Pulmonary atresia with intact ventricular septum and corrected transposition of the great arteries

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SUMMARY We report an 82-day-old infant with a rare combination of anomalies: pulmonary atresia associated with corrected transposition of the great arteries and intact ventricular septum; this is the second such recorded case.

By using echocardiography, chest radiography, and electrocardiography the correct anatomical diagnosis was obtained before invasive investigation and necropsy.

An association of pulmonary valve atresia with intact ventricular septum and corrected transposition of the great arteries is a rare occurrence. We present a second such case following that reported by Steeg et al. in 1971. In our case, non-invasive procedures, especially echocardiography, were diagnostic and were confirmed at necropsy as well as by angiography.

Case report

An 82-day-old Japanese female infant was admitted to the Heart Institute, Tokyo Women's Medical College, with a complaint of mild cyanosis since 4 days of age. She was the product of a 42-week uneventful gestation except for mild toxemia in a 32-year-old mother. The birthweight was 2940 g. The presence of congenital heart disease was first confirmed at 57 days of age. However, the general condition including the respiratory status was described as good.

The physical examination disclosed a well-developed infant with slight generalised cyanosis, tachycardia, and tachypnoea. The liver was palpable 2 cm below the costal margin. The peripheral pulses were normal. The first heart sound was of normal intensity, the second heart sound was single and of increased intensity. A grade 1 ejection murmur and click were heard in the fourth intercostal space at the left sternal border.

The haematocrit was 51 per cent and the oxygen tension of capillary blood was 34 mmHg; the base excess was -8.3 and the carbon dioxide tension 34 mmHg.

The electrocardiogram (Fig. 1) showed normal sinus rhythm and a frontal plane mean QRS axis of +80 degrees with peaked P waves in leads II, aVR, and aVF. The Q wave in lead III was deep and the R wave in lead V1 was 0.4 mV. A left ventricular dominant pattern was seen.

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\frac{RV1+SV5}{SV1+RV5} < 1.0
\]

Q waves were absent in the left precardial leads and lead aVL. Because of these findings corrected transposition was suspected.

The chest x-ray film showed situs solitus, a cardiothoracic ratio of 62 per cent, and severely decreased pulmonary vascularity. The left upper cardiac border was straight.

The echocardiogram (Fig. 2) disclosed a small atrioventricular valve on the right side, and a large atriocentric valve was detected on the left side. Only one great artery (the aorta) was detected, and an infundibulum was interposed between the aorta and the large left-sided atrioventricular (tricuspid) valve.

At cardiac catheterisation both venules were entered, the left-sided systemic (anatomically right) ventricle was entered via an interatrial communication. Radiopaque contrast material injected into this left-sided ventricle filled the aorta arising anteriorly via an infundibulum along the left heart border. Pulmonary blood flow was supplied via a persistent ductus arteriosus; the main pulmonary artery was not filled. Right-sided (anatomically left) ventriculography showed a hypoplastic chamber with no direct communication to the great arteries or
to the other ventricle. Dilated myocardial sinusoids in the wall of this chamber were opacified.

From these findings, we made a diagnosis of either pulmonary atresia with intact ventricular septum and corrected transposition of the great arteries, or pulmonary atresia with intact ventricular septum and double outlet right ventricle with atrioventricular discordance.

After cardiac catheterisation, a right-sided subclavian artery-pulmonary artery anastomosis and atrioseptostomy were performed but the infant died on the first day after operation.

**PATHOLOGY**

The right-sided (anatomically left) ventricle was small; the left-sided (anatomically right) ventricle was large and occupied about four-fifths of the whole anterior aspect. There was a single left coronary artery. The aorta arose anteriorly via an infundibulum from the left-sided right ventricle. The main pulmonary artery was vestigial, located right posterior to the aorta, and was above the right-sided left ventricle. Fig. 3A shows the right atrium and right-sided left ventricle. The volume of this ventricle was peanut-size with hypertrophied...
myocardium, measuring 1.2 cm in thickness. There were fine trabeculations and a smooth septal surface in this chamber. The right-sided atriorecordsvalve was a normal miniature of the mitral valve. No pulmonary outlet from this chamber was present and the ventricular septum was intact. The left-sided right ventricle had a large cavity and coarse trabeculation (Fig. 3B). The left-sided atriorecordsvalve was tricuspid. The atrial septal defect produced by atrioseptostomy was visible at the interatrial septum. The right-sided subclavian artery-pulmonary artery anastomosis was also recognised. Thus the pathological diagnosis was pulmonary atresia with intact ventricular septum and corrected transposition of the great arteries.

Discussion

Corrected transposition of the great arteries seldom exists without associated cardiac malformations.\(^1\) The most common associated cardiac malformations are malpositions of the cardiac apex (isolated dextrocardia, mesocardia, and mirror image dextrocardia), ventricular septal defect, pulmonary stenosis, anomalies of the left atriorecordsvalve (including regurgitation, stenosis, and atresia), and Ebstein’s anomaly.\(^3\)\(^4\) Pulmonary stenosis (or atresia) together with a ventricular septal defect is often seen in corrected transposition,\(^5\)\(^6\)\(^7\) but pulmonary stenosis with intact ventricular septum is rare in corrected transposition.\(^5\)\(^6\)\(^7\) Furthermore, pulmonary atresia with intact ventricular septum and corrected transposition of the great arteries is very rare, only one such case other than ours having been previously reported by Steeg et al.\(^1\)

In our case, the clinical features were similar to those of pulmonary atresia with intact ventricular septum not associated with corrected transposition, and the differential diagnosis was at first clinically difficult. The reasons we suspected the correct diagnosis before cardiac catheterisation and angio取消graphy are as follows. (1) The electrocardiogram showed absence of Q waves in leads I, aVL, and left precordial leads, and there were deep Q waves in lead III.\(^8\) (2) On x-ray examination of the chest, the left upper cardiac border was straight. (3) The echocardiogram indicated absence of fibrous continuity between a left-sided atriorecordsvalve valve (tricuspid valve) and the aortic valve (=presence of subaortic infundibulum). The correct anatomical diagnosis was verified by angiocardiography. Absence of Q waves over the left precordial leads in the electrocardiogram was noted in both our case and that reported by Steeg et al.\(^1\) The echocardiographic finding of the absence of fibrous continuity between the left-sided atriorecordsvalve valve (tricuspid valve) and the aorta in our case was diagnostically helpful in establishing atrioventricular discordance.\(^9\)

With meticulous application of non-invasive
techniques, correct anatomical diagnosis can be obtained even in such rare conditions as our present case.

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References


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