Double vascular ring: a case report of double aortic arch and concurrent pulmonary artery sling

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Background
Double aortic arch (DAA) and pulmonary artery sling (PAS) are vascular ring formations that present in neonates and infants with symptoms of respiratory stenosis.

Case summary
The patient was a girl with suspected ventricular septal defect (VSD), right aortic arch (AA), left patent ductus arteriosus, and bilateral superior vena cava (SVC) on foetal echography in the first day of life. The girl was delivered at 40 weeks and 4 days of gestation. Ventricular septal defect, DAA, coarctation of the left AA, and bilateral SVC were diagnosed. Contrast-enhanced computed tomography at Day 16 revealed PAS with concurrent anomalous tracheal branching in addition to DAA. The right A2 segmental artery, which supplies the right upper pulmonary artery, showed abnormal branching from the left pulmonary artery (LPA). At 3 months of age, VSD patching, left AA resection distal to the root of the left subclavian artery, arterial ligament dissection, and LPA replacement were performed.

Discussion
Pulmonary artery sling coexists with anomalous branching of the trachea and abnormal branching of the right pulmonary artery (RPA). Our patient had an extremely rare case of DAA concurrent with PAS and presented with anomalous tracheal and RPA branching. We were concerned that increased pulmonary blood flow caused by the VSD would exacerbate tracheal displacement. Radical surgery at 3 months of age resulted in good postoperative progress.

Keywords
Double aortic arch • Pulmonary artery sling • Vascular ring • Case report

Introduction
A double aortic arch (DAA) forms a vascular ring when either the left and right fourth pharyngeal arcuate arteries originating from the aorta do not regress. DAA surrounds the trachea and oesophagus, causing symptoms such as recurrent upper respiratory tract infections, wheezing, and dysphagia. A Pulmonary artery sling (PAS) is an extremely rare anomaly in which the left pulmonary artery (LPA)...

Learning points
- Double aortic arch can be concurrent with pulmonary artery sling, and we define two vascular rings overlap as double vascular ring.
- A case of double vascular ring has high risk of tracheal stenosis. Performing radical surgery in early infancy is likely to be effective.
originates from the right pulmonary artery (RPA),1 displaces the trachea, and causes symptoms of trachea and bronchial tree compression. We believe that this is the first reported case of a patient with concurrent DAA and PAS. Its interest derives from the achieving a good outcome with prompt surgical intervention.

### Timeline

| Event Description                                                                 | Date                  |
|----------------------------------------------------------------------------------|-----------------------|
| Foetal echography revealed findings suggestive of ventricular septal defect (VSD), right aortic arch (AA), left patent ductus arteriosus, and bilateral superior vena cava (SVC). | At 37 weeks and 6 days of gestation |
| The baby was born weighing 2662 g. Echocardiography led to a diagnosis of VSD, double aortic arch (DAA), coarctation of the left AA, and bilateral SVC. | At 40 weeks and 4 days of gestation |
| Computed tomography (CT) revealed DAA with concurrent pulmonary artery sling (PAS) and anomalous branching of the trachea. The right A2, which is part of the right upper pulmonary artery, showed abnormal branching of the pulmonary artery from left pulmonary artery (LPA). | At 16 days of age |
| Cardiac catheterization revealed Qp/Qs of 2.5, mean pulmonary artery pressure of 27 mmHg, and pulmonary vascular resistance of 1.5 WU m². Repeat CT examination revealed that the angle between the main pulmonary artery and LPA had decreased. | At 2 months of age |
| Ventricular septal defect patching, left AA resection distal to the root of the left subclavian artery, arterial ligament dissection, and LPA replacement were performed. | At 3 months of age |
| The patient has persistent inspiratory stridor when affected by a common cold; however, her progress is good with no achalasia. | At 1 year and 2 months of age |

### Case presentation

The patient was referred for treatment after polyhydramnios was detected on foetal ultrasound at 37 weeks and 4 days of gestation. Ultrasound also revealed duodenal obstruction and ventricular septal defect (VSD). Ultrasound at 37 weeks and 6 days of gestation were suggestive of VSD, right aortic arch (AA), left patent ductus arteriosus (PDA), and bilateral superior vena cava (SVC). The baby was born by normal delivery at 40 weeks and 4 days gestation. Her birth weight was 2662 g. Blood pressure was mmHg; heart rate was 110 b.p.m; respiratory rate was 40 per min, and SpO₂ was 97%. Clinical examination revealed that first heart sound and lung auscultation were normal. She didn’t have cyanosis and wheezing. Ultrasound led to a diagnosis of VSD, DAA, coarctation of the left AA, and bilateral SVC. Surgery at 1 day of age corrected the duodenal occlusion. Contrast-enhanced computed tomography (CT) at 16 days of age confirmed DAA (Figure 1) with a concurrent PAS (Figure 2A) and anomalous Wells classification IB tracheal branching (Figure 2B). Furthermore, the right A2, which is part of the right upper pulmonary artery (RUPA), showed abnormal branching of the pulmonary artery from LPA (Figure 3). Oesophagram at 18 days of age revealed the defect in the posterior wall of oesophagus at the level of DAA and the defect in the anterior wall at the level of PAS.

The patient developed mild inspiratory stridor at 2 months of age. There were no feeding issues, vomiting, and recurrent respiratory infections. Cardiac catheterization revealed a Qp/Qs of 2.5, mean pulmonary artery pressure of 27 mmHg, and pulmonary vascular resistance of 1.5 WU m². Repeat CT found that the angle between the main pulmonary artery (MPA) and LPA had decreased. Ventricular septal defect patching, left AA resection distal to the root of the left subclavian artery, arterial ligament dissection, and LPA replacement were performed at 3 months of age. LPA was detached from RPA and implanted into MPA with continuous sutures. She was treated by cardiopulmonary bypass through a median sternotomy. Postoperative extubation was followed by an uneventful recovery, and the patient’s progress was good.

Repeat CT examination at 9 months of age, 6 months after surgery, found LPA stenosis and residual stenosis at the site of tracheal bifurcation (Figure 5). At 1 year and 2 months of age, the patient has persistent inspiratory stridor when she was affected by a common cold. Achalasia is absent, and her progress is good.
Figure 2  (A) Left pulmonary artery originates from right pulmonary artery, forming pulmonary artery sling and surrounding the trachea. Right upper pulmonary artery supplying S1 + S3 (A1 + A3; black arrow) originates from right pulmonary artery, and right upper pulmonary artery supplying S2 (A2; yellow arrow) originates from left pulmonary artery. Right middle and lower pulmonary artery originate from right pulmonary artery (red arrow). (B) At the Th3 level, the right upper lobe bronchus branches alone. After the right upper lobe bronchus branches alone, the left and right bronchi branch and the trachea branching form takes on the shape of Wells classification Type IB. In the pulmonary artery sling area, the trachea is pressed by left pulmonary artery immediately above the tracheal bifurcation. DAA, double aortic aorta; dAo, descending aorta; L, left pulmonary artery; LAA, left aortic arch; M, main pulmonary artery; PAS, pulmonary artery sling; R, right pulmonary artery; RAA, right aortic arch; T, trachea.

Figure 3 Left pulmonary artery runs posterior to the trachea (arrowhead), and forms pulmonary artery sling. The right A2 supplying the posterior segment of the right lung (S2) shows abnormal branching of the pulmonary artery from left pulmonary artery. LPA, left pulmonary artery; MPA, main pulmonary artery; RPA, right pulmonary artery.

Figure 4 (A) Computed tomography at 16 days of age. (B) Computed tomography at 2 months of age revealed that the angle between main pulmonary artery and left pulmonary artery had decreased compared with that at 16 days of age. Tracheal displacement resulting from increased pulmonary blood flow and enlargement of the pulmonary artery caused by ventricular septal defect may have worsened the tracheal compression. LPA, left pulmonary artery; MPA, main pulmonary artery.
Discussion

A DAA forms a vascular ring when either the left and right fourth pharyngeal arcuate arteries originating from the aorta, or the eighth segment of the dorsal aorta, do not regress. DAAs account for about 42% of vascular rings and present in neonates and infants with respiratory symptoms, including stridor and respiratory distress, and dysphagia. The right AA is dominant in 80% of cases, the left AA is dominant in 10%, and the contribution is balanced in 10%. As the diagnostic modalities for vascular rings, Backer et al. prefer CT imaging to magnetic resonance imaging (MRI). CT imaging gives clearer picture of the tracheal anatomy and requires a shorter period to obtain than MRI. And they recommend an ultrasound in all patients with a vascular ring because there is a 12% incidence of associated cardiac pathology. Surgical repair is associated with no mortality in patients with DAA and 0-1.3% of these patients required reoperation.

A PAS develops when the LPA vascular bud forms between the oesophagus and trachea, originating from the RPA instead of the MPA anterior to the trachea. The LPA branches from the RPA and runs posterior to the trachea, thereby causing tracheal displacement. PAS accounts for 12% of vascular rings and presents in neonates and infants with symptoms such as stridor and respiratory distress. Postoperative mortality in patients with PAS is 12-21% and 0-12% of these patients required reoperation.

Wells et al. described four types of PAS (IA, IB, IIA, and IIB) depending on the level of tracheal bifurcation. Our patient had a Type IB PAS with a right upper lobe bronchus and tracheal bifurcation between the fourth to fifth thoracic vertebrae. This type of PAS is seen in 13% of cases. In this patient, we also observed abnormal branching of RPA from the right A2, which is a part of the RUPA, which originates from LPA. A retrospective evaluation of a series of PAS patients by Xie et al. found that the RUPA portion originated from the LPA in 10% of the Type IB cases. Consequently, we consider our case to be a rare type of PAS.

The coexistence of a PAS, right AA, and aberrant left subclavian artery in cases with two overlapping, or double vascular rings, has been previously reported. There are no previous reports of patients with concurrent DAA and PAS as seen in our patient. Although this condition has a very high risk of tracheal stenosis, this patient presented with relatively mild symptoms, which was attributed to the presence of a Type IB PAS. However, the angle of MPA and LPA decreased, raising concerns that an increase in the pulmonary blood flow would worsen the tracheal displacement. In such cases, radical surgery for VSD in early infancy is likely to be effective.

Figure 5 (A) Preoperative computed tomography (at 16 days of age). (B) Preoperative computed tomography revealed anteroposterior oppression of the tracheal bifurcation (arrowhead) caused by pulmonary artery sling. (C and D) Computed tomography at 6 months postoperatively (at 9 months of age) revealed left pulmonary artery stenosis and residual stenosis at the site of tracheal bifurcation. DAA, double aortic arch; LPA, left pulmonary artery; PAS, pulmonary artery sling.
Conclusions

To our knowledge, this is the first report of a vascular ring with concurrent DAA and PAS. In this patient, it was inferred that the coexistence of the two conditions carried a high risk of tracheal stenosis. Surgical intervention was performed early and resulted in a good outcome with no complications such as significant respiratory symptoms or dysphagia.

Supplementary material

Supplementary material is available at European Heart Journal - Case Reports online.

Slide sets: A fully edited slide set detailing this case and suitable for local presentation is available online as Supplementary data.

Consent: The author/s confirm that written consent for submission and publication of this case report including image(s) and associated text has been obtained from the patient in line with COPE guidance.

Conflict of interest: none declared.

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