Pulmonary artery anomalies: Interesting cases with a common embryonic basis

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Abstract
Isolated pulmonary artery anomalies are rare. We present two babies, one with absent left pulmonary artery (LPA) and the other with pulmonary artery sling (PA Sling), with similar clinical presentations. Both of these defects have a common embryonic origin.

We discuss the importance of a high index of clinical suspicion and clear delineation the left pulmonary artery during echocardiography to facilitate early diagnosis.

Introduction
The aortic arches or pharyngeal arch arteries (once referred to as branchial arches in human embryos) are a series of six paired embryological vascular structures which give rise to several major arteries (Figure 1). The branch pulmonary arteries arise from the proximal part of the sixth right arch and the proximal part of the sixth left arch. Defective development of the proximal portion of the sixth left arch gives rise to a disconnected left pulmonary artery or a pulmonary artery sling depending on where the distal left pulmonary artery gets its blood supply from [1-3].

Pulmonary artery anomalies can co-exist with other congenital heart defects, or can be an isolated anomaly.

Methods
Two babies who presented with unexplained desaturation were investigated. The results of investigation and management are reviewed. A Medline search using the keywords was performed and results summarized.

Results
Case 1 – Absent/disconnected left pulmonary artery
A term baby with 22q deletion presented on day 5 with shortness of breath, sweating during feeds and seizures (secondary to hypocalcaemia). Cardiovascular examination was normal except for a prominent systolic murmur. His oxygen saturation was 98% in all four limbs. Initial echocardiogram demonstrated a patent ductus arteriosus (PDA) and a foramen ovale (PFO).

In view of unexplained intermittent desaturations and a heart murmur, a repeat echocardiogram was organised, which documented in addition a right aortic arch, and absence of the left pulmonary artery. Cardiac MRI identified the disconnected LPA. At surgical exploration, a connection of the LPA to the left common carotid artery was identified through an atretic segment. A conduit was successfully constructed between the pulmonary trunk and the LPA, with relief to symptoms (Figure 2).

Case 2 – Pulmonary artery sling
A preterm 33 weeker exhibited shortness of breath, noisy breathing and blue episodes while feeding in the first few weeks of life. These

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symptoms persisted despite respiratory support with CPAP, treatment of sepsis and anti-reflux medication. An initial echocardiogram demonstrated a PDA and PFO. As symptoms persisted, a repeat echocardiogram demonstrated the distal origin of the LPA from the right pulmonary artery, that coursed round the origin of the left bronchus, with airway compromise.

Following surgical transection of the LPA at its origin and implantation into the main pulmonary artery, the baby achieved a good outcome with minimal respiratory symptoms (secondary to the bronchomalacia) (Figure 3 and 4).

Discussion

Both babies had pulmonary artery anomalies. Interesting, both these anomalies have a common embryonic origin related to the proximal segment of the primitive 6th branchial arch arteries. Medline search emphasized the importance of early diagnosis is such situations, and highlighted the role of careful look at the branch pulmonary arteries during any echocardiogram. Early diagnosis and surgical recruitment facilitates growth of the pulmonary artery and optimizes associated lung development.

Conclusion

Anomalies of the pulmonary arteries have a common embryonic origin, and could be overlooked at initial echocardiograms. The pulmonary artery branches need careful attention during all neonatal echocardiograms. Once identified and repaired, outcome is good.

References