Proximal Jejunostomy With or Without Myectomy-Myotomy Modification in Five Infants With Total Intestinal Aganglionosis: An Experience With Surgical Treatments in a Single Institution

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Background: Total intestinal aganglionosis is characterized by the absence of intramural ganglion cells, in which the disease’s involvement extends from the stomach to the anorectum. This disease was suggested previously to be incompatible with life, but recently an extended small bowel myectomy-myotomy has achieved some prolonged survivors.

Methods: Five patients with total intestinal aganglionosis underwent laparotomy at 1 to 5 days of age. Surgery was performed as a simple jejunostomy 60 to 70 cm below the ligament of Treitz in the initial 2, jejunostomy 30 cm below the ligament of Treitz in 1, and jejunostomy with myectomy-myotomy modification 30 to 35 cm below the ligament of Treitz in the remaining 2 infants.

Results: The initial 2 patients died of sepsis, possibly derived from frequent enteritis and bacterial translocation at 7 and 8 months of age. Another patient had prolonged survival but died of hepatic failure at 1 year, 4 months. The remaining 2 children have survived beyond 2 years of age without any liver dysfunction, receiving a combination of enteral and parenteral nutrition.

Conclusions: The more proximal site (30 to 35 cm below the ligament of Treitz) of jejunostomy with myectomy-myotomy modification appeared to be preferable for prolonged survival in these 5 patients with total intestinal aganglionosis.


INDEX WORDS: Hirschsprung’s disease, total intestinal aganglionosis, myectomy-myotomy.

TOTAL INTESTINAL aganglionosis is an extremely severe form of enteric nervous system maldevelopment.1-2 The aganglionosis involves the entire colon, small intestine, duodenum and stomach.3 Treatments for infants with total intestinal aganglionosis previously resulted in uniformly fatal outcomes, but recently extended myectomy-myotomy of the proximal jejunum (Ziegler) has achieved some prolonged survivors.4-5 However, the extreme rarity of the disease appears to make it difficult to assess the significance of this recently recognized surgical alternative for extensive aganglionosis without biases from differences in institutional equipment, clinical experience, or the practice of parenteral nutritional support.

Here, we describe the results of the surgical treatments for 5 such patients in a single institution.

MATERIALS AND METHODS

We treated 5 patients who had total intestinal aganglionosis between 1990 and 2000. Of the 5 children, 4 were sporadic cases and 1 had familial occurrence. In each case, the diagnosis of congenital aganglionosis was made based on histopathology, and the levels of involvement of aganglionic segments were determined by intraoperative leveling biopsies from the stomach, duodenum, jejunum, ileum, colon, and rectum.

Case 1

A male newborn baby, delivered after a 34-week gestation with a 2,100 g birth weight, underwent a laparotomy at the age of 1 day. His elder sister had total colonic aganglionosis and had been treated in our hospital. Intraoperative biopsy specimens all showed an absence of intrarumal ganglion cells from the stomach to the rectum. We made a jejunostomy 70 cm distal from the ligament of Treitz and treated with total parenteral nutrition (TPN). The volume of ostomy outputs remained less than 10 mL per day, and he repeatedly showed severe enteritis. He died of sepsis at the age of 7 months. Two years later, his younger sister was born. She had total colonic aganglionosis with extensive involvement of the ileum. She was treated uneventfully with a modified Martin’s procedure.

Case 2

The second patient, delivered after a 39-week gestation with a birth weight of 2,540 g, underwent a laparotomy at the age of 3 days. Intraoperative histopathology of the gut wall biopsy specimens confirmed total intestinal aganglionosis. We performed a jejunostomy at a site 65 cm distal from the ligament of Treitz, and treated with TPN. The ostomy did not function, and his abdomen was persistently distended. He died of sepsis complicated with liver failure at 8 months of age.

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Case 3

The third patient, delivered after a 41-week gestation with a 3,884 g birth weight, underwent a laparotomy at 5 days of age. Histopathology confirmed total intestinal aganglionosis during surgery. In this case, we performed a jejunostomy at 30 cm distal from the ligament of Treitz. His early postoperative course was favorable, and enteral feeding was started using an elemental diet (20 mL of 8 times a day; Elental-P, Morinaga-Lucel Co, Tokyo, Japan). The volume of ostomy output measured between 50 and 100 mL per day. He gained weight up to 6,500 g until 7 months of age. However, his liver dysfunction worsened at 8 months, and he died of liver failure at 1 year, 4 months.

Case 4

The fourth patient, of 2,270 g birth weight, underwent a laparotomy at the age of 4 days. Histology findings showed a small number of ganglion cells at 3 cm from the ligament of Treitz; but the distal gut was completely aganglionic. A jejunostomy was made 35 cm below the ligament of Treitz. At 8 weeks of age, he underwent myectomy-myotomy modification of a 30 cm proximal jejunal segment. A 10-cm myectomy with 5-mm width starting at the histology-proven transition zone was coupled with a 20 cm antemesenteric myotomy of the subsequent jejunum. The depth of the myectomy/myotomy incision was kept within the submucosa, and the cut edge was spread to a width of 8 to 10 mm. The final length of the treated jejunum was 30 cm, and it ended as a stoma. Enteral feeding was soon started orally using breast milk and an elemental diet (20 mL at 8 times per day). The volume of enterostomy output measured between 50 and 110 mL. Antibiotics and medications for hepatic protection were used to prevent cholestatic liver disease after the treatment for parenterally fed infants with short bowel syndrome. His growth consistently remained between the 5th and 95th percentile of the standard for weight and height. He currently is in good condition at 8 years old using a combination of home parenteral nutrition and continuing oral intake of an elemental diet (Elental-P, Morinaga-Lucel Co.). He receives 20% of his calories by oral intake and the other 80% via the parenteral route, respectively. She consistently remains within the 5th to 95th percentile of the standard for height and weight. She has not shown any signs or symptoms of liver dysfunction.

Case 5

A female newborn baby, delivered vaginally after a 39-week gestation with a 3,698 g birth weight, was referred to our hospital because of abdominal distension and bilious vomitus. At laparotomy at the age of 4 days, histopathology confirmed total intestinal aganglionosis. We performed a jejunostomy 30 cm distal from the ligament of Treitz. On the 8th postoperative day, we performed a myectomy-myotomy for the proximal jejunal loop. A 10-cm myectomy with 5-mm width starting at the ligament of Treitz was coupled with a 20 cm antemesenteric myotomy. The depth of the seromuscular incision was the submucosa and the cut edge was spread to a width of 10 mm. The treated jejunum was 30 cm and ended as a stoma. Her postoperative course was favorable, and enteral feeding soon was started using breast milk (20 mL at 8 times per day). The volume of enterostomy output remained between 40 and 100 mL per day. During early postoperative care, she showed no episode of enteritis, and no prophylactic antibiotics were used. She currently is in good condition at 2 years and 8 months of age using a combination of parenteral nutrition and continuing oral intake of an elemental diet (Elental-P, Morinaga-Lucel Co.). She receives 60% of his calories by enteral feeding and the other 40% from the parenteral nutritional support, respectively. She consistently remains within the 5th to 95th percentile of the standard for height and weight. She has not shown any signs or symptoms of liver dysfunction.

RESULTS

Intraoperative Findings

All 5 patients underwent laparotomy aged 1 to 5 days. At the initial surgery, gut caliber changes were found at a site between 90 and 110 cm distal from the ligament of Treitz in each case (Fig 1). The average ratio of the proximal bowel caliber to the distal one was 3.01 (Table 1). However, a histology-based transition zone was confirmed only in case 4 and was only 3 cm distal from the ligament of Treitz. Therefore, there was no correlation of the observed transition with the presence or absence of intramural ganglion cells in these 5 infants.

Surgical Outcomes

In cases 1 and 2, proximal enterostomy was not successful. The volume of ostomy output remained less
than 10 mL per day. They suffered from intraluminal stasis and cholestatic liver disease. Liver function measurements progressively deteriorated after repeated enteritis, and they died of sepsis at 7 and 8 months of age, respectively. In case 3, the output volume from ostomy measured between 50 and 100 mL per day. He grew up to the age of 7 months with enteral feeding and parenteral nutrition support. However, his liver function worsened after enteritis at 8 months of age, which led to a fatal result at 1 year, 4 months of age. The remaining 2 patients with myectomized proximal jejunal loop survived beyond 2 years of age without any liver dysfunction. They receive a combination of enteral and parenteral nutrition, in which cases 4 and 5 receive 60% and 20% of their calories by the enteral route (oral intake), respectively. Both of them remain within the 5th to 95th percentile of the standard for height and weight after the age of 1 year (Fig 2).

**DISCUSSION**

The majority of enteric ganglion cells are derived from the neural crest and migrate into the intestinal wall in a caudal direction between the 5th and 12th gestational weeks.6,7 Total intestinal aganglionosis is a rare condition in which the neural crest–derived cell development is disrupted severely.5–10 This type of congenital aganglionosis is estimated to occur in less than 0.1% of patients with Hirschsprung’s disease.11,12 The extreme rarity of the disease makes it difficult to assess the detailed pathophysiologic conditions. The pathology of total intestinal aganglionosis may be different from that of near-total intestinal aganglionosis, in which the ganglonated bowel often extends distally from the ligament of Treitz.13 In total intestinal aganglionosis, a lack of peristaltic propulsion by the ganglonated bowel theoretically is expected.14 In such pathologic conditions, the intraluminal stasis of the gastric, bile, and pancreatic juice may be harmful to the sphincter function of the papilla of Vater and the surrounding region resulting in severe cholestatic liver injury or dysfunction. An extended myectomy-myotomy for the aganglionic proximal small bowel produced encouraging results in children with both total and near-total intestinal aganglionosis.3 However, the questions of whether and how surgery for total intestinal aganglionosis should be modified from that for near-total intestinal aganglionosis have not been answered clearly.

Here we described the results of surgical treatment for 5 infants with total or near-total intestinal aganglionosis, 4 of whom had no ganglonated bowel below the ligament of Treitz (Table 1). In the initial 2 patients, we performed a jejunostomy 60 to 70 cm distal from the ligament of Treitz during the neonatal period, but could
perform no additional myectomy-myotomy surgical intervention because of their deteriorating general condition. Severe enteritis and intraluminal stasis in the proximal jejunum could not be controlled because the proximal jejunal loop was too long to be effectively irrigated and also because antibiotics could not relieve the overwhelming bacterial overgrowth. In previous studies, a jejunostomy with myectomy-myotomy modification 50 to 70 cm below the ligament of Treitz was reported to function in several cases with near-total intestinal aganglionosis. However, we performed it 30 to 35 cm below the ligament of Treitz in the subsequent 3 infants because a jejunostomy 60 to 70 cm below the ligament of Treitz appeared to us excessively long for total intestinal aganglionosis. Theoretically, myectomy can relieve the resistance to the entry of luminal contents into that segment of the intestine, but the resistance is never zero and will cause obstruction if the segment is too long. At an aganglionic segment with a length greater than 20 to 40 cm, the small bowel passive conduit length would be too long to permit proximal propulsive forces to push the luminal contents through, resulting in a functional obstruction and intraluminal stasis.

In these 5 infants, we found a gut caliber change at a site between 90 and 110 cm distal from the ligament of Treitz. However, the histologically proven transition was confirmed only in case 4. There was no correlation of the observed transition with the distribution of enteric ganglia in these patients. Based on most previous descriptions, the presence of gut caliber change may be inexplicable in the totally aganglionic gastrointestinal tract. This unique phenomenon may be derived from a passive fluid retention in the proximal gut during embryogenesis, which was aggravated from fetal amniotic fluid ingestion and functional obstruction of the distal gut. Attention should be paid to possible discrepancies between macroscopic and microscopic transition zones because the presence of a gut caliber change does not always indicate a location for which an ostomy should be made in this rare disease.

The absorptive capability of the myectomized aganglionic bowel remains to be assessed in children with total and near-total intestinal aganglionosis. In cases 3 through 5, we decided to perform a jejunostomy 30 to 35 cm distal from the ligament of Treitz for patients with total intestinal aganglionosis after the difficult clinical course in the initial 2 cases. A more proximal jejunostomy, as performed in the current patients, may be advantageous for the control of enteritis and intraluminal stasis, but may lead to poor absorptive activity. In cases 4 and 5, who had a 30- to 35-cm jejunal segment myectomized, body weight and height both remain between the 5th and 95th percentile of the standardized curve (Fig 2). However, they can receive only 20% to 60% of their calories by oral feeding, and the parenteral nutrition support has not been discontinued. Considering the findings reported on infants with short bowel syndrome, a 30- to 35-cm proximal jejunal loop must be too short eventually to support adequate enteral absorption to predict a TPN-free related survival. Alternatively, a second myectomized 30 cm of more distal jejunileum with a second stoma, which could be fed, may be a feasible and attractive option for such patients. Pursuit of the potential to be fully nourished enterally in infants with total intestinal aganglionosis remains a challenge for pediatric surgeons.
tion of the aganglionic bowel segment, the patient’s growth, and possible TPN-associated liver injury should be investigated further based on a greater number of children with total intestinal aganglionosis.\textsuperscript{20,21} However, considering that the outlook for gut nutritional absorption is hardly optimistic in many patients with total intestinal aganglionosis, this disease entity remains a candidate for intestinal transplantation.

REFERENCES