Occurrence of Tension Pneumothorax following Fogarty Embolectomy Catheter use for Lung Isolation in a Neonate with Congenital Pulmonary Airway Malformation

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Abstract

A pneumothorax can arise in a variety of clinical settings in the operating room, from bleb rupture to iatrogenic injury. Should it expand in size and cause increase in intrapleural pressures with resultant diminished venous return, it becomes referred to as a tension pneumothorax. We report a case of a premature infant presenting for repair of congenital pulmonary airway malformation who suffered an iatrogenic tension pneumothorax during lung isolation with fogarty embolectomy catheter used for single lung ventilation in a neonate.

Introduction

Congenital pulmonary airway malformation (CPAM), formerly known as congenital cystic adenomatoid malformation (CCAM), is one of the most frequent pulmonary malformations that presents as a multicystic mass of pulmonary tissue with proliferation of bronchial structures that fail to mature [1]. While the exact pathogenesis is unclear, it has been hypothesized that the cysts develop either because of failure of the bronchial buds to join alveolar mesenchyme [2,3] or from overgrowth of terminal bronchioles. Children presenting with CPAM bring challenges with regard to their anesthetic, including oxygenation and need for lung isolation. We discuss a neonate presenting with a right-sided CPAM lesion, who developed a tension pneumothorax of the contralateral side, secondary to attempted lung isolation with a fogarty embolectomy catheter.

Case Description

A 5-day-old, 2.6-kg male born at 36 weeks’ gestation via c-section was noticed to have respiratory difficulty shortly after birth. Radiologic investigations revealed a right-sided chest mass with slight mediastinal shift (Figure 1). A diagnosis of CPAM was made, and the child was scheduled for right upper lobectomy. He was otherwise healthy with no abnormal laboratory or clinical findings. The case was discussed with the surgical team, and it was decided to proceed with lung isolation and single-lung ventilation (SLV). Atropine was administered intravenously, and anesthesia was induced with propofol and rocuronium. Mask ventilation was instituted using minimal pressures with 2% sevoflurane and 100% oxygen. Direct laryngoscopy (DL) was performed and a 4 Fr Fogarty embolectomy catheter (FC) easily inserted until resistance was met. An initial attempt with

Figure 1: CT of the chest with contrast showing congenital pulmonary airway malformation in the right lung.
a 3.0 cuffed tube was met with resistance, and, subsequently, a 3.0 uncuffed endotracheal tube (ETT) was placed alongside the FC. A fiberoptic scope (FOS) was then advanced through the ETT, and we discovered that the FC had entered the unintended left main bronchus. We therefore withdrew the catheter and repositioned it into the operative (right) bronchus. The patient’s oxygenation and ventilation were optimal at this point and blood pressure and heart rate within normal ranges. Several minutes thereafter, the end-tidal CO₂ tracing diminished and then disappeared. Oxygen saturations then began to fall rather dramatically, and the child developed bradycardia. At this time, ETT position was confirmed and auscultation revealed minimal coarse sounds on the right and absent sounds on the left chest. Acute bronchospasm was suspected, and albuterol and epinephrine were administered. With no resultant improvement, pneumothorax was suspected. A stat portable radiograph was ordered; however, given the rapidly deteriorating condition of the child, it was decided not to wait for the radiograph but place the chest tube en-situ. Figures 2). Immediately upon decompression, we were able to reestablish ventilation and ETCO₂ tracing reappeared and oxygenation improved. Hemodynamic parameters returned to physiologic range. In view of the life-threatening episode that had occurred and nonurgent nature of the procedure, a decision was made to postpone surgery. The child was transported back to the neonatal intensive care unit intubated and ventilated in stable condition.

Discussion

Congenital pulmonary airway malformation usually presents in one of three ways: Respiratory distress of the newborn, recurrent pneumonia in young children, or incidentally in asymptomatic adults [4]. Should a newborn present with CPAM, initiation of breathing after delivery may lead to gas trapping and further cystic expansion with worsening respiratory compromise. Most CPAM lesions require resection of the mass either thoracoscopically or by open thoracotomy, and prognosis after surgical excision is generally good. With increasing popularity of thoroscopic approach to lung surgery, SLV is requested for an increasing spectrum of surgical procedures in infants and children. Advances in anesthetic equipment, lung blocking devices, monitoring, technology, and pharmacology have increased the safety of one-lung anesthesia. Despite these advances, injuries are associated with SLV devices, such as double-lumen tubes (DLT) and endobronchial blockers. To date, there is only one other documented case of accidental bronchial injury with FC during lung isolation likely related to overdistension of the FC balloon, which was noticed by the surgical team during thoracotomy. In this instance, there were no hemodynamic sequelae, and the lesion was repaired intraoperatively [5]. Literature comparing injuries in adult patients who have been intubated with DLT show more frequent complications such as hoarseness and vocal cord lesions compared with those who had single-lumen tubes with blocking devices [6]. To our knowledge, this is the first case where injury to the contralateral healthy lung has occurred because of FC used for lung isolation.

Infants and neonates, because of their size, are unable to accommodate either a DLT or Arndt blocker. Lung isolation in this age group is accomplished either by endobronchial advancement of the ETT to the ventilated nonsurgical lung or by placing an FC alongside the ETT to block the surgical lung [7]. The latter technique offers the advantage of accurate lung block and obviates the need for ETT withdrawal into the trachea when expansion of the collapsed lung needs to be re-established. The FC has been used at our hospital for SLV in infants for many years and has worked quite well without any complications. It is advanced into the trachea by DL, and the ETT is then placed alongside the blocker. The blocker is then accurately positioned in the desired main stem bronchus with the help of FOS advanced through the ETT, and the blocker balloon is inflated with air under fiberoptic view [8].

The tension pneumothorax in this scenario is the likely result of mechanical injury from the FC that was used for lung isolation. Another possible cause however is barotrauma secondary to positive pressure ventilation that was commenced after the child was intubated. We feel that trauma due to the FC to be more likely because pneumothorax occurred soon after the blocker was withdrawn from the left lung and positioned in the right lung. The FC has a considerably stiff metal stylet, and it is quite likely that, as we were trying to advance the ETT alongside the blocker, we may have inadvertently pushed the blocker deeper into the lung that led to perforation of one of the smaller more distal bronchi. This was not clinically apparent as long as the blocker was sealing the hole, but, when the blocker was withdrawn from the left and redirected to the correct side, the puncture site was no longer occluded and pneumo-
needed to ensure its safe use until a better alternative is available to accomplish lung isolation in this age group [10].

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References


