Congenital Colonic Atresia: 4 Case Reports

Abstract

Introduction: Colonic atresia is a rare cause of neonatal bowel obstruction. Choosing between intestinal diversion and primary anastomosis for the management of this atresia is still discussed.

Methods and results: We reviewed all cases of colonic atresia managed in our center between 2000 and 2012. We have recorded, summarized and analyzed clinical features, associated anomalies, location and type of atresia, management and outcome. We collected four cases of colonic atresia handled in our department during this period, one case of type I and three cases of type III according to the Grosfeld classification. Three of the cases had at least one associated anomaly. Initial restoration of bowel continuity with a protective diversion was performed in one case, Delayed anastomosis following a primary intestinal diversion was performed in two cases, and primary anastomosis was performed in one case. Death occurred in one case following a post-operative sepsis.

Conclusion: Atresia of the colon is a rare etiology of the neonatal bowel obstruction. Colonic atresia can be associated with other congenital anomalies, which affect the prognosis. The prognosis of Colonic atresia in absence of other serious associations is good if proper management is carried out, with an overall mortality of 10%.

Keywords

Neonatal occlusion; Colonic atresia; Newborn; Surgery

Introduction

Colonic atresia is an uncommon cause of neonatal intestinal obstruction accounting for 1.8% to 15% of all gastrointestinal atresia [1]. Binninger reported the first case of colonic atresia in 1673 [2]. Gaub reported the first survivor in 1922, an infant with sigmoid atresia who was treated with delayed colo colic anastomosis after a primary colostomy. Potts [3] reported in 1947 the first survivor with an atresia of the transverse colon who was managed with a primary anastomosis. We performed an retrospective analysis of four cases handled in the department of pediatric surgery B of the children hospital of Tunis. The aim of this study is to acquire and present more detailed information about colonic atresia.

Patients

We searched for all cases of colonic atresia managed in our department from January 2000 to December 2012. Four cases were identified. Family and maternal history were unremarkable in all cases, and prenatal examinations were normal.

Case 1

A girl weighing 3,050 Kg was delivered with a gestational age of 41 week. At the 2nd day of life she was admitted for neonatal occlusion with abdominal distension, bilious vomiting, and failure to pass meconium. Physical examination revealed a distended abdomen, no associated anomalies was found. Abdominal X-Ray showed distended bowel, air fluid levels, with no air in the rectum (Figure 1). Surgical exploration was indicated and revealed a total colonic atresia (Type III) located at 3 cm below the Bauhin’s valve (Figure 2). Colostomy was made with proximal colic biopsy that indicate normal colic segment with presence of neuroganglionary cells, colic continuity was restored 6 months later with resection of a segment of the proximal dilated colon, with uneventful follow up.

Case 2

A 3,300 Kg boy was delivered after full term pregnancy; he was admitted in our department 48 hours after birth for neonatal occlusion. Physical examination revealed abdominal distension, abdominal X-ray showed distended bowel and air fluid level. Water-soluble contrast enema showed distal microcolon with an interruption of the colic continuity (Figure 3). Surgical exploration revealed type I colonic atresia, located at 10 cm below the Bauhin’s valve, primary colic anastomosis with protective intestinal diversion were performed. The colostomy was closed...
Case 2

5 months later. Seven months later, the patient was readmitted for cholestasis jaundice. Abdominal ultrasound showed dilated intrahepatic biliary ducts. An MRI with cholangiography showed a distended principal biliary duct with a free stone gallbladder. Surgical exploration for choledochal cyst was indicated and Bilio Intestinal anastomosis was performed after resection of the cyst, with uneventful follow up.

Case 3

A 2,050 Kg girl was was delivered after a full term pregnancy, on her first day of life she was admitted for neonatal occlusion. Physical examination showed torticollis with a limb malformation. Abdominal X-ray showed air fluid levels. Water-soluble contrast enema showed distal microcolon with an interruption of the colonic continuity. Surgical exploration for choledochal cyst was indicated and Bilio Intestinal anastomosis was performed after resection of the cyst, with uneventful follow up.

Case 4

A 2,080 Kg girl was delivered after a normal gestation. 3 days after birth she was admitted for neonatal occlusion. Surgical exploration revealed a type III colonic atresia of the transverse and left colon (Figure 4). Primary intestinal diversion was performed. The patient was discharged 9 days after the intervention. Restoration of colic continuity was performed 12 months later by an elective colo-colic anastomosis. 5 days later appeared abdominal distention with no stool emission, re intervention was then indicated, no leakage or anastomosis fistula was found. An intestinal diversion was performed. Histopathology results of intestinal biopsy showed absence of neuroganglionary cells and concluded to an extended colic form of Hirschsprung disease. Duhamel pull through was performed with resection of the aganglionosis segment. Follow up was uneventful and the patient is now 6 years old with normal feeding and transit. The type of atresia was determined according to the classification of Grosfeld et al. [4],[5] (Figure 5).

a. Type I is represented by a transluminal membrane or short atretic segment causing complete intestinal bowel obstruction, such as one of our cases.
b. Type II has the blind ending proximal bowel attached to the collapsed and undeveloped distal bowel by a fibrous cord along the edge of mesentery. There is no mesenteric defect.

c. Type III is similar to type II except that the fibrous connecting cord is absent and there is a V-shaped mesenteric defect, such as 3 cases of our study.

d. Type IV represents multiple segment atresias like a string of sausages or a combination of types I and III (Table 1).

Discussion

Atresia of the colon is one of the rarest intestinal atresia. There are several theories that attempt to explain the etiology of colonic atresia, the main one concerns the intra-uterine extrinsic mesenteric vascular obstruction, which can be caused by an internal hernia, a volvulus, an intussusception, or a strangulation in tight gastroschisis [3,6-9]. Association between colonic atresia and choledochal cyst as mentioned in one of our cases, has been reported, colonic atresia is due in these cases to an intrauterine direct pressure effect of the cyst on the reverse colon mesentery [10]. Intraluminal vascular factors have also been reported as an etiology of colonic atresia, Erskine proposed the possibility of emboli originating from the placenta and reaching the mesenteric circulation by bypassing the pulmonary circulation [11]. Fetal infection with varicella has also been described as one of the causes of colonic atresia [12-14], injury of the enteric plexus by varicella may lead to poor vessel development and induce ischemic conditions resulting into colonic atresia.

Familial cases of colonic atresia have been reported [15], but most of the reports were lacking detailed family histories. The incidence of associated anomalies in cases of colonic atresia is lower than the other intestinal atresia, however colonic atresia may be associated with various congenital anomalies, like abdominal wall defects such in gastroschisis where the intrauterine closure of gastroschisis around a segment of midgut may be an overlooked cause of isolated colonic atresia [16]. Other unusual associations reported were ophthalmic anomalies, bladder extrophy, exomphalos, vesico-intestinal fistula, as well as choledochal cyst [17]. The association of colonic atresia and Hirschsprung’s disease have been reported in 20 cases in literature [18,19], we report in this article the 21th case of Hirschsprung’s disease associated with colonic atresia. Literature has estimated that coexistent Hirschsprung’s disease is present in 2% of patients with colonic atresia [20], incidence of simultaneous colonic atresia and Hirschsprung’s disease is estimated to be 1 in 10 million live births. The mean age of the diagnostic of Hirschsprung’s disease when associated with colonic atresia is 8 months (ranging from 1 week to 5 years) [21], and was 10 months in our case. Hirschsprung’s disease is diagnosed after one or more unsuccessful procedures for atresia. After intestinal anastomosis, the child presented post-operative complications such as anastomotic dehiscence or functional intestinal obstruction requiring staged intestinal diversion followed by resection of aganglionic distal colon and a pull through procedure.

Newborns with colonic atresia present usual signs of distal bowel obstruction. Plain radiography of the abdomen reveals dilated bowel loops with air fluid levels; we can find a huge distended loop representing the colon proximal to the atretic segment. Water soluble contrast is helpful, but care must be taken because of the risk of distal colon perforation. In the type I of colonic atresia, a “wind sock sign” can be seen as the contrast pushes against the membrane [22-24], or Hook sign on the microcolon side of the colonic atresia in type III [25].

Classical surgical approach was primary anastomosis in the right colonic atresia proximal to the splenic flexure and colostomy with delayed anastomosis for the other location [1,25]. Primary anastomosis or primary anastomosis with intestinal diversion depending on the condition of newborn is considered as an efficient approach for the management of colonic atresia. However, primary anastomosis may be technically difficult because of the very large difference between the sizes of the distal and proximal bowels, and the loss of length associated with resection of dilated proximal colon [26]. Many authors have reported the resection and primary anastomosis as a reasonable treatment option regardless of the location of colonic atresia, if the newborn’s condition allows it [27].

Patency of the distal colon segment should be tested by retrograde or antegrade instillation of air and saline. Further management recommendations vary widely, proximal colostomy is well tolerated in infants and delayed reestablishment of intestinal continuity can be done later. The proximal and distal colonic ends adjacent to the atresia are abnormal in both innervation and vascularisation, that’s why in addition to resecting the bulbous proximal end, a portion of the microcolon should also be resected [27].

Rectal biopsy included in the initial surgical procedure seems to be an important step in the management of colonic atresia to avoid delayed diagnosis of Hirschsprung’s disease and the appearance of complications. The morbidity and mortality depend on several factors such as delay in diagnosis and treatment, nutritional status, sepsis, or associated anomalies [2]. Unlike atresia of the proximal small bowel, the diagnosis of colonic atresia is commonly delayed until feeding have been started ,then abdominal distension, fecaloid vomiting, and failure

Table 1: Summary of the cases (surgical exploration and associated malformations).

| Cases | Gender/Age (Hours) | Weight (kg) | Type of Atresia | Associated Malformations | Initial Procedure | Evolution |
|-------|-------------------|-------------|-----------------|--------------------------|------------------|-----------|
| 1     | F / 48 H          | 3.050       | Type I          | -                        | Colostomy        | Uneventful |
| 2     | M / 48 H          | 2.050       | Type III (Figure 4) | Choledochal Cyst     | Colic anastomosis protective colostomy | Uneventful |
| 3     | F / 24 H          | 3.300       | Type III        | Torticollis and limb malformation | Primary anastomosis | Death |
| 4     | M / 96 H          | 2.080       | Type III (Figure 5) | Hirschsprung’s disease | Colostomy        | Uneventful |

Citation: Chouikh T, Charieg A, Mnd C, Ghorbel S, Siada S, et al. (2014) Congenital Colonic Atresia: 4 Case Reports. J Pediatr Neonatal Care 1(3): 00018. DOI: 10.15406/jpnc.2014.01.00018
to pass meconium are noted. This delay results in more severe dehydration, sepsis, and higher mortality rate [28]. In the absence of small bowel atresia, the normal length of small intestine should allow normal bowel function [2,25]. If colonic atresia diagnosed early and if other major pathologies are absent, overall mortality should be 10% or less [27], early diagnosed and individually treated isolated cases of colonic atresia are not likely to have an unfavorable prognosis. Significant high mortality rates in cases operated after 72 hours seem to underline the importance of early diagnosis and treatment. The diagnosis of colonic atresia is still likely to be easily missed and the treatment be delayed. Early perinatal diagnosis and individualized operation are the key steps of the management of colonic atresia [18]. Treatment should be adapted for each patient based on clinical status and associated anomalies to give optimal results with less morbidity (Table 1).

Conclusion

Atresia of the colon represents the least common cause of neonatal intestinal obstruction. The choice of the operative therapy in colonic atresia should be individualized, based on the condition of the patient and the presence of associated anomalies.

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Citation: Chouikh T, Charieg A, Mnd C, Ghorbel S, Siada S, et al. (2014) Congenital Colonic Atresia: 4 Case Reports. J Pediatr Neonatal Care 1(3): 00018. DOI: 10.15406/jpnc.2014.01.00018