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Case Report

Unilateral absence of pulmonary artery and azygous continuation of interrupted inferior vena cava

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ABSTRACT

Unilateral absence of pulmonary artery is a rare developmental anomaly. Infrahepatic inferior vena cava interruption is a well-recognized but uncommon developmental anomaly. Presence of both these anomalies in a single individual is extremely rare. A 58-year-old man with a history of recurrent lower extremity deep vein thrombosis and venous insufficiency presented to our emergency department with bilateral calf pain and swelling. Ultrasound demonstrated extensive deep vein thrombosis throughout bilateral lower extremities. Computed tomography angiography showed smooth tapering of the right pulmonary artery with absent distal most segment. To our knowledge, there is only 1 case report in the literature so far with both the abnormalities present in a single individual.

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Introduction

Unilateral absence of pulmonary artery (UAPA) is a rare congenital abnormality caused by the failure of the sixth aortic arch to connect with the pulmonary trunk during the embryologic development [1–3]. Intrapulmonary vessels and the distal portion of the affected pulmonary artery trunk can develop normally, and blood supply is achieved by systemic collaterals from bronchial, major aortopulmonary collaterals and other systemic arteries [2,4]. In most patients, UAPA is associated with other cardiovascular congenital abnormalities such as tetralogy of Fallot, arterial septal defect, and truncus arteriosus [2]. It is usually diagnosed in infancy or childhood due to symptoms related to other associated cardiovascular abnormalities. Few patients who remain asymptomatic until adulthood can present later with recurrent respiratory infections, dyspnea on exertion, high-altitude pulmonary edema, pulmonary hypertension in the contralateral lung, or hemoptysis [1].

 Interruption of the inferior vena cava (IVC) results from agenesis of a segment of the IVC or a fusion defect between the prerenal and hepatic segment of the IVC. Its prevalence is 0.6-2.0% in patients with congenital heart disease [4,5]. It results in termination of the IVC below the hepatic vein. The infrahepatic IVC may continue as the azygos vein or may continue as the hemiazygos vein to the left SVC, intrathoracic veins, or anomalous infrahepatic veins [2,3]. The hepatic seg-
Fig. 1 – (A) Smooth tapering of the right main pulmonary artery with absent distal segment. (B) 3D volume rendered image (posterior view) showing smooth tapering of the right main pulmonary artery with absent distal segment.

Fig. 2 – (A) Enlarged right internal mammary artery as a major collateral supply for right lung. (B) Thickened right pleura as evidence of additional systemic to pulmonary shunt.

Case report

A 58-year-old man with a history of recurrent lower extremity deep vein thrombosis (DVT) and venous insufficiency presented to the emergency department with bilateral calf pain and swelling. Ultrasound demonstrated extensive DVTs throughout bilateral lower extremities. Computed tomography angiography showed no pulmonary emboli, but demonstrated smooth tapering of the right main pulmonary artery and paucity of right intrapulmonary arteries. (Fig. 1A, and B). The left pulmonary artery was dilated while the pulmonary trunk measured within normal limits. The left lung showed compensatory hyperinflation. Dilated right internal mammary and right bronchial arteries were present as main aortopulmonary collaterals (Fig. 2A). Thickened right pleura showed evidence of additional systemic to pulmonary shunt (Fig. 2B). Incidental note was made of dilated azygos vein draining an interrupted IVC to the superior vena cava (Fig. 3A and b). Aortic arch was left sided and no other cardiovascular abnormality was seen.

Discussion

UAPA and azygos continuation of the IVC are rare congenital anomalies [1,2,4]. To our knowledge, this is the second case reported where both the anomalies are present in a single individual. Our patient did not have any other congenital cardiovascular anomalies like tetralogy of Fallot, arterial septal defect, and truncus arteriosus which are usually associated with unilateral pulmonary artery agenesis [2,3]. Patient did not endorse any common symptoms of UAPA such
as respiratory infections, dyspnea on exertion, high-altitude pulmonary edema, or hemoptysis. Rather patient presented with bilateral calf pain and swelling with a history of recurrent lower extremity DVTs and venous insufficiency. CT findings of both UAPA and interrupted IVC with azygous continuation were incidental. Interrupted IVC with azygous continuation is known to cause recurrent, unprovoked DVTs and venous insufficiency, possibly because the dilated vena azygos system cannot drain the lower limbs adequately [4]. This anomaly may be present in 5% of cases of idiopathic DVT in patients under 30 years [4,5]. Our patient had no other risk factors for DVT or venous insufficiency and interrupted IVC may be the potential cause. No known association of UAPA and azygous continuation of an interrupted IVC was found in literature. Awareness of the systemic collaterals in UAPA patients is important for ensuring the accurate selective embolization of bleeding vessels. Presence of interrupted IVC poses more problem in patients with UAPA during invasive procedures, such as right heart catheterization procedures, electrophysiological procedures, IVC filter placement, temporary pacing through the transfemoral route, and abdominal or thoracic surgery if need arises.

Knowledge of interrupted IVC with azygous continuation in a patient with UAPA is important before interventional or surgical procedures. In conclusion, we describe a unique case of 2 rare congenital anomalies occurring in a single individual, their radiological findings and potential management challenges.

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