Gastrointestinal Involvement in Spinal Cord Injury: a Clinical Perspective

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Abstract

Bowel problems occur in 27% to 62% of patients with spinal cord injuries (SCI), most commonly constipation, distention, abdominal pain, rectal bleeding, hemorrhoids, bowel accidents, and autonomic hyperreflexia. The acute abdomen, with a mortality of 9.5%, does not present with rigidity or absent bowel sounds but rather with dull/poorly-localized pain, vomiting, or restlessness, with tenderness, fever, and leukocytosis in up to 50% of patients. Fecal impaction may present with anorexia and nausea. Methods used for bowel care include laxatives, anal massage, manual evacuation, and enemas. Randomized, double-blind studies demonstrated the effectiveness of neostigmine, which increases cholinergic tone, combined with glycopyrrolate, an anticholinergic agent with minimal activity in the colon that reduces extracolonic side-effects. Improved bowel function occurs with anterior sacral root stimulators which may be combined with an S2 to S4 posterior sacral rhizotomy which interrupts the reflex arc by cutting the posterior roots carrying the spasticity-causing sensory nerves. For severe constipation, a colostomy reduces time for bowel care, providing a clean environment so decubitus ulcers may heal. Gallstones occur in 17% to 31% of patients, and acalculous cholecystitis in 3.7% of patients with acute SCI. A high index of suspicion is needed to properly diagnose bowel problems in SCI.

Keywords


Introduction

There are about 200,000 patients, mostly male, with spinal cord injury (SCI) in the United States with 10,000 to 15,000 new cases diagnosed annually [1]. One-third are tetraplegic, and 50% have complete lesions [2]. The mean age is 33 years old. With improvements in critical care and rehabilitation, SCI patients are living longer and becoming functional members of society.

There are analogies between the genitourinary and the gastrointestinal (GI) tracts. Both are organs controlled by a combination of smooth internal and striated external sphincters. Their innervation, blood supply, and response to cholinergic and adrenergic stimuli are also similar [3]. Patients with difficulty in micturition are likely to have difficulties with defecation.

Gastrointestinal tract involvement in SCI

Bowel problems occur in 27% to 62% of SCI patients [3, 4], associated with depression or with injuries of highest levels, most severity, and longest duration [5]. Complaints include constipation (42-81%), distension (43-53%), incontinence (0-75%), abdominal pain (14-38%), bowel accidents (27%), nausea (19%), diarrhea (14%), straining (20%), rectal bleeding (40%), hemorrhoids (36%-75%), autonomic hyperreflexia (AH) (43%), and headaches or sweats relieved by a bowel movement (13-27%) [5-11]. Constipation is most common with complete lesions [9], previous urinary outlet surgery, and use of anticholinergic agents [6, 8, 11]. Fecal incontinence is associated with complete lesions [10] or anxiety [12]. Whether constipation [7, 10] or fecal incontinence [7, 12] varies with the level of injury is controversial. Abdominal pain may be exacerbated by eating and often improves with a bowel movement or passing flatus; it is associated with constipation and distention [3]. Abdominal distension, if severe, may cause restricted ventilation particularly with high cervical cord injuries.

Gastrointestinal complications occur in 1.9% to 11% of patients [13, 14], and undiagnosed abdominal emergencies
account for 10% of all fatalities among patients with SCI [15]. Risk factors include increasing age, male sex, head injury, multisystem trauma, and cervical spine injury [14]. Most complications occur within days after the injury, mainly GI hemorrhage, perforation, ileus, and intestinal obstruction [13, 14]. The lifetime prevalence of peptic ulcer disease is 10%, similar to the general population [3].

Spinal shock occurs for several weeks after injury when all autonomic and reflex activities are lost below the level of cord transaction. During the first four weeks, 4.7% have acute abdominal pathology [16], while 4.2% have acute gastroduodenal ulceration and hemorrhage [17], mainly with SCI above the sympathetic outflow. This may be related to the increase in serum gastrin levels seen within the first 12 to 48 hrs after paralysis caused by penetrating CNS injuries but not after closed-head injuries without paralysis [18]. The abdomen is distended and flaccid with no sensation below the level of the injury. Acute gastric dilatation with high cervical cord lesions may occur at this time [19]. There are absent or hypoactive bowel sounds due to a paralytic ileus which resolves within a week in most cases. Acute abdominal pathology is difficult to diagnose during this period without a high index of suspicion. The rectum should be evacuated using manual disimpaction or enemas [20]. Enteral feeding is safe in patients with acute SCI, the most common reason for interruption being large gastric aspirates. When spinal shock ends, there is a return and then exaggeration of reflex activity below the level of the injury.

Autonomic hyperreflexia (AH) or dysreflexia, a condition often missed by medical personnel, is a potentially fatal syndrome in those whose SCI lies above T6 or the greater splanchnic outflow [21]. It results in intense reflex vasoconstriction below the level of the SCI, with tachycardia or bradycardia, hypertension, diaphoresis, headaches, bladder spasms, and diarrhea. It is triggered by processes distending a hollow viscus associated with pain or irritation with the colorectal area being the second most common trigger after bladder irritation. Anoscopy, with stretching of the anal sphincter, is a more potent stimulus than flexible sigmoidoscopy, which causes gaseous distension of the rectosigmoid, although both can raise blood pressure and trigger AH. Intersphincteric anal block with lidocaine before diagnosis results in a perforated appendix in 92% to 100% of patients. Laparoscopy may hasten the diagnosis, thus reducing mortality and morbidity.

Acute abdominal pathology includes a perforated viscus (such as peptic ulcer disease), cholecystitis [24], appendicitis, and bowel obstruction [23]. Appendicitis presents with abdominal discomfort and distention [26]. Signs of AH with SCI above T6, fever, and right lower quadrant abdominal mass may be present but rarely tenderness. The delay in diagnosis results in a perforated appendix in 92% to 100% of patients. Laparoscopy may hasten the diagnosis, thus reducing mortality and morbidity.

Abdominal operations are challenging in patients with SCI. Patients may experience a drop in blood pressure with induction of anesthesia. Some patients have deformities, contractures, stomas, or spasticity that can technically compromise the procedures. In addition, they may be debilitated and chronically ill so that they make poor operative candidates. Retention sutures may be needed to close abdominal wounds due to increased tension that can occur from abdominal spasticity. In addition, the patients may have hyperreflexic hypertension, decreased pulmonary excursion, particularly in high-level lesions, and prolonged ileus due to poor visceral muscle tone. Pulmonary complications increase the morbidity in those undergoing surgery for appendicitis [26]. Patients often have chronic septic foci, such as UTIs, bronchitis, and decubitus ulcers that increase their risk of wound infections. If carefully
managed, however, the morbidity may be similar to that of normal individuals.

Pathophysiology of bowel dysfunction

With rectal distension, the smooth muscles of the internal anal sphincter (IAS) normally relax (rectoanal inhibitory reflex or RAIR). With rectal contractions, continence is maintained by voluntary contraction of the skeletal muscles of the external anal sphincter (EAS). Defecation occurs when the puborectalis muscle and the EAS relax.

Neurogenic bowel consists of upper motor neuron (UMN) and lower motor neuron (LMN) lesions [27]. With UMN lesions above the conus medullaris (S2-S4), the defecatory maneuver cannot be performed due to lack of functioning abdominal musculature [28]. Patients with SCI above T5 generate abdominal pressure by intercostal and diaphragmatic contraction, while those with cervical lesions rely totally on diaphragmatic contraction [29]. Rectal distension causes brief EAS activity in high cord lesions, so that leakage of fluid may not occur. It also causes autonomic symptoms, such as flushing and sweating, above the level of the lesion. The bowel is hyperreflexic with a dysrhythmic or irregular electrical rhythm on electrorectogram [30]. The tone of the colonic wall increases. The voluntary control of the EAS is discontinued so the sphincter remains tight, thereby retaining stool. The result of the UMN bowel syndrome is constipation and mild incontinence. Half the patients have bowel movements on the toilet and almost half in bed.

The LMN bowel syndrome or the areflexic bowel is due to a lesion in the conus medullaris (S2-S4), the cauda equina, or the pelvic nerve. Patients have voluntary control of part or all of their abdominal muscles. Electrorectogram shows no electrical events [30]. There is no spinal cord-mediated peristalsis but rather slow stool propulsion. The myenteric plexus coordinates segmental colonic peristalsis resulting in dry, round stools. The EAS is denervated, increasing the risk of incontinence. The levator ani muscles lack tone so the rectal angle is reduced causing the lumen of the rectum to open. The final result of the LMN lesion is constipation and incontinence-increased frequency of defecation [3] but with increased time required for defecation. Patients use the Valsalva maneuver and manual removal of stool from the rectum. With the loss of sphincter tone, the rectum must be kept empty by removal of feces with the finger to reduce incontinence. Low paraplegic patients can empty their bowels in a sitting position which reduces the anorectal angle and facilitates defecation. They have partially intact abdominal muscles needed to expel feces [8].

Patients with SCI have normal [28, 32], decreased [33], or increased [34] anal sphincter pressures. RAIR is normal in SCI since it is an entero-enteric reflex dependent on the myenteric nervous system and independent of spinal control [28, 34]. Phasic giant rectal contractions increase in SCI, perhaps predisposing to fecal incontinence [35]. From 19% to 42% of patients have a desire to defecate [10]. Signals of an upcoming defecation include abdominal pressure with incomplete lesions and unspecified abdominal complaints with complete lesions, such as a prickling sensation, increased spasticity, cutis anserine (goose bumps), and sweating [36].

The gastrocolic reflex, the increase in colonic motility with food, may be present [32] or reduced [37, 38]. This response was seen only in the descending, not the rectosigmoid, colon [37] suggesting that differences in the studies may be due to placement of the recording electrodes. Other authors found an exaggerated response in the rectum and claimed that attaching recording clips to the mucosa in other studies inhibited bowel motility [32].

A normal pressure/volume relationship during ramp inflation of the rectal balloon causes a reversed-S shape with increase in pressure initially, followed by a plateau, and then further increase in pressure as the volume increases [29]. In SCI patients, however, this pressure/volume relationship is linear, indicating a tighter and less compliant rectum at high volumes. Using radio-opaque markers or pH-sensitive radiolabelled exchange resin pellets, prolonged transit time may be pancolic [39] or predominantly in the left colon and rectum [28]. Transit times may [40] or may not [41] correlate with symptoms.

Involvement of the esophagus, stomach, and duodenum in SCI

Dysphagia is present in 22.5% to 30% of patients with SCI, related to age, presence of a tracheostomy, mechanical ventilation, and cervical spinal surgery, especially the anterior approach [42]. Factors predisposing to heartburn include delayed gastric emptying (GE), recumbency, immobilization, and certain drugs (such as anticholinergics and meperidine). Although heartburn may not be more common in SCI, these patients may undergo endoscopy less frequently than normal individuals and esophagitis is seen more commonly [43]. While resting upper and lower esophageal sphincter pressures may be normal, there is incomplete upper esophageal sphincter relaxation in some patients, predisposing them to aspiration [44]. As the posterior wall of the pharynx lies adjacent to the cervical vertebrae, cervical trauma may impinge upon or even perforate the pharynx.

Gastric emptying is either delayed [45, 46] or normal [47] in SCI patients. This discrepancy may be due to the different test meals used and whether the patients are supine [45] or sitting [47]. Gastric emptying times increase directly with the higher level of the SCI [45, 46] and the increased duration of injury [45]. Intravenous metoclopramide, a potent dopamine receptor antagonist with prokinetic properties, corrects impaired GE [47]. Functional magnetic stimulation decreases GE times by more than one-third in SCI patients with preserved LMN function, particularly during the first 30 min [48]. Superior mesenteric artery syndrome is a condition in which the third portion of the duodenum is intermittently compressed by the overlying superior mesenteric artery.
Predisposing factors include rapid weight loss (with loss of mesenteric fat that shields the duodenum as it passes behind the superior mesenteric artery and in front of the lumbar spine and aorta), prolonged supine positioning, and the use of a spinal orthosis, all of which are common with acute quadriplegic patients [49]. Symptoms include epigastric pain and vomiting, especially when supine. A barium study shows a cutoff between the third and fourth portions of the duodenum in the supine position that disappears when the patient is upright. Treatment is upright body positioning, weight gain, and in resistant cases, a duodenojejunostomy.

**Involvement of the colon in SCI**

Fecal impaction is a common GI complication in SCI [13]. Patients may present with loss of appetite and nausea and are mistakenly given antinausea preparations, most of which have anticholinergic and constipating properties. Liquid stool may pass around the blockage. Plain films of the abdomen show the feces and abnormal air patterns. Digital removal of the stool should be done. Patients may develop stercoral ulcers in the rectum from pressure due to stool. Megacolon occurs in 18% or 73% of patients, depending on how it is defined [50, 51]; small bowel dilatation found in 39% of patients may be due to stool retention most prominent in the ascending colon [50]. Colonic volvulus, reported in 2.6% of hospitalized patients, is a common cause of intestinal obstruction in patients with chronic constipation due to the long redundant colon [52].

Hemorrhoids and hemorrhoidectomy are probably more prevalent in SCI than in normal individuals, particularly in those who strain and use suppositories, enemas, or chemical stimulation for bowel management [7, 9]; hemorrhoidal bleeding occurs in up to 76% of patients [3, 53]. Conservative therapy, banding, and sclerotherapy of hemorrhoids reduces bleeding and/or symptoms [54].

Colorectal carcinoma when discovered is more advanced in SCI than in non-paralyzed subjects and morbidity after surgery may be higher [55]. A delay in diagnosis may occur since the distension, constipation, and pain may be attributed to GI complications of SCI; in addition, the difficulty of bowel preparation in these patients may result in an avoidance of screening colonoscopies.

**Treatment of bowel problems**

Bowel management involves increasing fiber intake to maintain proper stool consistency, ensuring adequate fluid intake, and providing bowel training so that the colon is evacuated at regular intervals. This is best accomplished after breakfast to take advantage of the gastrocolic reflex which peaks about 30 minutes after eating. One study surprisingly showed that the mean colonic transit time increased and the evacuation time did not change with the addition of dietary fiber [56].

Methods used for bowel care include anal massage (35%), unaided self-defecation with or without oral medication and abdominal massage (29%), digital stimulation (18 to 82%), rectal suppositories (15 to 74%), oral laxatives (29 to 41%), supplemental fiber (17 to 43%), manual evacuation (65 to 68%), and enemas (11 to 89%), [4, 9,10]. A sequential schedule of evacuating stimuli reduces total but not segmental transit times, decreases the need for oral or rectal laxatives, and increases the frequency of bowel movements [57]. The bisacodyl suppository is the most commonly used laxative, while docusate is the most popular oral agent [6]. Polyethylene glycol-based bisacodyl suppositories or polyethylene glycol-based, glycerine, docusate sodium miniemas result in a decrease in the amount of time for bowel care, a shorter total defecation period, and stimulation of bowel reflex defecation sooner than hydrogenated vegetable oil-based bisacodyl suppositories [58]. Digital stimulation of the anal canal serves to manually disimpact the rectum, to dilate the anal canal, and to relax the puborectalis muscle; the last effect decreases the anorectal angle and reduces outflow resistance to the passage of stool, resulting in left-sided colonic activity [59]. A gloved finger in the rectum inducing sustained pressure towards the sacrum, relaxes the spastic EAS and pelvic muscles [60]. Rotation of the finger continues the stimulation until a reflex peristaltic wave is generated in the rectum, resulting in passage of flatus and stool. The resulting RAIR relaxes the IAS, while the rectocloitic reflex stimulates pelvic nerve-mediated peristalsis. When the abdomen is massaged, following the course of the colon in an aboral direction, intraabdominal pressure increases [20]. Massage may improve abdominal distension, fecal incontinence, number of bowel movements, and mean total colonic transit time [61]. Megacolon may contribute to reduced responsiveness to both laxative therapy and physical interventions [62].

Saline enemas increase the proportion of continent stools and decrease the number of constipated stools although there is a significant drop-out rate due to stressful life events, inability to retain the balloon, and AH [63]. Due to the difficulty in retaining enemas from the loss of voluntary control of the EAS, the enemas should be placed above the rectosigmoid junction. With transanal irrigation, performed daily or every other day, a rectal catheter with an inflatable balloon holds the catheter in place while the enema is administered [64]. It empties the rectosigmoid and descending colon [64]. In a randomized, controlled trial where it was compared to conservative bowel management, transanal irrigation improves quality of life and the neurogenic bowel dysfunction score which includes frequency of bowel movements, time for defecation, frequency of digital stimulation, and fecal incontinence [64, 65]. After 3 years, however, the rate of success is only 35% [66]. The main reason for failure is leakage around the catheter, expulsion of the catheter, or bursting of the balloon [64].

The Malone Antegrade Continence Enema is an operation whereby the appendix is brought out to the skin forming an appendicostomy. Through the small stoma, patients can introduce a catheter and administer an enema that washes out the colon and rectum [67]. The overall success rate in
patients with mixed neurological diseases including SCI who were dissatisfied with conservative treatment was 74% and 87.5% after a mean follow-up period of 38 and 75 months, respectively [67,68]. Complications occur in 12.5% and 67% of patients, including wound infection, small bowel obstruction, and stomal stenosis.

Cisapride acts locally on motor activity by facilitating the release of acetylcholine at the myenteric plexus. In two cross-over trials, cisapride did not alter the number of bowel movements, ease of evacuation, or oro-anal transit time [69, 70], although one study showed improvement in stool consistency [70]. Cisapride has since been taken off the market due to ventricular arrhythmias. Prucalopride, a serotonin 4 receptor agonist with enterokinetic properties, reduces total colonic transit time with an increase in bowel movements but does not affect other bowel care related parameters [71]. Adverse events include flatulence, abdominal pain, diarrhea, and headache, causing some patients to discontinue treatment. Neostigmine, which increases cholinergeic tone by blocking the metabolism of acetylcholine by acetylcholinesterase, is used most commonly for bowel care in SCI patients was tested in a randomized, blinded design comparing intravenously neostigmine alone, neostigmine with glycopyrrolate, or normal saline [72]. Glycopyrrolate is an anticholinergic agent that reduces the side effects of bradycardia and bronchoconstriction caused by neostigmine with limited activity on the muscarinic receptors of the colon. Prompt bowel evacuation occurs with neostigmine alone or neostigmine with glycopyrrolate. In a subsequent double-blind, cross-over study, intramuscular neomycin/ glycopyrrolate given during three consecutive bowel evacuation sessions reduced total bowel evacuation times compared to placebo [73]. Trials are in progress to test the efficacy of intranasal neomycin and glycopyrrolate [74]. Fampridine, a selective potassium channel blocker, increases the number of days with a bowel movement in about one-fifth of the subjects in a double-blind randomized placebo-controlled trial, also improving spasticity as well as bladder and sexual function [75].

For sacral nerve stimulation, an anterior sacral root nerve stimulator is implanted at the level of the cauda equina. This is usually preceded by a test period using a percutaneous needle electrode [76, 77]. This method causes increased perianal sensitivity and improved deliberate retention of feces [77]. It affects the left colon increasing frequency of defecation [78, 79], decreasing time dedicated to bowel movements [79], decreasing fecal incontinence [80], and improving quality of life. There may or may not be an increase in rectal and anal pressures [80, 81] and anorectal measurements do not predict which patients will benefit from this treatment [82]. Colonic transit times do not change [78]. Anterior sacral root stimulators can be combined with an S2 to S4 posterior sacral rhizotomy which interrupts the reflex arc by cutting the posterior roots carrying the spasticity-causing sensory nerves. It results in improved bowel function and increased bladder capacity and emptying [83]. With elimination by rhizotomy of spinal reflexes that control continence and defecation, incontinence is no longer a problem. Although reflex defecation is then impossible, evacuation of stools is more predictably induced by manual evacuation or the use of anterior root stimulators.

Noninvasive magnetic stimulation, which produces an electrical field sufficient to depolarize nerves, increases anal and rectal pressures, decreases rectal volume [84], decreases mean colonic transit time, and improves bowel habits [85].

Surgical placement of a colostomy or ileostomy, the main indication being constipation, reduces the time spent in bowel care, increases independence, prevents fecal incontinence, relieves abdominal distension, enhances or maintains quality of life, and reduces hospitalizations for chronic bowel dysfunction by 70% according to pre-post or retrospective studies [86-89]. It provides a dry, clean environment so that pressure decubitus ulcers may heal. Most patients wished that the colostomy had been offered sooner, and none wanted it reversed [88]. Colon transit time and anorectal manometry can be used to identify the severity of bowel dysfunction and the site for the colostomy [89]. A left-sided colostomy with a Hartman’s pouch is probably the most common approach used [87]. Management of complicated decubitus ulcers and perineal and pelvic wounds is the primary indication in 43% of patients with left-sided colostomies. A right-sided colostomy is less likely to have problems with emptying but results in more watery stools, more frequent stoma care, and increased risk of leakage compared to a left-sided stoma. The long segment of defunctionalized colon predisposes to diversion colitis and mucous impaction which can be prevented by a distal colonic resection, a procedure that increases operative morbidity. The most common complications of colostomies are leakage of mucus per rectum, stomal prolapse or parastomal hernias, bowel obstruction, and wound dehiscence [86, 89].

**Gallbladder involvement in SCI**

Gallstones or previous cholecystectomy occur in 17-31% of SCI patients [90-92], some of whom have risk factors such as diabetes, obesity and/or family history of gallbladder disease [90]. The prevalence may be high even if patients with risk factors are excluded [91, 92]. Ultrasound examinations found gallstones in 25% or 26% of asymptomatic males compared to a control of 9% or 10% [93, 94]. Ultrasound exams performed within the first 6 months after SCI demonstrates a high incidence of biliary sludge (19%) but not gallstones (8%); all stones developed after four months and were small and not strongly echogenic. Cholelithiasis does not correlate with neurological level of injury, duration of SCI, age, obesity, or diabetes mellitus [94] although it may correlate with the severity of the SCI lesion [90].

The reason for the increase in gallstones in SCI is unknown. Gallbladder fasting volume is low, probably from impaired gallbladder filling due to impaired relaxation, generally with SCI lesions above T10 [95]. Ejection fraction
is low especially in lesions above T10 [95]; this may be due to low resting volumes rather than a defect in contractility. By ultrasound, only 38% have abnormal emptying fractions but this was associated with diabetes and obesity. Contraction of the gallbladder is probably mediated by the vagus nerve and cholecystokinin, both intact in SCI patients. Gallstones may also be due to gastroduodenal and colonic motility disorders seen in these patients [96], altering bile acid pool size and biliary lipid secretion [97]. Gallstones may be due to events that occur around the time of the injury, such as rapid weight loss [90]. While there is no association between gallstone formation and body mass index (BMI) in SCI, the BMI may underestimate the percentage of body fat due to osteopenia and loss of muscle mass from severe atrophy of limbs.

Due to the loss of normal sensory perception, the symptoms and signs of gallbladder disease may be atypical or even absent so that pathologic processes are often advanced at diagnosis [11]. Other authors indicate that the presentation is normal [98] and that atypical GI symptoms (bloating, nausea, and abdominal pain not found in the right upper quadrant or epigastrium) are not associated with gallstones [98]. Localization of pain was found in one study but in others only occurred in those with low SCI [11, 99].

Acute acalculous cholecystitis, found in 3.7% of patients with acute SCI admitted to the Intensive Care Unit of a referral hospital [100], presents with a right upper quadrant mass (100% of patients), steady fever (85.7%), tachycardia (100%), and cholestasis (85.7%). Predisposing factors include narcotics, mechanical ventilation with positive end-expiratory pressure, and hyperalimentation. Complications include narcotics, mechanical ventilation with positive end-expiratory pressure, and hyperalimentation. Complications include a gangrenous gallbladder and an inflammatory mass.

The need for cholecystectomy is normal if operations done prophylactically are removed from consideration (90). Although mortality rates are comparable to control individuals, SCI patients have a higher complication rate and a higher conversion from laparoscopic to open procedures. Some authors, whose threshold for elective cholecystectomy is low [90, 99], support routine screening ultrasounds to alert the clinician to the presence of gallbladder disease, while others believe that no specific interventions are warranted based on the findings.

Conclusion

A high index of suspicion is required to diagnose gastrointestinal involvement in SCI due to impaired sensory and motor functions.

Conflicts of interest

The author has no conflicts of interest regarding this article.

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