Anomalous left pulmonary artery without pulmonary artery sling

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An anomalous left pulmonary artery always runs posterior to the trachea causing a pulmonary artery sling compressing the trachea and oesophagus. We describe a unique case of an anomalous left pulmonary artery passing anterior to the trachea without producing the pulmonary artery sling associated with a right aortic arch and an aberrant left subclavian artery.

Case report
A 5 year old asymptomatic girl presented with a heart murmur. Echocardiography established a clinical diagnosis of a large secundum atrial septal defect. A right aortic arch was also noted. The origin of the left pulmonary artery was not seen from the pulmonary trunk and was subsequently seen to arise from the right pulmonary artery anterior to the trachea (fig 1). Oesophagography showed an impression on the right upper lateral margin by the right aortic arch with an additional posterior oblique indentation, consistent with an aberrant left subclavian artery. There was no anterior oesophageal indentation. The anomalous origin of the left pulmonary artery was confirmed by angiography (fig 2) and by surgical inspection during closure of the atrial septal defect.

Discussion
A pulmonary artery sling, a rare anomaly, is formed when the left pulmonary artery originates anomalously from the right pulmonary artery and passes between the trachea and the oesophagus. Embryogenesis of this anomaly is based on the evaluation of the reported cases. Variants of this anomaly have recently been described emphasising the developmental hypothesis of Jue and associates. A case report of a partial anomalous left pulmonary artery has been published where the branch of the left pulmonary artery...
pulmonary artery supplying the left lower lung lobe passed behind the trachea forming a sling. The branch for the left upper lung lobe originated in a normal fashion from the main pulmonary artery. In another case of a partially anomalous left pulmonary artery the arrangement of the left pulmonary artery was similar except that the branch of the left lower lobe pulmonary artery ran in front of the trachea producing no pulmonary artery sling.

The distal parts of the pulmonary arteries normally develop from the capillaries of their respective lung buds. These are then joined with the ipsilateral sixth aortic arches. The embryology of the anomalous left pulmonary artery passing dorsally to the trachea has been described as a failure of the left lung buds' connection with the left sixth arch. Instead, the connection of the left lung buds occurs with the right sixth arch dorsally to the trachea forming the pulmonary artery sling. If this connection runs ventrally no sling is formed. The presence of the right aortic arch and the aberrant left subclavian artery in our patient is important because this anomaly of the aortic arch is believed to be a consequence of the involution of the left fourth aortic arch and loss of the left sixth aortic arch. Because of the loss of the left sixth aortic arch it is plausible that the left lung buds could not connect to it. A connection with the right sixth aortic arch has thus formed creating the anomalous left pulmonary artery (fig 3). We think that the involution of the fourth left aortic arch and loss of the left sixth aortic arch, if occurring before or at the same time as the faulty connection of the left pulmonary artery, could explain the coexistence of these anomalies.

The two cases with the partially anomalous left pulmonary artery, one running anteriorly and the other posteriorly to the trachea, and our patient with the completely anomalous left pulmonary artery passing in front of the trachea in combination with the right aortic arch and the aberrant left subclavian artery, lend strong support for the developmental hypothesis of Jue and associates.

5 Huntington GS. The morphology of the pulmonary artery in the mammals. Anat Rec 1919;17:165–90.