Hypoplastic right heart complex in a 46-year-old woman

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The case is reported of a 46-year-old woman with congenital right ventricular hypoplasia, severe tricuspid stenosis, pulmonary hypoplasia, and an intact ventricular septum. This cardiac malformation is extremely rare in adults. Pulmonary circulation was augmented by bronchial collaterals.

The recently designated 'hypoplastic right heart complex' (Khoury et al., 1969), consisting of various degrees of right ventricular hypoplasia, congenital tricuspid stenosis, and pulmonary stenosis, is extremely rare beyond childhood (Sackner et al., 1961; Williams, Barratt-Boyes, and Lowe, 1963). This report is of a woman who survived 46 years with this complex.

Case report
The patient had been cyanotic since birth. She was first evaluated at Duke University Medical Center when she was 20 years old at which time her heart was slightly enlarged but no murmurs were heard. At the age of 24 she developed pelvic thrombophlebitis which resulted in paradoxical emboli and an acute diaphragmatic myocardial infarction. Shortly thereafter she began to have occasional chest pain, but she did not require cardiac medication.

Deterioration of cardiac function, in fact, did not begin until six months before death when the chest pain became more frequent and signs of congestive heart failure appeared. A phonocardiogram one month before death disclosed an apical systolic murmur, a widely split second heart sound which moved slightly with respiration, and an early pulmonary ejection murmur.

Three weeks later the patient was admitted to hospital with congestive heart failure. Her blood pressure was 140/80 mm. Hg, and her pulse was irregular at 130 a minute. She was cyanotic with clubbing of her fingers and toes. The point of maximum cardiac impulse was diffuse in the sixth intercostal space at the anterior axillary line. A grade 3/6 blowing, mid-systolic murmur radiated from the apex to the left axilla. There was fixed splitting of the second sound along the left sternal border. No gallops were heard. Her haematocrit was 56 per cent.

An electrocardiogram indicated right axis deviation, biatrial enlargement, premature ventricular contractions, poor R wave progression, Q waves in leads II, III, and aVF, and T wave inversion in leads II, III, aVF, V5, and V6. Radiological examination of the heart showed left ventricular and biatrial enlargement. The pulmonary vasculature was diminished. There was evidence, however, of prominent bronchial circulation.

Cardiac catheterization revealed bidirectional shunting across a large atrial septal defect. Mean pressures in the right and left atria were 20 and 21 mm. Hg, respectively, with a left ventricular pressure of 115/4-22 mm. Hg. The catheter could not be passed through the tricuspid valve. Oxygen saturation was 56 per cent in the inferior vena cava, 70 per cent in the right atrium, 79 per cent in the left atrium, 77 per cent in the left ventricle, and 80 per cent in the aortic root.

Angiocardiography showed sequential opacification of the right atrium, left atrium, left ventricle, and the aorta. The right ventricle and a small pulmonary artery could be detected only on the last films of left ventriculography.

After injection of the contrast material the patient suddenly became unresponsive and more cyanotic. Three days later she died in ventricular fibrillation.

Necropsy findings The heart weighed 480 g. The great vessels arose normally, but the pulmonary artery was small and the aorta was dilated. An atrial septal defect, 2 cm. in diameter, was present. The ring of the tricuspid valve was hypoplastic, 6-9 cm. in circumference (Fig.). The valvular cusps had apparently never separated, so that a single circumferential leaf was present, leaving a stenotic orifice 1 cm. in diameter. The right ventricle was small. Though the pulmonary valve was hypoplastic with a ring circumference of 4-1 cm., the valve leaflets appeared normal.
septal defect and enough pulmonary outflow obstruction to prevent hypertension (Castleman and McNeely, 1969; Jordan and Sanders, 1966). However, we could find two case reports of right ventricular hypoplasia coupled with stenosis or hypoplasia of the tricuspid valve without interventricular communication. One patient was a 23-year-old man (Sackner et al., 1961), the other a 39-year-old man (Popper, Kushner, and Gasul, 1956). Similarly, the patient we are describing had no ventricular septal defect. Apparently the severe obstruction to pulmonary blood flow was largely overcome in this case through collateral bronchial circulation. Though evidence of these collaterals was not found at necropsy, they were shown by x-ray examination.

This patient’s long survival was especially remarkable because her myocardium had been weakened 22 years before death by a large diaphragmatic infarction.

References

Notice
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