Congenital fistula between the aorta and left atrium

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SUMMARY A child in whom abnormal vessels connected the descending thoracic aorta and the left atrium is described. This is a previously unreported congenital malformation. Clinically this condition differs from the more common aorto-cardiac fistulas in that the continuous murmur was better heard posteriorly as well as in the right parasternal area. Ligation of the aberrant vessels abolished the murmur.

Aorto-cardiac fistulas are well known. Aneurysms of the sinuses of Valsalva, usually of the right sinus, may rupture into the right ventricle, right atrium, or rarely the left atrium. The communications are rarely congenital. They rupture either spontaneously or as a result of infection, trauma, or operation involving the aorta or aortic valve.

An aorto-left ventricular tunnel is a congenital anomaly in which a defect in the aortic wall provides a tunnel opening on to the left ventricle via the septum bypassing the aortic valve. Coronary cameral fistulas are usually congenital and caused by an abnormal connection between a main or branch coronary artery and one of the cardiac chambers.

We report a patient in whom there were congenital fistulas between the abnormal branches of the descending thoracic aorta and the left atrium. We found no published reports of similar cases.

Case report

A four year old Iranian boy was admitted for investigation of a cardiac murmur. He was born after a normal pregnancy and was healthy until he had a "gland" infection at the age of two. Then a heart murmur was discovered.

On examination he was of normal size and build for his age, apart from a precordial bulge caused by an enlarged heart. He was intelligent and cooperative. He was cyanotic with no clinical signs of heart failure. The arterial pulses were regular and symmetrical; however, they were of high volume and ill-sustained. Blood pressure was 90/40 mm Hg. The venous pressure was normal. The left ventricle was greatly enlarged and showed a hyperkinetic impulse. A very loud continuous murmur was heard on auscultation. This was best heard posteriorly to the right of the spine at the level of T5 and T6, but it was also heard widely over the back and much more softly anteriorly. There was an apical third heart sound. The electrocardiogram showed sinus rhythm with voltage criteria typical of left ventricular enlargement. Chest x-ray showed considerable cardiomegaly and a large aortic knob. The main pulmonary artery was not enlarged and there was no evidence of a left to right shunt.

Cross sectional echocardiography showed situs solitus with normal intracardiac relations. There were no valvar abnormalities. The left ventricle was enlarged and contracting vigorously. The left atrium was also dilated, whereas the right heart chambers were normal. The ventricular septum was intact. There was no evidence of coarctation. The ascending aorta and the horizontal portion of the arch were enlarged and excessively pulsatile as far as the isthmus. At the isthmus a large vessel appeared to emerge from the descending aorta. This was taken to be a ductus arteriosus even though its continuity with the pulmonary artery was not visualised on the echocardiogram. Because the clinical findings were incompatible with a ductus arteriosus, cardiac catheterisation and angiography were performed. There was no pressure gradient across the valves. An aortogram showed a competent aortic valve and no ductus arteriosus. After the arch of the aorta became opacified the left atrium filled promptly through a large channel which appeared to lie posteriorly and parallel to the descending aorta (figs 1 and 2). A
idly tapered down to about half the size of the arch. There was a very strong continuous thrill in this area. After the third part of the arch and the descending thoracic aorta had been fully dissected and mobilised, it was possible to see a very large vessel passing directly to the right from the descending thoracic aorta just where it started to narrow and there was a further vessel about half it diameter immediately above it. Because the thrill only disappeared after trial occlusion of both aberrant vessels, they were evidently in direct communication. Both vessels were doubly ligated and this abolished the thrill. The patient’s postoperative course was uneventful. Figure 2 is a compositive diagrammatic representation of the angiographic, computed tomographic, and operative findings.

Discussion

Aorto-cardiac communications are not uncommon in children and neonates. The large gradient between the high pressure aorta and low pressure cardiac chamber causes a high flow for which the heart must compensate to maintain adequate perfusion. Most congenital aorto-cardiac fistulas communicate between one of the coronary sinuses, usually the right, and the right ventricle, right atrium, and much more rarely the left ventricle. These fistulas are often associated with other congenital heart lesions. Jaen et al described a patient with

fistula between the aorta and left atrium was diagnosed. Because neither the angiogram nor the echocardiogram had indicated the precise connections of this fistula the patient was examined by computed tomodraphy. This showed an anomalous vessel adjacent to the right of the descending thoracic aorta which appeared in images immediately below the arch of the aorta at a point where the calibre of the aorta abruptly diminished. This structure filled with the contrast to the same degree as the aorta and was of a similar size (fig 3).

Laboratory investigations were normal apart from an iron deficiency anaemia (haemoglobin 9.1 g/dl). The tuberculin test was negative.

Operation was performed through a left thoracotomy incision. The aortic arch was large and at the base of a large left subclavian artery there was an abnormal vessel of the same diameter as the left subclavian artery. After about 1.5 cm this anomalous vessel petered out into a fibrous chord and disappeared into the mediastinum in the region of the oesophagus. About 2 cm below the origin of the left subclavian artery the descending thoracic aorta rup-

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Fig 1 Contrast aortogram in the left anterior oblique projection showing the abnormal vessel connecting the descending thoracic aorta with the left atrium (arrows). Note also the diminishing calibre of the descending thoracic aorta.

Fig 2 Diagrammatic representation of the aortic arch viewed from the left lateral projection with the pulmonary artery removed for clarity (see text). IVC, inferior vena cava; LA, left atrium; LSC, left subclavian artery; PA, pulmonary artery; RSPV, right superior pulmonary vein.
Fig 3  Computed tomographic scan at the level of the heart (a) and great vessels (b, c). After contrast injection (b, c) the abnormal vessel became opacified simultaneously with the ascending and descending aorta (b) and the transverse portion of the arch (c). The calibre of the descending thoracic aorta at the level of the heart (a) is half the size of the arch (c).
communication between the ascending aorta, 3 mm above the origin of the left coronary artery, and the left atrium, which led to the formation of four large aneurysmal sacs.9

After birth, acquired communications occur either as varicose aneurysms or as direct arteriovenous anastomoses caused by trauma, operation, or infection.8 Apart from the unspecific history of a gland infection and the site of communication in our patient there is nothing to suggest that this communication may have been developed after birth. The smoothly communicating vessels with the presence of a second similar but much smaller vessel, the lack of fibrous tissue or adhesions, and the young age of our patient all indicate that these were anomalous branches of the aorta communicating directly with the left atrium. These vessels arose where one would expect the right sided intercostal vessels to arise from the descending thoracic aorta. The direct communication with the left atrium, however, is hard to explain embryologically.

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References