Distended neoesophagus in a man with congenital esophageal atresia

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Abstract: A 33-year-old man presented with abdominal fullness, chest tightness, and weight loss. He had undergone esophageal reconstruction at age 2 for congenital esophageal atresia. Sequelae of substernal interposed ileocolon gradually developed, including narrowing of cologastrostomy, subxiphoid compression of neoesophagus, redundancy of neoesophagus, and intestinal adhesion. In order to solve the aforementioned problems, he underwent revision surgery including enterolysis, release of subxiphoid compression, revision of cologastrostomy, and creation of colojejunostomy. Following the revision surgery, the alimentary tract achieved good patency and the neoesophagus yielded shrinking. He was discharged uneventfully on postoperative day 14.

Keywords: Congenital esophageal atresia; neoesophagus; esophageal reconstruction

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Introduction

A few cases have been reported requiring revision surgery more than 20 years after colonic interposition for congenital esophageal atresia. Revision surgery may relieve symptoms and improve quality of life. Further reconstructive options may be limited due to the lack of an available conduit (1). Here we report an interesting case with sequelae of esophageal reconstruction requiring complete preoperative evaluation and delicate revision surgery.

Case presentation

A 33-year-old married man presented with abdominal fullness and chest tightness. He was diagnosed as congenital esophageal atresia a few days after birth. The young man had undergone a total of 6 times of surgeries to solve the problem of congenital esophageal atresia at a medical center in northern Taiwan. In the newborn period, he underwent division of tracheoesophageal fistula, cervical esophagostomy, and gastrostomy. Subsequently, esophageal reconstruction was performed using ileocolon via retrosternum at age of 2. In recent 2 years, he had intermittent abdominal pain, abdominal fullness, and chest tightness, and lost 5 kg of body weight. Chest film showed marked dilatation of the neoesophagus with compression of right lung. Endoscopy showed stenosis of cologastrostomy. He searched our thoracic service for revision surgery. His body height was 169.8 cm, and body weight was 49.3 kg. Physically, the abdominal wall presented with irregular wound scar that related to previous wound infection. Laboratory data were all within normal limits. Computed tomography (Figure 1) revealed a distended neoesophagus (C-1) extended to abdominal cavity (C-2) and gastric dilatation (G). According to the preoperative evaluation, some sequelae of esophageal reconstruction were considered such as narrowing of colon-gastrostomy, subxiphoid compression of neoesophagus, redundancy of neoesophagus, and severe intestinal adhesion. To solve these problems, he underwent laparotomy with enterolysis, release of subxiphoid compression, revision of cologastrostomy (A1), and creation of colojejunostomy (A2)
At surgery, partial obstruction of the pylorus was detected by endoscopy and severe intestinal adhesion was confirmed. Following the revision surgery, chest films (Figure 3) showed the distended neoesophagus (left) yielded shrinking (right). Patient could drink water on postoperative day 10 and liquid diet on postoperative day 12. He was discharged on postoperative day 14 after examination of upper gastrointestinal series. He resumed regular diet on postoperative day 16 and was well in the out-patient-clinic follow-up.

**Comments**

In our experience of esophageal reconstruction, the common use of the esophageal substitute was ileocolon. Obstruction or redundancy of the neoesophagus was very rare (2). We have reported 14 cases requiring surgical revision for proximal anastomotic strictures after hypopharyngocolostomy or esophagocolostomy. These proximal strictures can be surgically corrected after excision of the scar and mobilization of the esophageal substitute through a cervical incision only or a cervical incision plus sternotomy (3). Some researchers reported a few patients requiring reoperation for late obstruction due to volvulus or kinking because of redundancy (4). Comparing to the proximal anastomotic stenosis, occurrence of the distal anastomotic stenosis is rare. Primary direct anastomosis is feasible in patients with congenital esophageal atresia with tracheoesophageal fistula. However, in case of without tracheoesophageal fistula required esophageal reconstruction because esophageal long gap (5). In the present case, he underwent 6 times of surgeries including esophageal reconstruction in the infancy because of unsuccessful primary anastomosis. Some authors emphasized that long-term follow-up of patients who underwent esophageal replacement in children is essential because of gradual changes in the function of the graft, strictures at the anastomosis and the unknown risks of Barrett’s esophagus (6). We kept this patient in the hospital until post-operative day 14 because we were concerned about the operative finding of intestinal adhesion.
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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Informed Consent: Written informed consent was obtained from the patient for publication of this manuscript and any accompanying images.

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