aspirated by enteric tube or gastrostomy before the start of the next meal to ensure that excess volumes do not accumulate in the stomach. Gastrostomy can be used, therefore, to aspirate liquid gastric contents (decompression of venting gastrostomy) as well as a conduit for feeding, particularly when there is a need to bypass a malfunctioning oesophagus and/or stomach. Pulmonary aspiration of large volume vomits is a very serious complication that may be difficult to prevent. A low antral site for gastro-jejunal tube placement is preferred to optimise drainage/venting and also stability of the jejunal extension.

9.4.4 Parenteral nutrition (PN)

Long-term PN should be reserved for patients with significant malnutrition or electrolyte disturbance who cannot tolerate enteral nutrition. Complications associated with total PN include infections, sepsis and cholestatic hepatic dysfunction.

If safe nutritional status cannot be maintained through the oral and enteral route, then HPN may be required. One problem of HPN in these patients is that they have more problems than do patients with a short bowel. They particularly have a higher incidence of catheter-related bloodstream infection, septicaemia and venous thrombosis. The reasons for this are not entirely clear. Procoagulation states sometimes exist, and it is possible that there is increased bacterial translocation from the gut. Opioid medication (which at high doses suppresses some aspects of immune function) and/or cyclizine increase the risk of catheter-related bloodstream infection, partly as the care taken by the patient in the management of their infusions at home, due to cognitive effects, is reduced. The use of feeding lines to administer any drug is to be strongly discouraged because of the risk of catheter infection. Such patients test the capabilities of the best organised nutrition teams to the full and should be managed in centres with a large experience. Vigilance for psychopathology and ongoing involvement of psychology and liaison psychiatry should be offered. There may be a benefit from the mutual support patients can give to each other in these situations, although patients with significant psychopathology can have a detrimental effect on others.

Howard et al. have emphasised that the clinical outcome on HPN, like the mortality risk, is to a large extent a reflection of the underlying condition. While about 70% of patients with Crohn’s disease or ischaemic bowel conditions are fully rehabilitated after the first year on HPN, only a third of those with chronic intestinal dysmotility are similarly rehabilitated and it is most likely if the gut is not dilated. Impairment of strength and of well-being as a result of undernutrition and fluid and electrolyte imbalance will be corrected by HPN, but if the patient continues to experience vomiting, diarrhoea or abdominal pain from the underlying condition, quality of life will remain suboptimal. The annual risk of catheter-related sepsis among HPN patients is consistently around 0.5 per 1000 catheter days but tends to be higher among those with chronic pseudo-obstruction, especially if they remain on opioid analgesia; by contrast, patients with systemic sclerosis who may tend to have lower opioid requirements have lower catheter infection rates.

Over half of those with pseudo-obstruction receiving HPN will be alive at 10 years.

9.5 Surgical options

Surgery is to be avoided in this group of patients who are at high risk of iatrogenic injury; however, judicious palliative surgical intervention (resection, bypass or stoma formation) can improve symptoms and quality of life. If constipation is difficult to manage and high volume saline washouts are needed, then colectomy with ileorectal anastomosis or ileostomy may be necessary but diarrhoea or continuing episodes of obstruction may remain a problem. Adhesiolysis in the absence of a clear focal obstruction carries a high risk of severe complications and morbidity with ultimately more adhesion recurrence and worsening pain. Urology is often needed to help with neuromuscular disorders of the urinary tract (dilated ureters and bladder) and stents and/or a suprapubic catheter may need to be inserted. Especially in women, there may be fertility problems due to dilated non-functioning fallopian tubes. Often the pains experienced result in gynaecological referrals.

9.5.1 Bypass surgery and enteric resections

There are several reports of surgery in adults to help these patients with pseudo-obstruction, although the clear separation into those with a myopathy and those with a neuropathy is not always made. After diagnostic laparotomy, bypass operations (gastro-enterostomy, duodeno-jejunostomy and jejunoo-enterostomy) can be performed in adults to reduce vomiting if there is dilated gut. If gastric surgery is being performed, a vagotomy must be avoided as this will further retard gastrointestinal transit. Many have an ileostomy often to treat constipation and some develop a short bowel from multiple resections. The reports of success are variable and any undertaking of surgery needs to be a multidisciplinary decision and based on each individual patient. Outcome is poorer in patients with evidence of small bowel dysmotility who undergo colectomy.

9.5.2 Small intestinal transplantation

Dysmotility is a rare indication for intestinal transplantation in adults with dysmotility needing HPN, but since the outcomes of HPN are currently better, transplantation should be reserved for those who develop complications related to PN including IF-associated liver disease, central vein thrombosis with reduced venous access, and recurrent catheter-related bloodstream infections. If other organs are damaged, a multivisceral transplantation may be considered. The role of small bowel transplantation solely to improve quality of life by ceasing PN is somewhat contentious, but as worldwide experience of transplantation increases with corresponding improvements in survival rates, the indications for transplantation may broaden in the future. It is vital that all patients considered for transplantation are reviewed by an experienced MDT with expertise in IF and a transplant centre. Pain is not a good indication for a transplant.

9.6 Psychosocial treatments

Psychological support from nurses, physicians and psychologists is important. Vigilance needs to be maintained for the presence of psychopathology even in patients with a strong suspicion of gastrointestinal neuromuscular disorder. In one case series, six patients diagnosed initially with IF had significant psychopathology requiring specialised psychiatric unit treatment. In addition to psychological distress including anxiety and depression, other psychological problems encountered can include somatisation disorder, personality disorders, substance misuse and disordered eating (see also section 5.4). Dysmotility disorders can also be associated with a risk of self-harm including suicide. Clinical psychology and liaison psychiatry provide
overlapping but complementary approaches and ideally a MDT involving both specialties should be available.

10.0 OUTCOMES
Outcome can vary from minor symptoms consistent with irritable bowel syndrome to problems resulting in home parenteral feeding, major analgesics and frequent hospital admissions. Causes of death in these patients include pulmonary aspiration, pulmonary embolism, cardiac failure and suicide. Cardiac failure may be the terminal event in hollow visceral myopathy. The relationship between 'megaduodenum' and upper gastrointestinal tract cancer seems tenuous if it exists. Death will often be related to the underlying condition—obviously so in the case of pseudo-obstruction occurring as a paraneoplastic phenomenon, and also particularly in the degenerative neuropathies, collagen vascular disorders and infiltrative conditions such as amyloid. Amiot et al reported 51 patients with a CIPO who required HPN, representing 26 years of experience, and found that surgery was required in 84% of patients and survival probability was 94%, 78%, 75% and 68% at 1, 5, 10 and 15 years, respectively. Higher mortality was associated with systemic sclerosis. The 20-year experience of HPN from the Mayo Clinic found that the survival for patients with dysmotility disorders was second worse only to cancer due to the progression of the underlying disease, which was similar to data from St Mark’s Hospital in the UK. Recently published data from the Salford IF Unit have demonstrated worse outcomes for patients with a CIPO than a non-CIPO dysmotility phenotype. However, it would appear that there is room for improved outcomes in this challenging patient population by cost-effective investment in tertiary multidisciplinary provision.

11.0 SUMMARY
Most cases of intestinal dysmotility will be without a clear diagnosis and thus labelled as idiopathic. Addressing the patient’s primary symptoms and treating malnutrition are the keys to management.

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Contributors JMDN wrote the manuscript. PP, JM, AE and SL contributed additional material and adjusted the text. JEM provided pathology input.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement There are no original data in this work.

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