The combination of pure oesophageal atresia with an associated missed H-type tracheo-oesophageal fistula

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DESCRIPTION
A 7-year-old girl presented with recurrent lower respiratory tract infections, coughing episodes associated with liquids and recurrent abdominal distension. There was maternal polyhydramnios and absent gastric bubble on antenatal anomaly scan with an antenatal diagnosis of pure oesophageal atresia (OA). There was no family history of OA for three generations on maternal or paternal side. At birth she had babygram showing arrested orogastric tube in the upper blind pouch and gasless abdomen, no vertebral anomalies and spinal ultrasound, echocardiogram and renal ultrasound were all normal. Voiding cystourethrogram with distal loopogram demonstrated bilateral vesicoureteral reflux (VUR). The chromosomal studies revealed normal 46XX karyotype and the patient had genetic counselling. She was born with partial VACTREL (Vertebral, Ano-rectal malformation, Cardiac anomalies, Tracheal anomalies, Esophageal anomalies and Limb anomalies) association comprising pure oesophageal atresia, anorectal malformation with rectovesibular fistula and bilateral VUR and treated elsewhere. She underwent gastrostomy and left descending colostomy at birth. The following week, she underwent posterior sagittal anorectoplasty with colostomy closure. At 2 months, she underwent repair of the long gap OA with gastric interposition, closure of gastrostomy and pyloroplasty as delayed primary closure proved impossible. Postoperatively she developed a leak at the gastro-oesophageal anastomosis which was treated conservatively. In the following 2 year she lived abroad and was only followed up within the UK on an intermittent basis. She underwent upper gastrointestinal contrast twice during this period which showed no evidence of a stricture but demonstrated oesophageal reflux and the recent one suspicious of an H-tracheo-oesophageal fistula (H-TOF, figure 1). She underwent combined bronchoscopy and an upper gastrointestinal endoscopy. This demonstrated a large H-type fistula, which could be easily cannulated,
that entered the oesophagus at 15 cm. She underwent ligation of H-TOF with interposition of a muscle flap via a right-sided neck incision after using the fibre optic light of the oesophagoscope to visualise the fistula intraoperatively. The postoperative period passed uneventful and she made a full recovery. She is asymptomatic, well and thriving at 6-year follow-up. Combined OA and H-TOF are very rare. The triad of symptoms, contrast study and endoscopy are diagnostic.1–3

**Learning points**

- Combination of pure oesophageal atresia (OA) and H-tracheo-oesophageal fistula (H-TOF) is very rare and should be suspected if there is a triad of symptoms of recurrent lower respiratory tract infections, coughing and/or choking episodes mainly with liquids and recurrent abdominal distension due to preferential high distal runoff of air from the tracheo-oesophageal fistula caused by low resistance in the oesophagus as compared with stiff lungs due to repeated aspiration pneumonia and associated fibrosis.
- Upper gastrointestinal contrast study needs special technique to visualise the fistula and bronchoscopy is diagnostic. This allows ureteric catheter cannulation which helps in easy identification at surgery.
- Use of fibre optic light source of an endoscope provides brilliant transillumination and easy identification of the fistula and recurrent laryngeal nerve during surgery avoiding injury to this vital nerve.

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**REFERENCES**