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Case report

Atypical presentations of alimentary tract duplications in children: two case reports

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Alimentary tract duplication is a rare congenital anomaly with an incidence of 1: 4,000 - 5,000 live birth and could occur anywhere along the gastrointestinal tract.\textsuperscript{(1)} The duplication was classified into 2 types, namely: cystic (80.0\%) and tubular (20.0\%). The former does not communicate with the adjacent lumen while the later might communicate and share the common adjacent lumen with the main alimentary tract. Ileum is the most common site (33.0\%) followed by esophagus (20.0\%) and colon (13.0\%).\textsuperscript{(2)} Clinical presentations are varied and determined by the type, site and size of the duplications.\textsuperscript{(3-5)} The preoperative diagnosis is usually difficult to establish especially in atypical presentations, due to the rarity of the anomaly. We herein report two cases of alimentary tract duplications in which clinical manifestations occurred beyond infant period, i.e., a girl with chronic constipation and a boy with acute intestinal obstruction.

Keywords: Case report, alimentary tract duplication, children, constipation.

Case presentation
Case report 1
Chief complaints
An 8-year-old girl with double inlet left ventricle post pulmonary artery banding and Blalock Hanlon procedure presented with acute abdominal pain and vomiting for 2 days during the regular visit at our hospital.

History of present illness
She had recurrent abdominal pain and chronic constipation for 4 years. The colicky pain had been worsened and she developed vomiting during the last 48 hours.
History of past illness
The patient had double inlet left ventricle which underwent pulmonary artery banding and Blalock Hanlon procedure.

Physical examination
The examination reveals mild abdominal distension, normo-active bowel sound with palpable, movable and soft-consistency mass size 6 x 8 cm² at the lower mid abdomen. Per rectal examination reveals empty rectum and minimal non-bursting soft yellowish feces without glove-like sensation implying that neither fecal impaction from functional constipation nor Hirschsprung’s disease was likely.

Imaging examinations
Abdominal radiography showed the localized dilatation of the descending colon with fecal impaction. But there was no proximal colonic nor small bowel dilatation. (Figure 1A). Disimpaction with polyethylene glycol and rectal enema initially failed but was finally successful by high rectal irrigation with glycerin-containing saline. Barium enema was done and revealed two colonic lumina from the ascending colon down to the distal descending colon, separated by the radiolucent line (white arrows) (Figure 1B). There was poor emptying function of the dilated part of descending colon and fairly good emptying function of the rectosigmoid colon after evacuation. The colonic transit study was performed with prolonged segmental and total colonic transit time (58 hours) (Figure 1C). To identify problematic segment, colonic manometry was studied. A colonoscopy with colonic manometry catheter insertion was performed. Colonoscopy showed two opening lumens of the colon at 38 cm from anal verge likely true diverticulum (Figure 1D). Colonic manometry study showed the normal colonic function with frequent high amplitude propagation contractions (HAPCs) after bisacodyl stimulation (Figure 1E). Further imaging with abdominal and pelvic computed tomography (CT) to evaluate this anatomical abnormality confirmed two laminae of the descending colon which one was dilated whereas the other was not. Moreover, there were two appendices at the right lower abdomen (Figure 1F).

Final diagnosis
The final diagnosis of this case was tubular duplication of nearly the entire colon.

Treatment
Exploratory laparotomy was performed to find a tubular duplication from descending colon to terminal ileum (Figure 1G). The descending colon was dilated and the involved ileum was 34 cm from ileocecal valve with normal caliber. There were two appendices and two Meckel diverticula. Left hemicolecction was performed followed by dividing common wall of transverse colon with surgical staplers. After spatulating the transverse colon’s distal end, primary end to end anastomosis with native sigmoid colon was done.

Outcome and follow-up
The postoperative course was uneventful. The patient started enteral feeding on postoperative day 7 and was discharged on day 10. At 18-months post-operation, the patient was very well without the symptoms of recurrent abdominal pain and constipation anymore.

Case report 2
Chief complaints
A previously healthy 13-month-old boy was referred from a private hospital due to suspected small intestinal obstruction.

History of present illness
He was a full-term baby of an uncomplicated pregnancy and well-grown since birth.

History of past illness
He was hospitalized due to 4 times of non-bilious vomiting and poor intake. During admission his abdomen became distended and he developed bilious vomiting. There was no history of constipation, diarrhea, or bloody stool.

Physical examination
On examination, the child looked sick without a fever. His abdomen was soft and distended. Bowel sounds were present and hyperactive. Per rectal examination showed yellowish stool without blood.
Figure 1. An 8-year-old girl with history of recurrent abdominal pain and chronic constipation for 4 years. 

(A) Anteroposterior supine abdominal radiography showed the localized dilatation of the descending colon with fecal impaction. But there was no proximal colonic nor small bowel dilatation. 

(B) Single contrast barium enema revealed two colonic lumina from the ascending colon down to the distal descending colon, separating by the radiolucent line (white arrows). 

(C) The distribution of sitz marks after 3-day ingestion. The segmental colonic transit time of right, left, rectosigmoid colon were calculated to 30, 16, 12 and 58 hours, respectively. 

(D) Colonoscopic view demonstrated the two colonic lumens of native colon and the true colonic diverticulum that shared the one-side colonic wall with the native colon. 

(E) Colonic manometry tracing showed high amplitude propagation contractions after bisacodyl stimulation (0.2 mg/kg/dose). 

(F) Axial post contrast enhanced CT of lower abdomen demonstrated two lumina of the descending colon, one was dilated (C1) and another was not dilated (C2). Moreover, there were two appendices at the right lower abdomen (white arrows). 

(G) Tubular duplication from descending colon to terminal ileum.
**Imaging examinations**

Abdominal radiograph revealed gasless abdomen pattern. There are a few abnormal small bowel loops were dilated at the mid and left upper abdomen (white arrows) (Figure 2A). Ultrasound abdomen showed a cystic lesion with the gut signature wall in mid-abdomen (Figure 2B), measured about 4.7 x 4.3 cm². The later abdominal and pelvic CT demonstrated the cystic lesion in mid-abdomen (Figure 2C) which compressed on the adjacent small bowel loop and caused abnormal dilatation of the multiple proximal small bowel loops with collapsed terminal ileum, entire colon, and rectum (Figure 2C).

**Final diagnosis**

Distal ileal obstruction due to cystic lesion was diagnosed in this patient.

**Treatment**

Emergent exploratory laparotomy was performed and revealed cystic duplication in the mesentery of the terminal ileum (10 cm from ileocecal valve). The duplication cyst was about 4 cm in diameter compressing the adjacent ileal lumen (Figure 2D). The residual bowel was in good condition. Segmental ileal resection with primary end to end anastomosis was done.

**Outcome and follow-up**

The patient improved and started enteral feeding on postoperative day 4. He was discharged on postoperative day 6. At 13-month follow-up, the patient was well and active.

**Discussion**

Alimentary tract duplications are rare congenital abnormalities that could occur in any area from the mouth to the anus. The pathogenesis of enteric duplication is unclear. Several theories propose the arising of the duplications. However, no single hypothesis can explain all sites and types of the anomaly. (2, 6) Duplications must be composed of 3 components, including continuity or adherence to a part of the alimentary tract, having a smooth muscle wall, and containing a mucosal lining of one or more cell types found in the alimentary canal. (2)

Our first case was a girl who presented with recurrent abdominal pain and chronic constipation that finally diagnosed long tubular colonic duplication. Duplication of the colon is extremely rare and first described in 1876. (7) The studies that indicated the high association of tubular colonic duplication with other abnormalities especially lower genital, urinary tracts, and lumbosacral vertebra. To the best of our knowledge, only 2 previous cases of the tubular colonic duplications associated with congenital heart diseases were reported. One was a girl with Tetralogy of Fallot, agenesis of the left kidney and ureter, and S-1 spina bifida occulta. (8) Another was a male infant with extensive vertebral defects, anal atresia, cardiac defects, tracheo-esophageal fistula, renal anomalies, and limb abnormalities.(VACTERL)-associated anomalies, namely, hemi-vertebrae, anal atresia with a recto-urethral fistula, hypospadias and common AV canal defect. (9) Unlike the present case, these two cases had entire colon duplication but they also had other obvious abnormalities as clues for the early investigations. As functional constipation and fecal impaction might be explained, the chief complaint of recurrent abdominal pain in the general population. Per rectal examination that showed “empty rectum” in the present case could not explain the functional cause of fecal impaction, hence mechanical obstruction from other causes should be sought out. However, it is difficult to diagnose colonic duplication before surgery, and misdiagnosis is common. (10, 11) Despite of imaging studies, some duplication cysts are diagnosed intraoperatively (38.0%). (12) Barium enema could be challenging to confirm the diverticulum if two colons’ opacification is overlap or in the same plane. Like our present case, the two colonic lumina from the ascending colon down to the distal descending colon, separating by the radiolucent line from barium enema study was retrospectively identified by an experienced pediatric radiologist later. Yousefzadeh DK, et al suggested using contrast media of two different densities during simultaneous evaluation of both colons in case suspected of colonic duplication. (7, 9) A colonoscopy might be valuable if there is the communication between the anomalous, and actual enteric lumens and CT whole abdomen is considered the most useful for the diagnosis preoperatively. (13) In the present case, the enlarged descending colon without any obvious mechanical obstruction from barium enema and the prolonged colonic transit time from colonic transit study misleading us to investigate more for colonic motility, and we accidentally diagnosed colonic diverticulum from colonoscopy. Again, to the best of our knowledge, this is the first case of an entire colonic diverticulum who received colonic physiologic study. The result showed normality of the entire colonic function, even
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Figure 2. A previously healthy 13-month-old boy with suspected small intestinal obstruction. (A) Anteroposterior supine abdominal radiography revealed gasless abdomen pattern with a few abnormal small bowel loops dilatation at the mid and left upper abdomen (white arrows). (B) Abdominal ultrasonography showed a cystic lesion with the gut signature wall lesion in mid-abdomen. (C) Two coronal images of post contrast enhanced abdominal and pelvic CT demonstrated the cystic lesion in mid abdomen (*) with abnormal dilatation of the multiple proximal small bowel loops and collapsed terminal ileum down to rectum. (D) Cystic duplication compressed adjacent ileal lumen.
long standing chronic obstruction that implied the intact enteric nervous system and intestinal muscle in patients with chronic intractable constipation or obstruction. In aspect of surgical treatment, to preserve the normal colon and intestine, left hemicolectomy with dividing common wall at transverse colon and primary end-to-end anastomosis to native distal sigmoid colon were fashioned. As a result, there was a residual duplicated descending, transverse colon and small intestine which possibly contains ectopic gastric mucosa, especially at 2 Meckel’s diverticula. Although the best option of treatment is the complete surgical removal of the duplicated part. Other considerations might be concerned, especially in this very young child with the long tubular duplication that involved the whole colon and terminal ileum. Complete surgical resection of the whole lesion as in one step might affect her nutrition status and the final long-term growth. Hence, we gave parental information of concerning symptoms, especially lower gastrointestinal bleeding and cancer surveillance as some evidence of a higher risk of adenocarcinoma of the residual duplication. Furthermore, long-term follow-up and careful monitoring are needed.

The second case, extrinsic compression by cystic duplication located at the terminal ileum resulted in intestinal obstruction. The diagnosis was missed due to a low index of suspicion. After a retrospective review of the ultrasonography with an experienced pediatric radiologist, the picture demonstrated the double layered wall, the so-called “gut signature” This finding consistent with located adjacent to the bowel wall should prompt the suspicious of duplication cyst. CT typically is not required to evaluate the duplication cyst. As the diagnosis could not be determined, CT could help us exclude some common causes of obstruction, including intussusception, ruptured appendicitis, and Meckel’s diverticulum with complications. CT may depict the location, extent of the cyst, and complications. Enteric duplications can present with gastrointestinal bleeding if the lesions contain gastric mucosa. Some patients are asymptomatic but the incidental finding was revealed during obtaining intra-abdominal workup for other purposes. Since the symptoms can overlap with other gastrointestinal abnormalities, it makes the diagnosis more difficult.

Conclusion
Alimentary tract duplications are rare and could present asymptomatic or with diverse manifestations. The high index of suspicion is needed, especially the clinical signs and symptoms that could not be served other more common diseases. Imaging studies are suggested as contrast study and CT can raise a clue for tubular duplications as well as ultrasound can help diagnose cystic duplications. A colonoscopy might be helpful, only the duplication with opening orifice to the main alimentary tract. Although the definite diagnosis could be made intraoperatively, clues from those investigations could help surgeons for the well-planned operation. Options of surgical procedure depend on the locations, types and involved structures. If possible, complete surgical resection is the treatment of choice.

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