Long-Segment Aganglionosis: a 15–year Experience

Ali Reza Mirshemirani1*, Sina Sadeghian, and Jafar Kouranloo

1 Department of Pediatric Surgery, Pediatric Surgery Research Center, Mofid Children's Hospital, School of Medicine, Shaheed Beheshti University of Medical Sciences, Tehran Iran

Received: 11 June 2007; Received in revised form: 25 Oct. 2007; Accepted: 12 Nov. 2007

Abstract- To access the surgical progress of patients with long segment aganglionosis and long-term evaluation. During the past 15 years, we reviewed the therapy and outcome of 15 patients with Long-Segment Aganglionosis (LSA) in Mofid children's hospital. 2 with Aganglionosis to the midtransvers colon (MTA), 8 to the ascendant and cecum (A/CA), 4 with total colonic and distal ileum involvement (TCA), and one child with total intestinal Aganglionosis (TIA). The patient with TIA had only laparotomy and staged biopsies. 12 were treated by Swenson technique and 2 patients by other procedures. All patients received total parenteral nutrition up to 10 days postoperative to ensure appropriate fluid and electrolyte status. The patients with TCA needed supplementary therapies (Diet, Drugs) up to 4 months to reduce the stool frequency. 2 patients died. The patient with TIA died 6 months after Biopsies. Another one died 5 months after pull-through: small bowel ileus with total intestinal necrosis. 9 Of the 13 alive patients have normal stool pattern: 1-3 times per day. 2 patients have 3-6 stools per day with 1-2 soiling per week. An adolescent patient has occasionally nightly soiling (1-2 per month). No actual data about one patient (A/CA). One TCA patient doing well immediately postoperative becomes obstipated 3 months later (after having Entrocolitis). He had washout enemas for 9 months. Now stools regularly. Awareness of long-segment Aganglionosis should lead to earlier diagnosis. Appropriate surgery and improvements in supportive care led to a significantly increased rate of survival. Patients with LSA even involving ileum (but no more than 50 cm of ileum) should attain a normal stooling behavior. Long-term outcome is quite favorable.

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Key words: Long-Segment Aganglionosis, total colonic aganglionosis, intestinal aganglionic.

Introduction

The incidence of Hirschsprung's disease (HD) was found to be between 1 in 4400 (1) and 1 in 7100 (2), live births with a male/female ratio of 4.1:1 in short segment, and 2.4:1 in long segment HD (2).

The incidence of LSD (Including total colonic Aganglionosis), in different studies, has a wide range of 5% to 20% from all HD patients (pts) (3-5). The breakthrough in the surgery of congenital megacolon occurred in 1948: Swenson and Bill identified the distal segment of the colon as a benign spastic lesion producing megacolon (6). They resected the rectum and the rectosigmoid with preservation of the sphincter and pulled through the normal colon (6). Although other procedure like Duhamel, Soave, Rehbein, (or the modification brought by boley, Martin or Kimura mostly for long-segments have been instituted to improve functional results, Swenson procedure proved to be as good as the other if the procedure is carried out correctly.

Patients and Methods

Mofid children's hospital records of 162 pts with biopsy-proven aganglionosis from 1990 to 2005 were reviewed. We found 15 pts with LSA: 2 with aganglionosis to the midtransverse colon, 8 to the ascendant and cecum, 4 with total colonic and distal ileum (15 to 45 cm) involvement and one child with total intestinal aganglionosis (TIA) extending to the duodenum (Figure 1,2) all but the last had pull-through procedure, and the pt with TIA had only laparotomy and staged biopsies.

There were 11 boys and 4 girls. All pts with TCA and the pt with TIA were boys.

10 pts were primarily operated (all TCA included) and the other 5 were reoprated in our hospital. From primarily operated pts, 6 were under one year of age, the other 3 pts were 2,3 and 5 years old at the time of pull-through. The PT with TIA had no pull-through operation but laparotomy and staged biopsies.

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Preoperative 5 pts had enterocolitis and 3 pts had ileus. The five reoperated pts were: one under one year, one 3, one 5 and two 7 years old. All these pts had the primary pull-through when they were younger than one year. Primary procedures used in these last 5 cases were Rehbein (N=3), Duhamel (N=1) and laparoscopic Swenson (N=1). The 3 pts formerly operated by Rehbein technique presented with aganglionosis rectum 6 to 10 cm proximal to dentate line. The pt operated by Duhamel technique had a 7 cm aganglionic rectum pouch with colorectal anastomosis 3 cm proximal to dentate line. The laparoscopic pull-through (MTA-PT) was complicated by anastomosis fistula with pelvic abscess and totally stricture of the anastomosis. Also in this case was found an aganglionic distal segment 20 cm long.

Before pull-through, 5 primarily treated pts (3 with TCA) had ileostomies and the resection of aganglionic bowel as a first operation followed by pull-through 2 to 6 months later. Also 2 from 5 reoperated pts had colostomies before pull-through. The following pull-through procedures were achieved: Swenson in 12 pts, Duhamel in one and a combined approach (laparotomy and posterosagittal) in one. The last procedure was achieved in the PT with primary laparoscopic pull-through. The reason for posterosagittal approach was the cicatricial tissue in the pelvis after abscess by anastomosis fistula.

Regarding only the TCA pts, 3 of them were treated by Swenson (total colectomy with ileum-rectum end-to-end anastomosis) and one by Duhamel.

The leaving rectum segment in these pts was 1 to 4 cm long.

Postoperative management consisted of total parental nutrition 5 to 10 days, followed by gradual introduction of oral feeding beginning with a low-osomolarity low-residue diet.

Three TCA pts needed loperamide, and 2 of them washouts with saline for 3 and 6 months respectively.

Results

The pts were followed-up 6 months to 15 years postoperatively. There has been no operative-related mortality in these series. Two pts died: the pt with TIA died 6 months after biopsies, and another pt died 5 months after pull-through by small-bowel ileus with total intestinal necrosis. There were two major and 8 minor complications in 15 surviving pts. The major complications occurred both after a secondary pull-through. They consisted in an anastomosis fistula 3 weeks postoperative needing a temporary ileostomy and a volvulus developed 3 years postoperative needing surgery.

As minor complications were registered subileus in one pt with spontaneous resolution, enterocolitis (one to three episodes) in 4 pts (2 TCA pts) and postoperative perineal skin irritation in 3 pts up to 1 year postoperative. Nine of 13 alive pts have normal stool pattern: 1-3 time per day. Two pts have 3-6 stools per day with 1-2 soiling per week. An adolescent pt has occasionally night soiling (1-2 per month).

There are no actual data about one pt (A/CAPT).

The pts with TCA needed supplementary therapies (diet, drugs) up to 4 months to reduce the stool frequency. One TCA pt doing well immediately postoperative becomes obstipated 3 months later, after having enterocolitis. Thereafter he needed washout enemas for 6 months. Now he stools regularly (3 time per day), and has no symptoms.

The levels of serum iron, serum ferritin, folic acid, and B12 vitamin were determine in all TCA pts. Two of them needed iron, B12 vitamin and folic acid supplementation up to 1 year postoperative. The ferritin level was normal in all pts.

Discussion

The major advance that have occurred in the management of hirschsprung’s disease include the definitive
management of the newborn, our understanding of hirschsprung’s-associating enterocolitis and the treatment of this entity, and the successful management of the very complex form of this disease, total colonic aganglionosis (7).

The most common surgical techniques used in hirschsprung’s disease are Swenson (introduced in 1948), Duhamel (introduced in 1956), Soave (introduced in 1963), the modification of the last one (Boley 1968), and recently transanal pull-through.

In total colonic Aganglionosis these operations have been performed without or with modifications: most known are Martin modification of Duhamel procedure, and Kimura procedure. Both of them preserve a part of the aganglionic colon aiming water resorption leading to the formed stools.

There is no agreement about the need to preserve 8-10 cm of aganglionic colon in TCA pts. In our opinion there are some arguments against preservation of the aganglionic patch: the aganglionic patch alternating the motility with subsequent stasis and obstruction, and postoperative enterocolitis seems to be a frequent occurrence in pts having a Patch of aganglionic Bowel.

There are four mainly causes leading to failure in surgical therapy of hirschprung’s disease:
• Incomplete resection of aganglionic segment.
• Ischemia of pull-through segment.
• Associated intestinal neuronal dysplasia.
• Postoperative enterocolitis.

In four of reoperated pts we found a high made anastomosis, more than 4 cm above the dentate line.

Ischemia may have a pathogenetic role in some cases of the acquired hirschsprung’s diseases.

Vascular compromise of the distal bowel segment at the time of the initial pull-through procedure may contribute to the selective loss of ganglion cells postoperatively because neural tissues are most sensitive to hypoxia.

This finding must be considered when recurrent obstructive symptoms develop following pull-through for aganglionosis (15).

The pts developing recurrent symptoms of hirschsprung’s disease, after adequate surgical correction, have to be fully studied with sequential biopsies in order to gain a better understanding of their condition (16).

The 5th reoperated pt (status post laparoscopic pull-through) had the anastomosis 1 cm above the dentate line.

The proximal end of the resected colon at first operation was normal in histological examination, by redo operation a 20 cm aganglionic colon was found, very probably due to intraoperative ischemia. The key of success in the therapy of hirschsprung’s disease is to pull-through a healthy bowel doing the anastomosis down within 1-2 cm to the dentate line.

The TCA pts we preserved a little more from aganglionic rectum (2 to 3 cm). Even in this situation the pts developed diarrhea, needing diet and loperamide up to 4 months postoperatively. One of 4 TCA pts becomes obstructed 3 months postoperatively, after having enterocolitis (17).

He had washout enemas for 6 months. However the long term results are good, showing us preservation of 2.3 cm of rectum in TCA pts obviates the need to use an aganglionic colonic patch.

The major postoperative complications occurred both in the pts having a second pull-through operation: anastomotic fistula leading to ileostomy (1 pt) and volvulus needing surgery (1 pt).

Two of these 5 pts had soiling 1-2 times per week.

All reoperated pts had aganglionic segments.

We did not find skip aganglionic area (18) in these children. In conclusion, awareness of long segment aganglionosis should lead to earlier diagnosis. Appropriate surgery and improvements in supportive care led to a significantly increased rate of survival.

Regularly followed up, early recognition and aggressive therapy of enterocolitis lead to good outcomes.

Patients with long segment aganglionosis even involving a part of ileum, should attain a normal stooling behavior. Long-term outcome is quite favorable.

Acknowledgement

The authors thank Mrs. P. Pourang and M. Saeedi for kind help and preparation of this manuscript.

References

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