Educational Gap

A recent study in Pediatrics concluded that 25% of children with functional constipation continued to experience symptoms at adult age, suggesting that referral to specialized clinics at an early stage for children who are unresponsive to first-line treatment may help improve outcomes. (1)

Objectives

After completing the article, the reader should be able to:

1. Know that constipation is a common problem in childhood with a diverse clinical presentation.
2. Understand that functional constipation is a symptom-based diagnosis that does not require extensive testing.
3. Recognize that most children who present with fecal incontinence or encopresis have associated constipation.
4. Describe the treatment of constipation and encopresis, which should include a medical-behavioral approach that focuses on maintaining soft and regular bowel movements and improving toileting behavior.

INTRODUCTION

What do the following children have in common?

- A 12-month-old girl with hard pellet-like stools.
- A 3-year-old girl with frequent complaints of dysuria and hard stools.
- An 8-year-old boy with a weekly stool that is large enough to clog a toilet.
- A 12-year-old boy with daily loose stools in his underpants.

Answer: They share a familiar diagnosis: functional constipation.

Constipation is a common pediatric problem and parental concern. In general, a complaint of constipation accounts for 5% of general pediatric office visits and 25% of all referrals to pediatric gastroenterologists. The estimated worldwide prevalence is 0.7% to 29.6%. (2) Constipation rarely signifies a serious disease, but it has an unfavorable impact on patient quality of life, parental satisfaction, and health-care costs. Children with constipation often complain of abdominal pain, decreased appetite, and painful stooling, which can be distressing to both the child and the parents. Common transient problems with defecation, if unrecognized and untreated, can develop into disruption of toilet training and

AUTHOR DISCLOSURE Drs Colombo, Wassom, and Rosen have disclosed no financial relationships relevant to this article. This commentary does not contain a discussion of an unapproved/investigative use of a commercial product/device.
interference with achieving and maintaining bowel continence. The chronic nature of constipation and common misconceptions about the symptoms and pathophysiology of constipation can lead to frustrating experiences for patients and families. Multiple office visits, emergency department visits, and unnecessary diagnostic testing contribute to the rising cost of health-care.

DEFINITION

Defining constipation remains a challenge because stooling patterns are highly variable in childhood. Generally, infants have an average of three to four stools per day and a toddler may have two to three stools per day. (3) By the age of 4 years, children have a pattern and frequency of bowel movements that are similar to those of adults. (4) A consistent, reliable method for diagnosis allows for better understanding of and communication about the disorder. Constipation can be roughly defined as infrequent passage of hard, uncomfortable stools that are distressing to the child.

Encopresis is the repeated passage of feces into inappropriate places (usually the underpants). Some have suggested replacing the term fecal incontinence with the term encopresis in the literature to clarify that most children treated for this problem have either current or intermittent constipation. However, we continue to reference the term encopresis along with fecal incontinence due to the prevalence of this term in the literature. Fecal incontinence/encopresis is often the result of liquid/soft stool leaking around a large mass of stool in the rectum, which clinicians should describe as constipation with overflow.

Encopresis differs from delayed bowel training in that children with encopresis pass liquid/soft stool in their underpants unknowingly because of constipation with overflow and difficulty feeling the indication to stool. Children with encopresis also generally do not have accidents of formed stool whereas children with delayed bowel training simply refuse to use the toilet and have regular bowel movements in their diapers or underpants. Children with encopresis often also use the toilet to pass formed or semiformal stool.

Children with delayed bowel training may refuse to use the toilet because of fear, anxiety, oppositional behavior, skill deficits, or lack of interest or motivation. Bowel continence is expected to occur by the age of 4 years. Encopresis is not a developmental variation after the age of 4 to 5 years.

PATHOGENESIS

Causes for the development of constipation include inadequate hydration, low-fiber diet, slow intestinal transit, minimal activity level or inactivity, and behavioral factors. Because some or all components may play a role in the development of constipation and encopresis, these conditions should be conceptualized in the biopsychosocial framework. Constipation can manifest at any age and most commonly presents during a period of transition in the child’s life. In infancy, constipation may present when the breastfed infant is transitioned to formula or whole milk or when transitioning from pureed to solid foods. In toddlers, constipation may arise when toilet training begins. In childhood, constipation is more likely when a child enters school and is using a toilet away from home.

Normal Anatomy and Physiology

The internal anal sphincter, external anal sphincter, puborectalis muscle, and rectum must work together for a productive bowel movement. The internal anal sphincter and the rectum are composed of circular smooth muscle. The external anal sphincter and puborectalis muscle are made up of skeletal muscle. When the rectum is empty and collapsed, the internal and external anal sphincters are tonically contracted, maintaining continence. The puborectalis muscle forms a sling around the rectum, pulling the rectum forward when it is contracted and increasing the angle acuity between the rectum and the anus.

When a bolus of stool reaches the rectum, distension of the rectal wall signals the urge to defecate. The internal anal sphincter reflexively relaxes and the external anal sphincter contracts. There are two options at this time: 1) squatting or sitting on the toilet, relaxing the puborectalis muscle, straightening the anorectal angle, relaxing the external anal sphincter, and increasing intra-abdominal pressure to evacuate stool or 2) maintaining and increasing contraction of the external anal sphincter and gluteal muscles to force stool back into the rectum. When the stool is pushed back into the rectum, the sensation or urge to have a bowel movement disappears.

Constipation and Withholding

When children do not recognize or respond to the urge to defecate, stool is retained in the rectum, the urge to defecate subsides, and the rectal wall stretches to accommodate the fecal load. Repeated withholding or avoidance of defecation leads to larger stool load in the rectum, causing further stretching and potential thinning of the rectal wall. The retained stool becomes larger, harder, drier, and more difficult to pass the next time the urge arises.

Prolonged and repetitive stool withholding and avoidance of defecation leads to large amounts of retained stool in the rectum. The large fecal mass becomes impacted and
exceedingly rare. Functional constipation is most common in childhood.

**Functional**
Infant dyschezia describes healthy infants younger than 6 months of age who strain excessively with bowel movements. They appear to be in significant discomfort, often crying or screaming, turning red in the face, and bringing their knees up to their abdomens, before eventually passing soft stools. Symptoms abate following the stool passage. Infants of this age have not yet coordinated increasing intra-abdominal pressure with relaxation of pelvic floor muscles to have a bowel movement. Infant dyschezia often spontaneously resolves around 6 months of age.

Functional constipation and functional fecal retention are synonymous and the terms are often used interchangeably. Functional constipation refers to hard or infrequent stools. Symptoms abate following the stool passage. Infants of this age have not yet coordinated increasing intra-abdominal pressure with relaxation of pelvic floor muscles to have a bowel movement. Infant dyschezia often spontaneously resolves around 6 months of age.

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**TABLE 1. Differential Diagnosis of Constipation and Defecation Disorders**

<table>
<thead>
<tr>
<th>FUNCTIONAL</th>
<th>NEUROLOGIC</th>
</tr>
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<tbody>
<tr>
<td>Infant dyschezia</td>
<td>Hirschsprung disease</td>
</tr>
<tr>
<td>Functional constipation</td>
<td>Neuronal dysplasia</td>
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<tr>
<td>Nonretentive fecal soiling</td>
<td>Anal achalasia</td>
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<td></td>
<td>Disorders of the spinal cord</td>
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<tr>
<td>Obstructive</td>
<td>Endocrine/Metabolic</td>
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<tr>
<td>Anal stenosis</td>
<td>Hypothyroidism</td>
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<tr>
<td>Anterior displacement of the anus</td>
<td>Celiac disease</td>
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<tr>
<td>Small left colon syndrome</td>
<td>Diabetes</td>
</tr>
<tr>
<td>Meconium ileus</td>
<td>Cystic fibrosis</td>
</tr>
<tr>
<td>Colonic stricture</td>
<td></td>
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</table>

**Medications**
- Opioid narcotics
- Anticholinergic agents
- Tricyclic antidepressants
- Decreased fluid intake for age

**Other**
- Sexual abuse
- Chronic intestinal pseudo-obstruction
stools in the absence of any other disorders, including neurologic, obstructive, endocrine, or metabolic, and is discussed in more detail throughout this review.

**Neurologic**

Hirschsprung disease (HD) is a congenital form of constipation in which the infant or child cannot evacuate stool due to a lack of ganglion cells in the myenteric and submucosal plexus of the intestinal wall. Without ganglion cells and nerve fibers to innervate the intestinal musculature, the affected colonic segment remains in a chronic contracted state.

HD should be considered in any newborn who has delayed passage of meconium (ie, beyond 48 hours after birth). Age at presentation may depend on the length of the affected intestine. Infants with long-segment HD develop signs of distal intestinal obstruction, which may include abdominal distention, vomiting, irritability, lethargy, and failure to pass meconium or stool. Enterocolitis with or without bowel perforation must be considered if the infants develop fever, bloody diarrhea, and continued abdominal distention. One of the most serious and possibly fatal complications of Hirschsprung enterocolitis is progression to toxic megacolon and overwhelming sepsis.

Infants with shorter segments of HD may not be diagnosed until childhood. They may experience intermittent abdominal distention and severe constipation that is refractory to standard treatment. They may also have poor growth or failure to thrive due to decreased caloric intake. Only rarely do children who have HD experience encopresis or inadvertent leakage of stool.

Anal achalasia is the failure of the internal anal sphincter to relax despite the presence of ganglion cells on biopsy. It is unclear if anal achalasia is a variant or mild form of HD.

Neuronal dysplasia and hypoganglionosis are rare disorders involving inadequate or inappropriate numbers of ganglion cells. These conditions are infrequent and incompletely understood. They can be associated with neurofibromatosis, multiple endocrine neoplasia type IIb, or Chagas disease.

**Obstructive**

Anal stenosis presents with painful and difficult defecation in infancy. This is due to the presence of a tight anal opening or ring.

Anterior displacement of the anus is a congenital variation in the placement of the anus. External anal inspection can reassure the clinician that anterior displacement is not present. Theoretically, the anogenital index can be calculated after careful examination and measurement of the perineum by dividing the distance (in centimeters) from the vagina or scrotum to the anus by the distance (in centimeters) from the vagina or scrotum to the coccyx. The normal anogenital index in females is 0.30 ± 0.09 and in males is 0.56 ± 0.2. Affected children have difficulty with defecation because they are not able to straighten the anorectal canal completely due to malposition of the anal sphincter complex in relation to the anus.

Meconium ileus causes delayed passage of meconium in children with cystic fibrosis. It is one of the earliest signs of cystic fibrosis and is almost always associated with pancreatic insufficiency. The meconium in infants with cystic fibrosis is much thicker than the meconium in unaffected children, which is attributed to an altered ratio of albumin and water concentrations. The viscous meconium and increased mucus production can lead to partial or total bowel obstruction. This is often recognized at birth or within the first few days of birth.

Strictures can occur anywhere in the intestinal tract. If they occur more distally, children often have symptoms of distal obstruction, including lower abdominal pain, abdominal distention, and infrequent or total lack of bowel movements. Strictures can be congenital or acquired due to necrotizing enterocolitis in infants or inflammatory bowel disease in children and adolescents.

Small left colon syndrome is a rare diagnosis that is most closely associated with infants born to women who have diabetes. Infants with signs of distal intestinal obstruction and a small left colon noted on barium enema should be screened for HD and cystic fibrosis because a small-caliber left colon is a common finding on barium enema due to the aganglionic segment in HD or as a result of meconium ileus in cystic fibrosis.

**Other**

Hypothyroidism may slow the motility of the gastrointestinal tract, leading to constipation. Other symptoms of hypothyroidism include fatigue; weight gain; shortness of breath; and changes in the skin, hair, or nails. This constellation of findings can help direct appropriate testing.

Celiac disease is an autoimmune sensitivity to gluten and gluten-containing products in genetically susceptible individuals. The clinical presentation of this disease is so variable that the “atypical” or nonclassic presentation of celiac disease is becoming more commonplace. Children may present with diarrhea, constipation, bloating, abdominal pain, poor weight gain, short stature, skin rash, or iron deficiency anemia. Evaluation for celiac disease should be considered in children who have constipation that does not respond to laxative therapy.
Chronic intestinal pseudo-obstruction manifests as severe altered motility of the intestinal tract. This rare disorder can be congenital or acquired. Children with this disorder experience recurrent signs and symptoms of bowel obstruction, such as vomiting, abdominal pain and distention, and constipation or diarrhea without an anatomic obstruction.

**CLINICAL ASPECTS**

Diagnosis of constipation relies foremost on an appropriate definition, including the symptom-based Rome III criteria for functional constipation. Rome III defines functional constipation as two or more of the following (fulfilled at least weekly for 2 months) in a child older than 4 years who does not have irritable bowel syndrome:

1. Two or fewer defecations in the toilet per week.
2. At least one episode of fecal incontinence per week.
3. History of retentive posturing or excessive volitional stool retention.
4. History of painful or hard bowel movements.
5. Presence of a large fecal mass in the rectum.
6. History of large-diameter stools that may obstruct the toilet.

Similar age-appropriate criteria are also available for children younger than 4 years. A variety of supplemental tests should be applied only in the presence of warning signs or symptoms or with failure of constipation to respond to typical therapy. Test results should be interpreted in the context of the patient's history and physical examination findings, which are sufficient for diagnosis in most cases.

The clinical history should include a description of stool frequency and quality, associated symptoms such as abdominal pain and rectal bleeding, growth pattern, continence and toilet training, presence or absence of withholding behavior, and symptom onset and duration. Delayed passage of meconium should raise suspicion for HD. Thin, ribbonlike stools also may suggest HD compared to the large bulky stools that often are found with functional constipation. Fecal incontinence should be directly assessed in the context of the patient's history and physical examination, which are sufficient for diagnosis in most cases.

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Standardized measures such as the modified Bristol stool form scale (6) or Amsterdam infant stool scale (7) allow for a common language and description of stools. These measures can be used in combination with age-appropriate questions that engage the patient and caregiver in an effort to overcome barriers of symptom anxiety, embarrassment, or denial (eg, Does it take you a long time to push poop out of your body? Does the poop hurt your bottom when it comes out?). Physically exemplifying withholding behavior during the interview sometimes provides a moment of clarity for caregivers who thought such behavior indicated an attempt at defecation.

Physical examination should explore both the severity of constipation and potential causes. Ideally, a growth curve contains data spanning the onset of constipation to determine current parameters as well as past growth velocity. Observation of the patient should not be underestimated in its ability to elicit information; interaction between the patient and caregiver, willingness of the patient to engage in toileting discussion, and ability to sit and climb on or off the examination table can provide important diagnostic clues and sometimes direct treatment strategies. Abdominal distension, tenderness to palpation, and presence of fecal mass as well as perianal examination for skin tags, fissures, and anal appearance and location are important for all new patients in whom constipation is suspected. External anal inspection can assess for anal atresia and displacement and may identify anal fissures, skin tags, or external hemorrhoids. It may also be useful to assess sphincter tone visually or identify fecal material around the anus or in the underwear. In addition, examining the back for sacral dimples or spinal deformities and assessing lower extremity motor tone, strength, and deep-tendon reflexes can indicate whether additional assessment for neurologic pathology is indicated.

Digital rectal examination (DRE) is important in specific circumstances but is not always necessary to diagnose functional constipation. Palpation of a firm or large rectal stool mass on rectal examination often confirms clinical suspicions, abnormalities in sphincteric tone may indicate anal stenosis, and an empty rectal vault with expulsion of stool on finger withdrawal is a classic but infrequently seen finding in HD. Performing a DRE should be left to the discretion of the clinician. DRE may provoke anxiety or fear in children who have had past experiences with painful stool passage. It is important to avoid reinforcing this negative association when possible and limiting frightening or painful interventions, which can aid in building a therapeutic alliance between the patient and caregiver.

Laboratory evaluation is not warranted for constipation unless warning signs are present (Table 2) or other aspects of the history or physical examination suggest systemic
disease. Constipation rarely is the sole presenting symptom of hypothyroidism, electrolyte abnormalities, lead toxicity, or celiac disease, and routine screening for these diseases is not recommended. Routine allergy testing is also not recommended in evaluation of constipation, and cow milk protein restriction in young children for a limited time to assess the clinical response remains controversial.

History and physical examination generally precludes the need for radiography to diagnose functional constipation. Although the presence or absence of a fecal mass and determination of stool burden are important to direct therapy, abdominal radiography is usually not necessary. An abdominal radiograph may help parents visualize the amount of retained stool, allowing for a better understanding of constipation (with or without overflow) and the proposed treatment plan. A single abdominal radiograph is an inexpensive, low-risk test, but even when using standardized scales to determine stool burden, it is not clearly reliable, sensitive, or specific.

The indications for barium enema are extremely limited. Barium or other contrast enema is suggested but not required when constipation is accompanied by “red flag” symptoms (Table 2). It provides information about the caliber of the rectum and colon and may be useful if obstruction in the colon is suspected. It does not require any specific preparation and does not subject the patient to risk aside from radiation exposure. Gastrografin enemas in the setting of suspected meconium ileus may be diagnostic as well as therapeutic.

Further diagnostic tests when the clinician suspects HD depend on patient characteristics (age, health status) and test availability. Full-thickness rectal biopsy remains the gold standard for diagnosis and is performed under anesthesia, but rectal suction biopsy can be performed at the bedside without adjunct medication and is recommended in lieu of surgical biopsy as initial evaluation. Although rectal suction biopsy traditionally is performed in infants, it may also be sufficient in older children and teenagers. Aganglionosis or hypertrophied nerves on rectal biopsy hematoxylin and eosin staining can indicate HD. Although the presence or absence of these findings is typically sufficient for determining or ruling out the diagnosis of HD, supplementary analysis may demonstrate aganglionic intestine with altered acetylcholinesterase morphology or absent calretinin expression. Anorectal manometry uses a small rectal balloon and anorectal pressure sensors to determine the presence or absence of the rectoanal inhibitory reflex (relaxation of the internal anal sphincter in response to rectal distension). Although not used as an isolated test to diagnose HD, clear demonstration of the rectoanal inhibitory reflex is sufficient to remove HD from diagnostic consideration.

Anorectal manometry may also have a role in determining rectal sensation threshold and the presence of anorectal dyssynergia, potentially directing therapy, including the addition of physiotherapy or biofeedback. High-resolution anorectal manometry with increased number of pressure sensors and enhanced computer analysis may better delineate anorectal sensory and motor function, but this test is not widely available in pediatrics.

Spinal imaging, including magnetic resonance imaging, should be considered in the child with constipation and other neurologic signs or symptoms, including lower motor dysfunction, lower urinary tract symptoms, and lumbosacral spinal abnormalities. The neurologic examination may yield normal results in constipated children with spinal cord abnormalities, but routine spinal imaging of constipated children is not recommended.

**MANAGEMENT**

Constipation and fecal incontinence are clinical issues that require a thorough understanding of physiology, biology, behavior, and psychology for effective management. Therefore, a combined treatment approach is recommended. Although no objective clinical trials and data support a single treatment approach to constipation and fecal incontinence, we discuss a general management protocol. The management protocol can be divided into four major treatment components: (1) education about constipation and encopresis, (2) disimpaction or cleanout of stool, (3) maintenance laxative therapy and establishing regular bowel movements, and (4) behavior modifications to improve daily toileting behaviors.

**Education**

Education and reassurance comprise the first component in the management of functional constipation and encopresis.
or fecal incontinence and should continue throughout all stages of management. Education includes talking with the parents and the child about constipation and its influence on lower gastrointestinal tract functioning and overflow and fecal incontinence. Developmentally appropriate discussion of the anatomy and physiology of the lower gastrointestinal tract and defecation is important and visual diagrams can aid in this education. Many parents of children presenting with encopresis incorrectly assume that the child is soiling on purpose. Parents need to be educated that fecal incontinence is often involuntary and the result of overflow from constipation, deconditioning, or altered function of the rectum and pelvic floor as well as learned withholding behaviors in some children. Counseling can be provided to parents to help them establish a positive and supportive attitude toward their child during treatment. Counseling can help remove blame from the child and encourage the parents and child to join the clinician in addressing the symptoms. Clinicians should recognize and empathize with families about the stress and frustration surrounding constipation and fecal incontinence.

Disimpaction or Cleanout
The second component in the management of functional constipation is removal of the fecal impaction. Such removal decompresses the rectum, allows for the normal passage of stool, and prevents liquid stool from leaking around the fecal mass. If the fecal impaction is not removed, a child with functional constipation cannot achieve a normal stooling pattern, and fecal soiling may be exacerbated, which is highly frustrating to parents and children.

Among the approaches to disimpaction are high-dose oral laxatives, enemas, manual disimpaction, or admission to the hospital for nasogastric administration of a bowel cleansing agent. High-dose oral laxatives and enemas are equally efficacious, but the preferred method for evacuation of fecal impaction is via the oral route. Minimizing attention to the anus and rectum via oral laxatives can be important because these children have a history of unpleasant and painful experiences associated with defecation. Current recommendations (Table 3) suggest the use of polyethylene glycol solution (PEG 3350) at doses of 1 to 1.5 g/kg per day for 3 consecutive days (up to 6 consecutive days if necessary) to achieve disimpaction. If PEG 3350 is unavailable, once-daily sodium phosphate, saline, or mineral oil enemas for 3 consecutive days are acceptable. Suppositories may be used in combination with high-dose oral laxatives to help promote evacuation of the fecal impaction. Manual disimpaction is rarely necessary and generally not advised except in cases of severe impaction and obstipation. If manual disimpaction is required, general anesthesia should be used to decrease the trauma associated with this procedure.

Maintenance Therapy
The third component in the management of functional constipation is maintenance laxative therapy to ensure regular passage of soft, appropriate-sized stools. Such maintenance can eliminate painful defecation and prevent the recurrence of fecal impaction. This component of treatment may last many months to years and requires ongoing close follow-up evaluation. Having families use a bowel symptom tracking form or calendar to monitor the child’s response to treatment may be helpful. Clinicians should emphasize the need for close monitoring and long-term treatment with parents because nonadherence to prescribed medications or discontinuing medications too early can result in the development of hard stools and relapses of withholding, leading to fecal impaction. The most common medication used for maintenance therapy is PEG 3350 due to its ease of use, titratability, low adverse effect profile, and efficacy.

When full evacuation of the rectum consistently occurs with stooling for 1 to 2 months without any development of hard stools or withholding behaviors, the laxative medication may gradually be reduced. Early recognition of relapse by both parents and clinicians is vital to long-term treatment of functional constipation. Increasing therapy and aggressively treating a relapse can avoid prolongation of the maintenance phase.

### TABLE 3. Medical Therapy for Disimpaction

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>Oral (preferred)</td>
<td>Polyethylene glycol solution 1–1.5 g/kg/day x 3–6 consecutive days</td>
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<tr>
<td></td>
<td>Magnesium citrate 4 mL/kg/day x 2 consecutive days</td>
</tr>
<tr>
<td>Rectal</td>
<td>Normal saline enema 10 mL/kg x 3 consecutive days</td>
</tr>
<tr>
<td></td>
<td>Sodium phosphate enema x 3 consecutive days</td>
</tr>
<tr>
<td></td>
<td>Mineral oil enema x 3 consecutive days</td>
</tr>
<tr>
<td></td>
<td>Polyethylene glycol solution 25–40 mL/kg/hr until rectal effluent is clear</td>
</tr>
</tbody>
</table>

- Oral (preferred)
- Rectal
- Nasogastric (requires hospital admission)

- Polyethylene glycol solution 1–1.5 g/kg/day x 3–6 consecutive days
- Magnesium citrate 4 mL/kg/day x 2 consecutive days
- Normal saline enema 10 mL/kg x 3 consecutive days
- Sodium phosphate enema x 3 consecutive days
- Mineral oil enema x 3 consecutive days
- Polyethylene glycol solution 25–40 mL/kg/hr until rectal effluent is clear (24–48 hr)
Laxatives used for maintenance therapy should be individualized for each patient. Dosing guidelines and recommendations are suggestions for starting therapy (Table 4). In practice, laxative doses should be titrated to achieve at least one soft bowel movement every day. Understanding the mechanism of action of each laxative can aid clinicians in customizing maintenance therapy because combination therapy can be useful and necessary.

Osmotic laxatives increase the osmotic load within the lumen of the intestine, allowing for fluid retention. The retained fluid is incorporated into the stool and distends the colon, promoting peristalsis. Children may experience bloating, but these laxatives are generally safe; the most common adverse effect is diarrhea. Examples of osmotic laxatives include PEG 3350, lactulose, and magnesium products such as magnesium hydroxide and magnesium citrate.

**TABLE 4. Maintenance Therapy for Chronic Constipation**

- **Osmotic laxatives**
  - Polyethylene glycol 1 g/kg/day
  - Lactulose 1–3 mL/kg/day divided into 2 doses
  - Magnesium hydroxide
    - <2 years: 0.5 mL/kg/dose
    - 2–5 years: 5–15 mL/day once before bedtime or in divided doses
    - 6–11 years: 15–30 mL/day once before bedtime or in divided doses
    - >12 years: 30–60 mL/day once before bedtime or in divided doses
- **Stool Softeners/Lubricants**
  - Docusate 5 mg/kg/day (up to 400 mg/day)
  - Mineral oil 1–3 mL/kg/day divided into 2 doses
- **Stimulant Laxatives**
  - Senna
    - 1 month-2 years: 2.2–4.4 mg/day at bedtime or in 2 divided doses
    - 2–6 years: 4.4–6.6 mg/day at bedtime or in 2 divided doses
    - 6–12 years: 8.8–13.2 mg/day at bedtime or in 2 divided doses
    - >12 years: 17.6–26.4 mg/day at bedtime or in 2 divided doses
  - Bisacodyl
    - 3–12 years: 5–10 mg/day
    - >12 years: 5–15 mg/day

Stimulant laxatives such as bisacodyl or senna irritate smooth muscle of the colon and stimulate the myenteric plexus to produce peristaltic activity within the colon. Children may experience abdominal cramping with the peristaltic activity. The abdominal cramping is self-limited and can be reduced by decreasing the dose. Although stimulant laxatives are safe, no studies have assessed dependency with chronic daily use. Stimulant laxatives can generally be reserved for intermittent use and rescue therapy.

Stool softeners decrease the surface tension of the stool, which allows integration of more water into the stool, thereby softening it. Docusate has a modest stool softening effect. It is generally safe, with minimal adverse effects.

Mineral oil may ease the passage of stool by lubricating the intestine and decreasing water absorption. A common complaint with use of mineral oil is leaking of the oil from the rectum, which can be unpleasant. Palatability of mineral oil is also a challenge for many children. Oral mineral oil is contraindicated in children younger than age 1 year or with known or suspected aspiration.

Dietary modification is frequently considered for treatment, but increasing fluid or fiber intake has unclear efficacy in constipated children. Maintaining adequate hydration is important for a variety of physiologic functions and, in most cases, is a safe recommendation. However, solely increasing fluid intake should not be expected to alter stooling frequency or consistency. Fiber is often chosen as first-line therapy in constipated adults, but recent reviews of evidence cast some doubt on its effectiveness. Dietary fiber intake may be reduced in constipated children compared to those without constipation, but increasing fiber does not clearly improve symptoms and is not recommended as therapy based on current evidence. Fiber also may be tolerated less well than other therapies for constipation in children. Although complications related to fiber therapy are unusual, use of fiber as monotherapy may delay implementation of effective treatment and prolong patient symptoms. Growing evidence supports probiotic and prebiotic use as treatment in adult constipation. These agents may reduce whole-gut transit time, increase stool frequency, and reduce constipation-associated symptoms. However, evidence in pediatrics demonstrates mixed efficacy and their use is associated with additional patient expense, possibly to the exclusion of other effective therapies.

**Behavior Modification**

The fourth treatment component is behavior modification to improve daily toileting habits and routines. This component should be started at the time of bowel disimpaction or cleanout and continue throughout maintenance treatment.
Research regarding specific behavioral treatments for encopresis is disjointed and difficult to synthesize based on recent reviews. However, the use of operant procedures (incentive/reward programs and positive reinforcement) for goals related to toileting and cleanliness have empiric support. Incentive/reward systems can be used to target various goals related to successful toileting. The targets of intervention may vary with individual children. Rewarding the patient for cooperation with the components of the treatment regimen and NOT just for proper elimination in the toilet is important. Children must achieve important goals or behavioral skills on the way to successful toileting, such as gaining confidence and compliance with toilet sitting, responding appropriately and honestly to soiling accidents, and learning effective pushing techniques to produce complete and emptying bowel movements.

An important part of the standard medical-behavioral treatment of encopresis is improving toilet sitting behavior. However, stool withholding and toileting refusal behaviors may interfere with progress toward toilet sitting goals and sometimes must be addressed before implementing a toilet sitting plan. Stool withholding and toileting refusal are believed to be related to the history of difficult-to-pass or even painful bowel movements and are often conceptualized as an anxiety or phobia about passing bowel movements, especially into the toilet. The initial focus of stool withholding management should be to ensure soft and easy-to-pass bowel movements so that the child can gain comfort in passing a bowel movement on a daily basis. In early stages of treatment, bowel movements in a pull-up or diaper may need to be reinforced for the child to gain confidence and voluntarily relax the pelvic floor to achieve a bowel movement. Toilet refusal behavior should also be treated with interventions that gradually desensitize children toward toileting. Desensitization to the toilet may include rewarded trips to the bathroom to look at the toilet, stand by the toilet, sit on a closed lid fully clothed, and eventually sit on the toilet with open lid and pants down. Once the child is having bowel movements comfortably in the diaper or a pull-up and able to sit on the toilet without significant anxiety, parents can use a shaping procedure to encourage bowel movements closer to the toilet and eventually into the toilet. Reward systems or incentives are used to encourage children to take a next step toward successful toileting behavior.

Once the child is comfortable and compliant with sitting on the toilet, the overall goal is to improve daily toileting habits and routines. Empiric evidence suggests that operant procedures or reward systems should be an active part of a toilet sitting schedule. Scheduled toilet sits can occur 20 to 30 minutes after meals to take advantage of the gastrocolic reflex. In addition, pairing toilet sitting with meals is easier to build into the family routine and can create a behavioral stimulus condition for bowel movement success. The time on the toilet should be unrushed and positive. It may include special activities that are only available while on the toilet (special books, toys, or handheld electronics). Parents can also be counseled to provide modeling and coaching during toilet sitting, which includes the parents showing the child when they sit on the toilet and that they are pushing to help get bowel movements out in the toilet. Toilet sits should generally last 5 minutes, but some children need to gradually work their way up to longer sits if there is initial resistance. Scheduled, rewarded toilet sits should include small stepstools to assist the children in getting on the toilet and to use as leverage for their feet. We recommend a wider stepstool or potty stool to allow the child to spread out the feet and knees for better posture to allow successful defecation and for them to feel more comfortable and balanced on the toilet. Once children are having more productive bowel movements in the toilet and soiling has stopped for a 1 month, the number of daily toilet sits can be reduced. Often parents can observe which toilet sits during the day are most productive and begin focusing on those sits. As treatment progresses, children can start to earn incentives/rewards for independently going to the toilet when they feel the urge to have a bowel movement rather than strictly relying on the schedule and parental prompt.

Involving the preschool, kindergarten, or school in scheduled and rewarded toilet sitting is important. Children who are apprehensive about completing toilet sits at school benefit from a more private restroom so that they can take their time and be comfortable with toileting. It also is helpful to allow an “anytime bathroom pass” for children when they start school so that they do not withhold stool when they need to have a bowel movement.

Due to the biopsychosocial nature of functional constipation and fecal incontinence, multidisciplinary or even interdisciplinary care is becoming more common and is highly recommended when available. The combination of medical therapy, behavioral modification, and supportive counseling has the greatest success in the treatment of constipation and encopresis. When multidisciplinary or interdisciplinary care is not readily available, clinicians can still effectively treat this condition with a basic understanding of behavior modification techniques, such as the use of incentives/rewards and gradual setting of goal related to effective toileting.
PROGNOSIS

The overall prognosis for functional constipation has not been completely established. However, a general message for families is that the treatment of constipation and encopresis often requires many months of medication and behavior modification. In addition, relapse of symptoms is very common. According to a recent systematic review, approximately 60% of children with functional constipation are symptom-free between 6 and 12 months after beginning treatment regardless of laxative use, with the remaining 40% of children still experiencing symptoms. (8) In addition, a study in Pediatrics concluded that 25% of children with functional constipation continue to experience symptoms into adulthood. (9) Older school-age children and adolescents who have ongoing constipation and encopresis are even more difficult to treat. All these points highlight the need for aggressive treatment as early as possible as well as close follow-up evaluation and adjustments to the treatment plan. Nonetheless, most children with constipation and encopresis can be managed effectively by the general pediatrician. Indications for referral to a pediatric gastroenterologist include medical red flags, trouble with disimpaction, trouble establishing maintenance therapy, and lack of improvement after 6 months of therapy. Referral to a pediatric behavioral specialist should be considered if significant conditions are interfering with treatment, such as attention-deficit/hyperactivity disorder, oppositional behaviors, anxiety or mood disorders, family conflict or parent-child conflict, or problems with adherence to recommendations.

Summary

The following summary statements are based primarily on consensus and expert opinion due to the lack of relevant clinical studies. A recent comprehensive review of the literature by Tabbers et al, in the Journal of Pediatric Gastroenterology and Nutrition, identified no moderate- or high-quality evidence regarding therapeutic interventions for the evaluation and treatment of functional constipation in infants and children.

- The presentation of constipation varies, but constipation should be identified according to an appropriate definition, which includes the symptom-based Rome III criteria.
- Constipation is prevalent in children and infrequently a result of underlying intestinal or systemic disease.
- Based on limited evidence as well as consensus, history and physical examination are sufficient to provide a diagnosis of functional constipation; digital rectal examination, laboratory tests, and abdominal radiography are generally not necessary.
- Treatment of constipation requires four components: education, disimpaction, maintenance therapy, and behavioral modification.

References for this article are at http://pedsinreview.aappublications.org/content/36/9/392.full.
1. By what age do children have a pattern and frequency of bowel movements similar to those of adults?
   A. Three years.
   B. Four years.
   C. Five years.
   D. Six years.
   E. Seven years.

2. Which of the following is more characteristic of behavior by children with delayed bowel training versus children with encopresis?
   A. Have difficulty feeling the indication to stool.
   B. Generally do not have accidents with formed stool.
   C. Often use the toilet to pass formed or semiformed stool.
   D. Usually have regular bowel movements in the diaper or underpants.
   E. Pass liquid/soft stool in their underpants.

3. A 4-month-old girl presents with substantial straining with bowel movements. She cries and turns red in the face just before she passes a soft stool, after which she relaxes. Which of the following is the most likely diagnosis for this infant’s signs and symptoms?
   A. Anal achalasia.
   B. Functional constipation.
   C. Hirschsprung disease.
   D. Infant dyschezia.
   E. Neuronal dysplasia.

4. A 3-day-old term infant has had delayed passage of meconium. At 48 hours, he passed a small, thick stool. Which of the following is the most likely diagnosis?
   A. Anterior displacement of the anus.
   B. Celiac disease.
   C. Cystic fibrosis.
   D. Hirschsprung disease.
   E. Hyperthyroidism.

5. An 8-year-old boy with chronic constipation and encopresis has been successfully treated with education and disimpaction. Which of the following medications is most commonly used for maintenance therapy due to its ease of use, titratability, low adverse effect profile, and efficacy?
   A. Lactulose.
   B. Magnesium hydroxide.
   C. Mineral oil.
   D. Polyethylene glycol solution.
   E. Senna.

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