Management of complete tracheal rings in a neonate with tetralogy of Fallot

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DESCRIPTION

A 3-day-old full-term female infant was postnatally diagnosed with tetralogy of Fallot with severe subpulmonary and pulmonary valve obstruction, ventricular septal defect with right to left shunt and patent ductus arteriosus (PDA). She was stable on room air with an intravenous prostaglandin infusion while awaiting surgical repair. On day 3 of life, she was found unresponsive with significant desaturation to 30%. Following successful bag-mask ventilation and recovery of her saturation, multiple attempts to intubate the patient yielded a grade I view (figure 1). However, 3.0, 2.5 or 2.0 endotracheal tubes were unable to be passed due to an obstruction immediately distal to the vocal cords (figure 2). Mask ventilation was maintained without difficulty between each attempt. Transthoracic echocardiogram was performed at bedside in order to confirm adequate pulmonary blood flow through the PDA. The patient was transported to the operating room with mask ventilation for emergent direct laryngoscopy and bronchoscopy.

Otolaryngology was unable to insert a rigid 2.5 mm bronchoscope into the trachea though a 1.9 mm bronchoscope was able to be passed. Complete tracheal rings were seen immediately distal to the vocal cords down to carina (figure 3). The surgeon was also unable to intubate the patient with a 2.0 endotracheal tube. Tracheostomy was not an option due to complete tracheal rings throughout the entire length of the trachea; therefore, the airway was temporised with a laryngeal mask airway to maintain ventilation. With no other options for a more definitive airway, the cardiac surgeon cannulated the patient’s right internal jugular vein and carotid artery for arteriovenous extracorporeal membrane oxygenation (ECMO). To prevent fluid overload to the lungs, the arterial cannula was inserted with the tip in the descending aorta to minimise flow through the PDA, and the prostaglandin drip was discontinued. Two days

Figure 1 Normal vocal cords are visualised on direct laryngoscopy.

Figure 2 Obstructive tissue can be visualised beyond the vocal cords.

Figure 3 The severity of the complete tracheal rings is indicated by airway narrowing to less than 1.9 mm in diameter.
later, the patient was taken to the operating room for complete repair of her tetralogy of Fallot and slide tracheoplasty on cardiopulmonary bypass.

Oxygenation and ventilation were particularly concerning in this patient due to potential risk of tet spells triggered by hypoxia, hypercarbia and acidosis. In addition to complications of ECMO, including infection, bleeding and neurological injury, there were other factors to consider for this patient. First, in the setting of a PDA, ECMO would result in significant pulmonary overflow at the expense of systemic blood flow. By inserting the arterial cannula into the right common carotid artery and advancing it past the PDA to the descending aorta, the PDA could be bypassed and prostaglandin could be discontinued. Second, the patient would require cardiopulmonary bypass for the tetralogy of Fallot repair and slide tracheoplasty. Surgeons note improved surgical exposure and more precise surgical repair without an endotracheal tube obstructing the field.

Congenital tracheal anomalies usually present as respiratory distress, stridor or respiratory tract infections during the first days of life, but can be asymptomatic in some patients.1 Long-segment tracheal stenosis can be life-threatening and can require emergent airway management with mechanical ventilation. If this is not possible prior to surgical repair, ECMO or stenting may be considered as temporising measures to oxygenate and ventilate.2

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