Isolated absence of right pulmonary artery in a 4-year old child: a case report

Abstract: Isolated unilateral absence of the pulmonary artery is a rare anomaly with congenital absence of the left or right pulmonary artery in the absence of other cardiac malformation, and diagnosis is usually made during adolescence. We report a 4-year old male child with isolated absence of the right pulmonary artery, who presented with recurrent chest infection and wheezing since infancy.

Keywords: pulmonary artery, child, wheeze, chest infection, Ethiopia

Case presentation

A 4-year old male child was referred to our hospital (St Paul’s Hospital Millennium Medical College located in Addis Ababa, Ethiopia) from a district hospital for further evaluation of recurrent chest infection and wheezing in 2018. He was admitted to the referring hospital eight times for chest infection in the first 2 years of his age. The first admission happened at the age of 2 weeks. Cough, fast breathing, and wheezing were the main presenting symptoms during the exacerbations. At each admission, he was treated with parenteral antibiotics and bronchodilators and was discharged from the hospital after he got improvement. He was apparently healthy in between the episodes of the respiratory illness. He did not have history of choking episodes. He was delivered in a hospital at term, and pregnancy was uneventful. He had optimal development and was a kindergarten student at the time of writing this report.

On examination, he was not in respiratory distress and SPO2 was 94%–96% with room air. He weighed 14 kg, with a height of 106 cm (his anthropometric indices lie above –2 SD except his weight-for-height which lies between –2 and –3 SD). Chest examination revealed trachea shifting to the right side, decreased air entry over the right lung field, and relative dullness over the same area. Cardiac auscultation revealed normal heart sounds with regular rate and rhythm with no gallop. He was alert and playful, having good interaction during the examination.

Basic laboratory examinations were within normal limits, and HIV status was negative. Chest X-ray showed hyperinflated left lung and reduced right lung volume with decreased vascularity and trachea shifted to the right (Figure 1).

Echocardiography demonstrated absence of the right pulmonary artery with no other cardiac lesion (Figure 2). Subsequent computed tomography (CT) scan also showed hypoplastic right lung with absent right pulmonary artery (Figure 3).

The case was evaluated by a team comprising pediatrician, pediatric pulmonologist, and cardiologist, and the team decided to follow the child regularly.

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Unilateral absence of pulmonary artery (UAPA) is congenital absence of the left or right pulmonary artery caused by involution of the proximal sixth aortic arch with persistence of intrapulmonary part to the distal sixth aortic arch. As a result, “proximal interruption of pulmonary artery” has been suggested as a preferred term.1,2

UAPA was first described in 1886 and is a rare anomaly with prevalence in the general population ranging from 1 in 200,000 to 1 in 300,000. It rarely presents as an isolated lesion and is more commonly associated with other heart defects. Associated defects include patent ductus arteriosus, tetralogy of Fallot, atrial septal defect, aortic coarctation, truncus arteriosus, and pulmonary atresia. Collaterals to the affected side commonly arise from bronchial arteries, but may also arise from intercostal, subclavian, and rarely coronary arteries.1–3

Common presenting symptoms include recurrent respiratory infections, dyspnea, exercise intolerance, hemoptysis, chest pain, and rarely wheezing. Diagnosis can also be incidental in an asymptomatic child. Infants may present with pulmonary hypertension (PHT) and congestive heart failure, whereas hemoptysis is the commonest manifestation in adults.1,2,4–7 Though there is a recent report of neonatal diagnosis of UAPA,2 diagnosis is usually made during adolescence (median age of 14 years), and risk of PHT, hemoptysis, and death significantly increases with age.2,5

The mechanism(s) for recurrent pneumonia include hypoperfusion with poor blood flow and decreased inflammatory

Figure 1 Posteroanterior chest radiograph showing hyperinflated left lung with reduced right lung volume.

Note: There is also decreased vascularity and hila not well visualized on the right hemithorax (RH).

Abbreviations: L, left; T, trachea; RH, right hilum.
isolated UAPA 4-year-old child
cell migration, impaired ciliary function, ventilation–perfusion mismatch, alveolar hypocapnea with poor mucosal defense, and protease–antiprotease imbalance.\textsuperscript{1,2}

Diagnosis of UAPA is based on a complete history, physical examination, and imaging. Absent pulmonary artery usually causes secondary lung hypoplasia, as in our case, and examination reveals asymmetrical chest with abnormal breath sounds on the affected side. Chest X-ray demonstrates reduction in the volume of hemithorax, elevation in the diaphragm, reduced intercostal spaces, and mediastinal shift to the same side. UAPA should also be suspected in an asymptomatic child with suggestive chest X-ray.\textsuperscript{1,9}

Echocardiography, CT of the chest, and magnetic resonant imaging can give the diagnosis with certainty, and invasive procedures including angiography are rarely performed except for intervention or to screen for complications; ventilation–perfusion scan is not necessary for diagnosis.\textsuperscript{1,10}

Thrombotic embolus of pulmonary artery, anomalous venous drainage of the right lung into the inferior vena cava (Scimitar syndrome), and Swyer–James syndrome should be considered in the differential and should be ruled out.\textsuperscript{9}

Therapeutic strategies for UAPA remain unclear. Presence or absence of complication, pulmonary artery anatomy, associated aortopulmonary collaterals, cardiovascular anomalies, and PHT guide the decision to treat. Regular follow-up is all that the child needs if she/he has normal cardiopulmonary function. Most recommend treatment for the small number of patients with hemoptysis, recurrent respiratory tract infection, or PHT. Treatment options include revascularization, pulmonary vasodilator therapy for PHT, partial or total pneumonectomy, primary or staged pulmonary artery anastomosis, and embolization of collateral hemorrhage.\textsuperscript{1,11–15}

The prognosis depends on associated cardiovascular anomalies and the degree of PHT as well as presence or absence of hemoptysis.\textsuperscript{1,4,11}

Conclusion
We presented a child with recurrent respiratory symptoms who was finally diagnosed to have isolated unilateral absence of the pulmonary artery (IUAPA). Though IUAPA is a rare condition, it should be considered as a differential diagnosis on a child presented with recurrent chest infections and wheezing. Diagnosis can be made based on history, physical examination, chest X-ray, echocardiography, chest CT, and magnetic resonance imaging without employing invasive diagnostic studies. Treatment is unclear and is usually individualized based on symptoms and associated conditions and is decided by expectant medical or surgical approach. The case described on this case report is being followed by a team consisting of a pediatrician, pulmonologist, and cardiologist.
on an outpatient basis. As stated above, therapeutic strategies for IUAPA remain unclear and initiation of treatment should be guided by the presence or absence of cardiac or respiratory complications. Therefore, the team decided to follow the child regularly since currently he does not have the anticipated complications of IUAPA, except the recurrent chest infection which he has had in the past and which was treated.

**Ethics statement**

Written informed consent was obtained from the parent of the child to publish this case report. According to the standard of operating procedure of the Institutional Review Board of our college (St Paul’s Hospital Millennium Medical College), publishing a case report is exempted from the board review.

**Author contributions**

All authors contributed toward data gathering and compilation, drafting and critically revising the paper, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**
