Case Report

Anomalous origin of the left pulmonary artery from the ascending aorta in two children with pulmonary atresia, sub-aortic ventricular septal defect and right-sided major aorto-pulmonary collateral arteries

L PEPETA, FF TAKAWIRA, PE ADAMS, NH NTSINJANA, BJ MITCHELL, AM CILLIERS

Abstract

We report two rare cases of an anomalous origin of the left pulmonary artery (AOLPA) from the ascending aorta, associated with pulmonary atresia, a ventricular septal defect and a left aortic arch. The cases are unusual because AOLPA is more commonly associated with a right aortic arch and it is more usual for the right pulmonary artery to originate anomalously from the ascending aorta. The pulmonary blood supply to the right lung in both patients was absent and provided instead by major aorto-pulmonary collateral arteries which were stenosed at multiple levels. The AOLPA in both patients originated from the postero-lateral aspect of the ascending aorta just distal to the sino-tubular junction. Only one patient showed the more common association of an unusual aortic arch branching pattern in the form of an anomalous right subclavian artery.

Neither patient was in heart failure and the chest X-ray in both revealed differential pulmonary perfusion with prominent vascularity of the left lung. Cardiac catheterisation showed systemic pressures within the anomalous left pulmonary artery. Karyotyping revealed normal chromosomes, and fluorescent in-situ hybridisation done in one patient was negative for chromosome 22q11.2 microdeletion. Both patients have been managed conservatively.

Keywords: pulmonary atresia with ventricular septal defect, anomalous origin of pulmonary artery, collateral arteries, pulmonary hypertension, branchial arches, CATCH22 syndrome

Submitted 23/3/10, accepted 31/8/10

Cardiovasc J Afr 2010; 21: October online publication www.cvja.co.za
DOI: CVJ-21.048

An anomalous origin of the pulmonary artery from the aorta (AOPA) is very rare, with the first case of this nature reported by Fraenzel in 1868.1 The incidence of anomalous left pulmonary artery from the aorta (AOLPA) was reported in one study to be around 17%,1 with anomalous origin of the right pulmonary artery (AORPA) from the ascending aorta as the more common variant of AOPA, seen in about 83% of cases. Without surgery, the mortality in patients with AOPA is extremely high, reaching as high as 80%.2-5 This is due to severe, rapidly progressive pulmonary vascular disease and congestive cardiac failure. Both isolated AOLPA and that associated with other defects have been well described.6-8

The commonest associated congenital cardiac lesion is tetralogy of Fallot, which is seen in about 75% of cases.1 AOLPA is also associated with a right arch of the aorta in 63% of cases, and in about 38% of cases, an anomalous origin of the right subclavian artery from the descending aorta has been reported as well.1 Of note, both our cases had a left aortic arch and had no main and right pulmonary arteries, with the only source of blood supply to the right lung being partially stenosed major aorto-pulmonary collateral arteries (MAPCAs).

To the best of our knowledge, this series represents the first report in the literature of the anomalous origin of the left pulmonary artery arising from the aorta, associated with pulmonary atresia, with no central pulmonary arteries and MAPCAs supplying the whole of the right lung.

Case 1

This 10-month-old male patient was referred for assessment of a complex cyanotic heart lesion and a centrally positioned heart, namely mesocardia. Clinical assessment showed the patient to have central cyanosis, a loud single second heart sound, and 1/6 ejection systolic murmur over the left upper parasternal border. There were no abnormal facial features, no clubbing of the digits
or features of heart failure.

The chest X-ray showed a cardiothoracic index of 66%, mesocardia, a left-sided aortic arch, a boot-shaped heart with right ventricular predominance and an absent pulmonary artery segment. Differential perfusion to the lungs was evident on the chest X-ray, showing prominent pulmonary vasculature on the left and reduced pulmonary vasculature on the right. Situs solitus was present (Fig. 1).

The electrocardiogram revealed right-axis deviation and right ventricular hypertrophy by voltage criteria. An echocardiographic assessment confirmed the presence of pulmonary atresia with a large mal-aligned peri-membranous (sub-aortic) ventricular septal defect, and an overriding aorta. A surprising finding was the anomalous origin of the left pulmonary artery from the left postero-lateral aspect of the ascending aorta, just above the sinotubular junction, and an aberrant right subclavian artery arising from the descending aorta distal to the left subclavian artery. In addition, numerous major aorto-pulmonary collateral vessels were seen arising from the descending aorta, and the ascending aorta arched to the left (Fig. 2). The central pulmonary arteries were not visualised.

Angiography confirmed the above findings (Figs 3, 4). Of importance were the total absence of native pulmonary arterial vasculature and the presence of a bipartite right ventricle with an absent apical portion. Haemodynamic studies revealed very...
low systemic saturations of 79% and, as was expected, very high systolic pressures equalling that of the aortic pressures of 61 mmHg in the aberrant left pulmonary artery. By contrast, the pressures measured within the major aorta-pulmonary collateral vessels which supplied all three lung segments on the right side were slightly elevated with a systolic pressure of 35 mmHg, and a mean of 19 mmHg. The patient was not offered any further treatment.

Case 2

The second case was a 6½-year-old female patient who presented with anomalous origin of the left pulmonary artery from the ascending aorta, pulmonary atresia with a ventricular septal defect (VSD), again associated with MAPCAs as the only supply to the right lung. She was diagnosed with this problem as a newborn baby. Unfortunately, she was lost to follow up and presented at 6½ years with worsening cyanosis and decreased effort tolerance. Clinically, she had no dysmorphism. She was cyanosed with pulse oximetry saturations of 85% at room air. There was associated digital clubbing. The precordium was bulging with a laterally displaced apex. The second heart sound was loud and single, and there was a 3/6 ejection systolic murmur over the left upper parasternal border. There was no evidence of congestive cardiac failure.

On chest X-ray, the cardiothoracic ratio was slightly increased at 62%, there was a left arch of the aorta, and the pulmonary artery segment was absent. Of note in this patient was differential perfusion with plethoric left lung fields, compared to the right.

The electrocardiogram revealed features of left atrial and left ventricular hypertrophy. On echocardiographic assessment, there was a large sub-aortic mal-aligned VSD with 50% aortic override and bi-directional shunting across the VSD. The ventricles were balanced in size and the left atrium was dilated, with a left atrium:aortic index of 2.6. There was associated pulmonary atresia. The left pulmonary artery was noted to be arising from the ascending aorta just above the sino-tubular junction and both the main and the right pulmonary arteries were not seen.

At cardiac catheterisation, the systemic arterial saturations were low at 67%, in keeping with a mixing situation. There were balanced ventricular pressures with systolics of 82 mmHg. As expected, the AOLPA pressures were systemic at 82 mmHg.

Angiography confirmed the echocardiographic findings of pulmonary atresia with a VSD, associated with an anomalous origin of the left pulmonary artery from the ascending aorta. There were no native pulmonary arteries supplying the right lung that were seen, and the only source of blood supply to the right lung was the partially stenosed MAPCAs arising from the descending thoracic aorta. This patient was conservatively managed as well.

Discussion

The presentation of a single pulmonary artery arising directly from the ascending aorta is very rare, and even less common is the entity of AOLPA. Besides the well-known association with tetralogy of Fallot, AOLPA is usually associated with a right-sided aortic arch and an absent patent ductus arteriosus (PDA). By contrast, our case study patients presented with a left arching aorta. A frequent association of AORPA is the CATCH 22 syndrome, which was not found in one of our cases, based on the absence of the 22q11 deletion anomaly following a fluorescent immuno-sorbent hybridisation (FISH) chromosomal analysis. The other patient was not tested for this chromosomal aberration.

Only one case has been reported in the literature, where AOLPA and tetralogy of Fallot with membranous pulmonary atresia were diagnosed. In this patient, there were well-developed central pulmonary arteries and the patient underwent successful complete surgical repair. The aetiology of AOLPA is unclear, but one case of AORPA has been reported to be associated with fetal valproate syndrome.

The pathogenesis of an anomalous pulmonary artery originating from the aorta is said to be related to an abnormality in the development of the aortic or branchial arches. The failure of resorption of the fifth branchial arch and presence of the sixth brachial arch usually lead to the anomalous origin of the right pulmonary artery from the aorta. By contrast, the anomalous origin of the left pulmonary artery from the aorta is reported to be due to developmental absence of the fifth and sixth branchial arches. Abnormalities of the branchial or aortic arches are associated with neural crest-cell aberrations and therefore may be associated with the CATCH 22 syndrome. Precursors of the trunco-aortic sac are mainly of mesodermal origin, therefore AOPA is unlikely to be associated with the trunco-aortic sac abnormalities.

The origin of AOLPA anatomically is usually very proximal, arising from the ascending aorta just distal to the aortic sinus. Other origins of the pulmonary arteries, such as at the base of the innominate artery, from a patent ductus arteriosus, or from the thoracic aorta distal to the subclavian artery have been described and can be labelled as ‘distal types of AOPA’. They are not the same entity that is described in this case study.

The clinical presentation is that of a significant left-to-right shunt with early and rapidly progressive pulmonary hypertension, with or without right-sided heart failure. The pulmonary vascular disease affects both lungs, and may be seen as early as three months of age. The mechanism by which both lungs are affected involves neurogenic crossover from the involved lung to affect the protected lung, thereby producing reflex vasoconstriction in the protected lung. Diastolic run-off due to shunting of blood away from the aorta into the aberrant pulmonary artery may affect coronary perfusion and result in left ventricular dysfunction.

The diagnosis of AOPA can be easily achieved by careful echocardiography. Other imaging modalities such as cineangiography or magnetic resonance angiography have also been utilised. Due to the rapid development of irreversible pulmonary vascular disease in infancy and associated high mortality in the patients presenting with AOLPA, re-implantation of the pulmonary artery is an operation that needs to be performed within the first year of life.

Conclusion

Anomalous origin of the pulmonary arteries from the ascending aorta, and more specifically the left pulmonary artery is a rare entity, which can be diagnosed with readily available imaging modalities. The management is surgical re-implantation of the
anomalous vessel, but may provide a management challenge if complicated by pulmonary hypertension or other associations such as pulmonary atresia with collaterals vessels.

References