Double Aortic Arch: An Uncommon Cause of Stridor in an Infant

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SUMMARY
A male infant six months of age presented with stridor. Echo, MRI, cardiac catheterization confirmed the diagnosis of double aortic arch. The child underwent successful surgery. The case is reported as an uncommon cardiac anomaly and an uncommon cause of stridor in an infant.

Keywords: Stridor, Double Aortic Arch, Subclavian Artery, Thoracotomy, Ligamentum Arteriosum, Aortopexy, Stenosis, Trachea

ETIOLOGY
The classical double aortic arch anatomy develops when involution of the distal right fourth arch does not take place. The fourth right and left arches both persist and form the left sided descending thoracic aorta. The right and left arches encircle the trachea and oesophagus. The double aortic arch then forms a ring around the trachea and oesophagus. The defect is rarely associated with other intra cardiac defect. Factors responsible for aberrant persistence of double arch segment have not been clearly identified. Band 22q11 deletion was found in 3 of 22 patients (14%) in double aortic arch.

Anatomy
In a double aortic arch both arches may be patent or an atretic segment may be present at one of the segments located in either arch. When both arches are patent one may be larger than the other or both may be similar in size.

In 75% of patient with double aortic arch the right arch is larger than the left. Associated cardiac anomaly, one fifth have other congenital cardiac defect; tetralogy of fallot, ventricular septal defect, coarctation of aorta, truncus arteriosus, patent ductus arteriosus, transposition of the great arteries.

Fig. 1 MRI larger Rt. Aortic and smaller Lt. Aortic Arch

Fig. 2 MRI Double Aortic Arch forming a complete circle

Fig. 3: Angiogram showing stenosis in Lt. Arch distal to origin of Lt. Subclavian Artery

Fig. 4: Angiogram showing small Lt. Arch in lateral view
The risk of the procedure is extremely low\(^4\). Outcome is usually good. However in some patients obstruction of the airway persists. Symptoms may improve after aortopexy is performed\(^5\). Robotic assistance division has been performed\(^6\), as well as video assisted thoracoscopic vascular ring division\(^7\).

**DISCUSSION**

The treatment of double aortic arch is undertaken by division of the non dominant arch after it is established as the cause of compression of the trachea. In our case the cardiac catheterization had shown stenosis distal to the origin of the left subclavian artery therefore the non dominant left arch was divided distal to the origin of the left subclavian to ensure that there is no compromise in blood flow. Thus the site of division is decided according to the anatomical variation as dominance of a arch, stenosis or atretic segment ensuring the blood flow to the head and neck vessels remains normal.

**CASE REPORT**

6 month old boy referred to us with stridor. Echo cardiography suggested double aortic arch. MRI Fig. 1 & 2 and cardiac catheterization Fig. 3& 4 confirmed the diagnosis. In cardiac catheterization it can be seen that there is stenosis distal to the origin of the left subclavian artery.

**Treatment:** Under general anaesthesia, central line and bilateral radial arterial line were placed. Left lateral thoracotomy was done the Lt Aortic Arch, Rt. Aortic Arch and Lt. Subclavian Artery were identified Fig. 5; the ligamentum arteriosus was divided after being ligated. The left arch was divided distal to the origin of the left subclavian artery and after minimal mobilization the trachea could be seen Fig. 6. Chest drain was placed. In ICU post extubation the stridor disappeared completely and the child was discharged home four days later.

**REFERENCES**

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