Acquired Jejunal Atresia in a 2-Month-old Infant

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ABSTRACT
Acquired intestinal atresia in non–necrotizing enterocolitis affected patients is very rare. We report a 2-month-old male infant who presented with bilious vomiting, abdominal distension and constipation. He was exclusively breast fed, gained weight and was asymptomatic till six weeks of age. Exploratory laparotomy revealed blind-ended, dilated proximal segment and collapsed distal segment of jejunum at approximately 30 cms from duedeno-jejunal flexure, with a V-shaped mesentery defect resembling a type III-a congenital jejunal atresia.

CASE REPORT
A 2-month-old male infant was admitted with bilious vomiting, abdominal distension and constipation that were ongoing for 2 weeks. The child was born to a 24-year-old mother by spontaneous vaginal delivery at 38 wk of gestation, weighing 2400g. The pregnancy and delivery were uneventful. There was no evidence of intra-uterine ischemic insult including placental abruption, preeclampsia, intrauterine growth restriction, chorio amnionitis or placental infarction. There was no history of birth asphyxia. The child passed meconium within the first 6 h of birth, and his stooling pattern was normal until 6 wk of age. He was exclusively breast fed, gained weight and was asymptomatic till 6 wk of age.

Physical examination revealed a male infant who was pale, anicteric, and severely dehydrated. He had a soft and distended abdomen with visible peristalsis. The laboratory studies including complete blood count, serum electrolytes, and renal function tests were normal. An erect abdominal plain radiograph revealed dilated bowel loops, multiple air fluid levels and flecks of calcification in the right lower quadrant [Table/Fig-1a]. An upper GI contrast study done at the referring hospital revealed dilated small bowel loops with non visualization of colon even in delayed images [Table/Fig-1b]. A barium enema study revealed normal sized colon visualized till the level of ascending colon. A working diagnosis of healed intrauterine meconium perforation with postnatal adhesive obstruction was made.

Exploratory laparotomy revealed dense inter-loop adhesions with foci of calcification on the small bowel serosa. Adhesiolysis revealed blind-ended, dilated proximal segment and collapsed distal segment of jejunum at approximately 30 cms from duedeno-jejunal flexure, with a V-shaped mesentery defect resembling a type III-a congenital jejunal atresia [Table/Fig-2]. Jejunal resection and end-to-end anastomosis was performed.

Histopathological examination revealed chronic inflammatory changes. Normal gastrointestinal motility was achieved in three weeks. The patient had an uneventful course and was discharged on the 8th postoperative day.

DISCUSSION
Strictures and occasionally atresia of the bowel are known to occur in premature infants after recovery from necrotizing enterocolitis (NEC). This is particularly true in the segment of the intestine distal to an established enterostomy, most commonly the colon [1,2]. It is postulated that passage of stool through a partially damaged intestinal segment will render some protection against the development of strictures or atresia, whereas diversion of the fecal content will cause mucosal adherence and stricture or atresia formation [3].

Michaelis et al., reported a case of a 19-month-old boy in whom intestinal atresia developed after drainage of a large intra-abdominal abscess [4]. The proximal segment fistulized to the umbilicus which eventually healed without surgical intervention. Two months later at the time of surgical exploration, complete atresia with a V-shaped gap defect in the mesentery was noted. Sigge and Wurtenberge reported a 1430 gm premature infant in whom NEC developed in the 1st week of life and who 2 wk later was found to have a 7-mm mid-ileal atresia [5]. Jona described a similar case in a premature low-birth-weight neonate who developed ileal atresia as a sequel to NEC and intestinal continuity was established via an ileo-ileal fistula [3].

Acquired intestinal atresia in non–necrotizing enterocolitis affected patients is very rare. We could find only four case reports of acquired atresia of the small bowel in non-NEC patients in the English literature [6-9]. Puvabanditsin et al., reported a case of postnatal intussusception that caused jejunal atresia in an infant born at 26 wk of gestation [6]. Elemen et al., reported acquired ileal atresia in a 2-year-old girl because of chronic entrapment of the ileal segment together with its mesentery by an adhesive band interrupting the blood supply resulting in a V-shaped mesenteric defect between the referring hospital revealed dilated small bowel loops with non visualization of colon even in delayed images [Table/Fig-1b]. A barium enema study revealed normal sized colon visualized till the level of ascending colon. A working diagnosis of healed intrauterine meconium perforation with postnatal adhesive obstruction was made.

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CONCLUSION

The acquired atresia in our patient may be because of chronic entrapment of the jejunal segment together with its mesentery by an adhesive band interrupting the blood supply resulting in a V-shaped mesenteric defect between the atretic segments. Local ischemia and ongoing inflammation with further resorption of the devitalized segment and subsequent healing were likely involved resulting in complete healing of the atretic ends. We speculate that the reason for adhesions and calcification would have been a healed intra-uterine meconium perforation.

This brief report suggests an adhesive band, which may entrap a jejunal ileal segment together with its mesentery, may be a potential cause for acquired small bowel atresia in non-NEC patients.

REFERENCES