CASE REPORT

Trilogy of foregut, midgut and hindgut atresias presenting in reverse order

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SUMMARY

We present a case of triple atresias (foregut, midgut and hindgut) with multiple congenital anomalies presenting as imperforate anus. Abdominal radiography showed the double bubble sign. Upper gastrointestinal study through a nasogastric tube confirmed duodenal obstruction. Exploratory laparotomy, duodeno-duodenostomy for duodenal atresia and a left descending colostomy for anorectal malformation were performed. During extubation, the nasogastric tube came out and could not be reinserted by the anaesthetic team under laryngoscopic guidance. A chest radiograph showed the tube curled in the upper pouch. Bronchoscopy and oesophagoscopy confirmed oesophageal atresia (OA) with a distal tracheoesophageal fistula (TOF). The patient underwent right-sided extrapleural thoracotomy and division of the fistula with primary repair of OA uneventfully. Triple gut atresias presenting in reverse order with multiple anomalies is rare and passage of a nasogastric tube into the stomach in the presence of OA+TOF is exceptional. Alimentary tract obstruction should be corrected in proximal to distal direction.

BACKGROUND

A trilogy of gut atresias involving all three embryological segments of the alimentary tract in the same patient associated with multiple congenital anomalies but no chromosomal or genetic abnormalities is very rare.1 We report such an exceptional case which presented in reverse order and also had an exceptional nasogastric passage to the stomach delaying the diagnosis and treatment of oesophageal atresia (OA) and tracheoesophageal fistula (TOF).2–4

CASE PRESENTATION

A term baby girl had been born by elective lower segment caesarean section for breech presentation. Multiple abnormalities had been detected antenatally. These were hydrocephalus, phocomelia with oligodactyly, atrio-ventricular septal defect (AVSD) and intra-uterine growth retardation.

The baby was born in good condition, was cardiovascally stable and breathing in air. Meconium-stained liquor was noted at delivery but no meconium was noted to be passed following delivery. She did not pass meconium in the first 24 h and no anal opening could be seen in the perineum but urethral and vaginal openings appeared normal.

The baby had excessive salivation and some frothing, so an associated OA with a TOF was suspected. With some difficulty an attempt was made to pass a nasogastric tube into the stomach and the radiograph confirmed it was in the abdomen in the region of the stomach. However, there was a double bubble appearance with a possibility of duodenal atresia or associated malrotation and midgut volvulus so the patient was urgently referred to us after stabilisation.

The patient was referred to the paediatric surgical team as having an imperforate anus with a possible rectovestibular fistula, and transferred from the district general hospital. On examination, she had a urethral orifice and a vestibular orifice but no separate anus. The abdomen was soft, non-tender and scaphoid with epigastric fullness, but no mass was palpable.

INVESTIGATIONS

Chest and abdominal radiographs showed a small amount of gas in the stomach but no gas in the rest of the gastrointestinal tract (figure 1). It was unclear whether this was an atresia or a malrotation/volvulus. An upper gastrointestinal contrast study under fluoroscopy was suggestive of a possible midgut malrotation and volvulus (figure 2). A cranial ultrasound scan showed lobar holoprosencephaly and an abdominal ultrasound scan showed bilateral small dysplastic kidneys. An echocardiogram was performed which showed a complete AVSD with an additional secundum atrial septal defect. There was a moderate sized ventricular septal defect with a dilated coronary sinus.

Other abnormalities were as follows: microtia with absent external auditory meatus, complete AVSD, OA, TOF, duodenal atresia, anal atresia, bilateral radial ray abnormalities with oligodactyly, crowded mid-face features, and bilateral radial clubbing of the hands with only three digits.

A genetics review was performed by the clinical genetics team. The infant was the second child of consanguineous, Bangladeshi parents. Multiple congenital abnormalities identified in pregnancy were consistent with a diagnosis of trisomy 18. However this constellation of syndromes would also fit a VACTERL-D association. Therefore, an urgent fluorescence in situ hybridization (FISH) study for chromosome 18 trisomy was performed which was normal.

DIFFERENTIAL DIAGNOSIS

Anorectal malformation with a rectovaginal fistula with either duodenal atresia or midgut malrotation with volvulus neonatorum with multiple congenital anomalies with a possible VACTERL-D association.
was considered. The possibility of chromosomal anomaly of trisomy 18 and the Martinez-Frias syndrome were considered before the clinical genetic team review and investigations.

**TREATMENT**

Following investigations and resuscitation, the patient underwent an exploratory laparotomy, a duodeno-duodenostomy for duodenal atresia and a left descending colostomy for anorectal malformation. During extubation, the nasogastric tube came out and could not be reinserted even by the anaesthetic team under laryngoscopic guidance. Chest radiograph showed the tube curled in the upper pouch. Bronchoscopy and oesophagoscopy confirmed OA with distal TOF. The patient underwent right-sided extrapleural thoracotomy, division of the fistula with primary repair of OA and a gastrostomy uneventfully.

**OUTCOME AND FOLLOW-UP**

The patient was transferred to the neonatal intensive care unit post-operatively. A number of discussions were held with her parents by consultants and the family liaison nurses. In summary, the infant had severe underlying abnormalities, multiple intestinal atresias, with a complete AVSD and holoprosencephaly. This constellation of abnormalities was considered incompatible with survival. Her parents understood the very poor prognosis in view of the multiple severe congenital anomalies involving all major body systems and the prospect of major

**Figure 1** Chest and abdominal radiographs. Note the nasogastric tube (NGT), proximal pouch shadow and small gastric air bubble with the rest of the abdomen gasless.

**Figure 2** Upper gastrointestinal contrast study showing complete duodenal obstruction beyond the duodenal cap. NGT, nasogastric tube.

Reference:
invasive recurrent surgery in order to attempt to address the infant’s problems as listed above. It was felt that withdrawal of intensive care treatment would be in her best interests. Therefore, following family visits she was held in her mother’s arms, the endotracheal tube was removed and death was pronounced. A death certificate was issued, with the cause of death as multiple congenital abnormalities (probably VACTERL) and multiple alimentary tract atresias (which were operated on, with the infant recovering from anaesthesia). The staff extended their deepest sympathy to her parents at this most difficult time and offered an appointment for bereavement follow-up in the future.

DISCUSSION

Alimentary tract atresias involving the foregut, midgut and hindgut in the same patient are very rare and it is presumed that the embryopathogenesis of each is different. Various mechanisms proposed for these atresias include differential growth of the tracheal bud and the oesophageal tube, recanalisation of the duodenal lumen, and formation of the urogenital diaphragm and cloacal membrane at the hind gut, respectively. Antenatal diagnosis of OA and TOF and duodenal atresia is possible but these conditions were missed in our case although multiple other congenital anomalies were detected. Nor were these conditions suspected in the postnatal period as the nasogastric tube was not obstructed by the proximal oesophageal blind pouch and radiographs confirmed its position in the abdomen and not curled up in the upper pouch. Various plausible mechanisms explaining such an exceptional event have previously been considered by Patel et al.4

With duodenal atresia, the double bubble sign is very obvious with significant dilatation of the duodenum. However, if there is an associated OA+TOF, then the infant may not be able to swallow meconium directly into the stomach and the distal TOF may lower the pressure thereby preventing the stomach from significantly dilating, which may explain why the antenatal diagnosis was missed in this case.

Alimentary tract atresia can be an isolated anomaly, a chromosomal anomaly (most commonly a trisomy or the Martinez-Frias syndrome) or can present in the form of an association. VACTERL (vertebral, anorectal, tracheal, oesophageal and renal anomalies) association has been modified to VACTERL-D association with the addition of cardiac and limb anomalies, and we have amended it to VACTERL-D to incorporate duodenal obstruction due to atresia, stenosis, and malrotation or volvulus neonatorum to include the midgut in addition to the foregut and hindgut.6–8

Even with the advanced technology of the 21st century, there are limitations to antenatal diagnosis and postnatal treatment of multiple severe congenital anomalies. If all had been detected prenatally, the option of termination of pregnancy should have been given to the family if acceptable to them. The prognosis of numerous multisystem anomalies is poor with very reduced quality of life with both the family and society carrying the burden of the disease. Palliative care management avoids such a massive waste of resources and saves time, money, energy and ongoing distress for all parties including the patient.

In alimentary tract obstruction, the proximal obstruction should be corrected first. In our case, OA+TOF would have been treated first in order to manage the airway and breathing, followed by tracing the continuity of the lumen from the proximal to the distal direction and addressing the duodenal obstruction, and then colonic diversion for the anorectal malformation.6–7

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Competing interests None.

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REFERENCES
