Unilateral Lung Aplasia Presenting with Acute Respiratory Distress Possibly due to milk aspiration in Young Toddler

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Abstract

A 13 month female child with h/o recurrent respiratory tract infection developed cough and breathlessness nearly 30 min after breastfeeding. Chest x-ray showed homogeneously opaque right hemithorax with hyperlucency of left lung field suggestive of lung agenesis without any features of foreign body. Bronchoscopy revealed oedematous mucosa, thick bronchial secretions, mucus plug and white coagulates (apparently from milk) which were removed from left bronchus; trachea continued as left main bronchus without any stenosis or compression with absent right bronchus. Pulmonary agenesis was confirmed by CT scan in the post-operative period. The child made full recovery.

Key words: Lung aplasia, Respiratory distress, Recurrent respiratory tract infection.

Introduction

Lung aplasia is often associated with acute respiratory distress and has a high mortality rate as this is usually associated with anomalies like coarctation of aorta, aortic stenosis, transposition of great vessels and septa defects [1]. Fifty percent born with pulmonary aplasia are stillborn or die within the first five years of life [2]. Bilateral congenital pulmonary agenesis is a rare lethal anomaly, first described by Morgagni [3]. In unilateral lung agenesis, the trachea continues directly into the main bronchus of the normally developed lung. Foreign body (FB) inhalation is a potentially life threatening event, in these patients. A small reduction in airway radius due to oedema will result in a large increase in resistance to air flow. Hypoxia may rapidly occur because of high oxygen consumption of infants. [4] Bronchoscopy is done for a wide variety of diagnostic and therapeutic procedures. Bronchoscopic procedures poses a number of challenges in paediatric age group and requires the paediatrician to be fully familiar with airway and medical management,[5] We describe a case of right lung agenesis presenting with respiratory distress due to aspiration of breast milk which was misdiagnosed as a case of foreign body bronchus.

Case report

A 13-month-old female child, weighing 10 kg, presented with severe respiratory distress and peripheral cyanosis. The child had breast-feeding about 2 hours before onset of symptoms. She started to cough and became breathless 30 min after feeding. She had past history of recurrent respiratory tract infections. On examination, the child had cough, dyspnea, and tachypnea with peripheral cyanosis. Her heart rate was 140/min, respiratory rate was 45/min and oxygen saturation was 85% in room air. Air entry was absent on right side of chest while heezing was noted on left side of chest. Chest wall was bilaterally symmetrical without any skeletal deformity. Plain X-ray chest showed homogenously opaque right hemithorax with mediastinal shift and hyperlucency of left lung field. Any definite diagnosis could not be made, and emergency bronchoscopy was planned under general anaesthesia.
Bronchoscopy revealed oedematous mucosa, thick bronchial secretions, mucus plug and white coagulates (apparently from milk) which were removed from left bronchus. Bronchoscopy also revealed that trachea continued as left main bronchus without any stenosis or compression with absent right bronchus. A 4 mm plain oral endotracheal tube was placed after bronchoscope removal, and the lung was well expanded. Air entry improved on the left side. The child improved dramatically in the postoperative period.

Medications included hydrocortisone 50 mg and dexamethasone 6 mg. Heart rate, ECG, oxygen saturation, temperature and BP was monitored. Intraoperative heart rate and oxygen saturation were in the range of 140-170 per minute and 97-99%, respectively.

In subsequent evaluations, contrast enhanced CT-scan thorax showed absence of right bronchus and pulmonary parenchyma with normal hyper-inflated left lung extending anteriorly across midline to right, and slightly posterior deviation of trachea. In the absence of any other pathology and FB, the diagnosis of milk aspiration leading to acute respiratory distress in a patient with unilateral lung agenesis was made. After 5 days of treatment child was discharged from hospital.

Discussion

Pulmonary agenesis is defined as complete absence of bronchus, parenchyma and vessels. The present case has isolated unilateral lung agenesis of right side without any other associated anomalies.

In the present case, although there was h/o breastfeeding a while back, it was not sufficient lead for any diagnosis alone or along with the chest X-ray. Radiological features of lung agenesis mimic that of foreign body aspiration and should be considered in unilateral hyperinflation and mediastinal shift.

The challenges of anaesthetic management in such cases include efficient sharing of the small airway for bronchoscopy and anaesthesia, prevention of systemic arterial desaturation, achievement of adequate depth of anaesthesia, minimization of airway secretions, stabilization of hemodynamics, rapid recovery of airway reflexes as well as minimization of sedation beyond the procedure[6]. The underlying pulmonary pathology may pose additional challenges such as compression of airways due to shifting vascular structures, limitation of pulmonary reserve, increased right ventricular afterload, pulmonary hypertension due to absence of right pulmonary artery and predisposition to pulmonary oedema due to reduced residual volume of lung and pulmonary vascular bed [7].

In the present case, flexible bronchoscope was used. Nandalike, et al presented a case of FB aspiration in a child with unilateral lung aplasia where FB was extracted by basket forceps via a flexible bronchoscope after failure to do so with rigid bronchoscopy[8]. Especially in select patients with abnormal airway anatomy, flexible bronchoscopy may be advantageous.

Conclusion

In summary, congenital pulmonary agenesis is an extremely rare anomaly and may present acutely with severe respiratory distress requiring lifesaving urgent intervention. High index of suspicion and meticulous management in consideration of underlying pathology and
associated anomalies are warranted for favourable outcome.

References


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