Unilateral Agenesis of Pulmonary Artery

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Abstract

Isolated unilateral absence of pulmonary artery is uncommon. Although not certain, genetic, teratogenic and mechanical factors are held responsible. The anomaly results from the involution of the proximal sixth aortic arch. In this report we present an asymptomatic 24 yrs old male patient. Unilateral right-sided absence of pulmonary artery was diagnosed by thoracic computerized tomography (CT) and angiographic evaluation.

Keywords: Congenital pulmonary disease; Pulmonary artery agenesis; Pulmonary artery disease

Introduction

Isolated unilateral agenesis of pulmonary artery was first defined at 1868. Although the exact prevalence is unknown, in a study it is estimated as 1/200,000. Congenital unilateral agenesis of pulmonary artery is a rare anomaly which is frequently associated with other congenital cardiovascular anomalies such as tetralogy of Fallot or septal defects. In 37% of cases left pulmonary artery is absent while in 63% the right one is absent. Nearly 30% of patients are asymptomatic. When symptomatic, patients may experience chest pain, pleural effusion, recurrent pulmonary infections, dyspnea or limited exertion or hemoptysis. Several methods are used in diagnosis such as computerized tomography, bronchography, perfusion scintigraphy, echocardiography and cardiac catheterisation. Massive hemoptysis, cardiac insufficiency, bronchectasis and necrotizing bronchopnemonia may be included among the complications. The following case report presents a patient who was not diagnosed until adult ages.

Case Report

A 24-year-old man was admitted to our hospital with dyspnea. He smoked cigarettes 8 pcks/year and currently he was a light smoker. His medical and family history was unremarkable. The physical examination and laboratory findings (biochemical markers, C-reactive protein (CRP), erythrocyt sedimentation rate (ESR)) were normal as well. Respiratory function test revealed slight restriction. Chest x-ray confirmed the right sided shrunken lung and hyperinflation of the left one. 3-D rotational angiography was performed following the thorax CT which revealed atresia of right pulmonary artery. [(MIP) minimum intensity projection images seen Figure 1 and 2, volume rendering (VR) images seen Figure 3 and 4 and Thorax CT angiography image seen Figure 5 of the patient].

Figure 1. MIP coronal mediasten window. Left arrow: Left pulmonary arter and its branches; in right absence pulmonary arter.
Discussion

Bilateral agenesis of pulmonary artery is incompatible with life [1-6]. Isolated agenesis of pulmonary artery is a rare congenital anomaly [6, 7], 108 cases were reported during the timeframe between 1974 and 2000 [8]. Cardiovascular, gastrointestinal, neuromusculoskeletal and genitourinary system anomalies may coexist in 50% of cases [6]. Symptoms become evident at very early ages in cases in which other congenital anomalies associate. There wasn’t any additional system anomaly in our case. Since this anomaly progresses asymptomatically in adults, the cases are diagnosed by chest radiographs taken incidentally. Contrary to Komatsu, Walter and Gulhan’s cases, our case did not present with hemoptysis [5, 7, 9]. Despite the other cases reported in literature, bronchectasis was not determined in our case although there was a volume loss at the right lung. The absence of coexisting bronchectasis and cardiac or another congenital anomaly is probably the reason for the negative findings in previous health screenings. The prognosis of isolated right pulmonary agenesis is better than the left one and it is less associated with other congenital anomalies [6], 20% of cases of isolated right pulmonary agenesis is associated with pulmonary hypertension. Asymptomatic patients with pulmonary hypertension may become symptomatic at pregnancy or high altitudes [10, 11]. Echocardiographic evaluation of our patient revealed pulmonary systolic hypertension. Since the effected side provides the blood from systemic circulation, the progression of left to right shunt may lead to heart failure in future. CT angiography of our case revealed that the right lung supplied blood from systemic circulation. Nevertheless he hadn’t symptoms of heart failure for the time being.

When asymptomatic, unilateral absence of pulmonary artery presents at incidental chest x-rays with small hilar shadow and hyperlucent lung tissue at the effected side. In our case, posteroanterior chest radiography taken for general examination revealed the absence of a pulmonary arterial vena hilar shadow which was suggestive of a vascular

Figure 2. MIP coronal image, lung parenchyma window; right upper and lower arrows: shows the systemic collateral vessels.

Figure 3. VR axial image; the right pulmonary artery is atresic. Right arrow shows systemic collateral vessels.

Figure 4. VR image; Volume loss at the right lung with pulmonary atresia compared to the left one; deviation of the mediastinal and bronchial structures to the right and increased interstitial density at the parenchyme.

Figure 5. Thorax CT angiography, axial plan, mediastinal window; arrows; systemic collateral vessels surrounding the trachea.
pathology in differential diagnosis. Definitive diagnosis in this type of cases is made by pulmonary arteriography. Eventually, we diagnosed our case by CT angiography. Hyperlucency was determined in most cases presented in literature [6, 7, 12-14]. Unlike the cases with Swyer-James-MacLeod syndrome, in our case, the right main pulmonary artery was not detected at pulmonary CT angiography [12, 13]. The upper right lung was fed by the branches of thyrocervical trunkus while the medium and lower lung’s blood supply came from the branches of intercostal arteries. Asymptomatic cases do not require treatment while surgical treatment may be planned for coexisting cardiac anomalies in cases with absence of left pulmonary artery. Pneumonectomy of the affected side may be indicated in case hemoptysis becomes life-threatening or there are cystic bronchectasic formations.

Revascularisation is performed in selected cases [6]. Surgical treatment was not an option in this present case since bronchectasis and cardiac or any other genetic vascular malformation was not determined at the hypoplastic pulmonary tissue [14-17]. Isolated absence of pulmonary artery may not be diagnosed until they reach adulthood because of the asymptomatic benign course or misdiagnosis [8, 18, 19]. The presence of any hilar pathological image at chest roentgenograms taken for any other purpose should suggest vascular anomalies. In addition, unilateral atresia of pulmonary artery should be regularly followed up closely for progression of possible complications.

**Conflict of Interest**

There was no conflict of interest among authors.

**References**