BICUSPID PULMONARY VALVE IN ASSOCIATION WITH CALCIFIC AORTIC STENOSIS

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The aetiology of aortic stenosis in the presence of a normal mitral valve is obscure, the major possibilities being old rheumatic infection and calcification of a congenitally abnormal valve. The present report describing a healthy pulmonary valve with only two cusps occurring in association with a typical extreme calcific aortic stenosis, therefore, throws some light on this subject.

Case Report. A married woman of 56 had a history of 5 years of increasing breathlessness. Eighteen months before she died she had a pain in the calf and chest, and a year later swelling of the ankles first appeared. She next developed pains in the abdomen and in the chest and then had transient congestive cardiac failure. Two weeks before death there was more chest pain and retention of urine; this contained protein and urobilinogen. The blood urea rose to 153 mg. per 100 ml. while the serum sodium and chloride fell to 290 mg. and 455 mg. respectively. Her E.S.R. was 21 mm. in the first hour (Wintrobe). She died in uremic coma.

Necropsy. There was slight icterus and some dependent edema. The heart weighed 575 g. and was covered with a fibrinous exudate from a small effusion. This cardiomegaly was due to hypertrophy of the left ventricle (average thickness 28 mm.) and some dilatation of the right ventricle (5 mm.) and atrium. The tricuspid and mitral valves were somewhat opaque; the chordae of the former were normal while those of the latter showed some thickening and shortening but no matting or grouping to suggest rheumatism. Both orifices were normal. The most significant abnormality was that the pulmonary valve had only two cusps, the posterior one being slightly larger and bearing at its mid point a transverse ridge-like elevation. They were quite transparent and there was no fibrosis. The circumference was 75 mm. at the level of the commissures. The aortic valve was grossly diseased being irregularly calcified and ulcerated so that it formed a dome-shaped protrusion into the aorta. Its slit-like orifice was nearly antero-posterior and measured 19 x 7 mm.; the front end abutted the nodule on the posterior pulmonary cusp. It seemed originally to have been bicuspid, too, its right-hand cusp bearing a nodule similar to that on the pulmonary valve. The right coronary artery arose from two orifices 2 mm. apart in the anterior part of the right sinus in front of the nodule. There was a small mural thrombus in the right atrium.

There were multiple organizing pulmonary infarcts with effusions and evidence of chronic passive congestion in the other organs. Organizing thrombi of considerable age were present in the edematous calf of the right leg. There were no histological abnormalities in the myocardium, nor, apart from congestion and infarction, in the lungs. No organic cause for the uremia was found.

Discussion

Aortic valves apparently composed of only two cusps are relatively common and though they are generally found to be stenosed there is little direct evidence of the cause of this distortion. Neither rheumatic infiltrations diagnosed histologically (Karsner and Koletsky, 1947) nor a clinical history reaching back into childhood (Campbell and Kaoutzze, 1953) are more than suggestive. Bicuspid malformation of the pulmonary valve, on the other hand, is extremely uncommon and a healthy bicuspid valve in a heart that does not have some fairly gross abnormality of septa or origin of great vessels may well be unique. Most examples occur in some form of the tetralogy of Fallot. 7 of the first 18 cases in the Guy's Hospital series of necropsies showing this abnormality: one of these had also aortic stenosis (Campbell and Brinton, 1953). In isolated pulmonary valvular stenosis apparent bicuspid malformation of the contracted pulmonary valve is much less common, none of 14 consecutive cases showing it, but one of these had a bicuspid aortic valve (Campbell, 1954).

Since a bicuspid pulmonary valve is usually associated with some other error of cardiac...
development the associated abnormality in this heart was probably congenital in origin. This was, however, typical of the sort of aortic malformation that would, if present alone, commonly be considered to have developed after birth. This heart is described as a piece of evidence supporting the suggestion that many of these lesions, although first responsible for symptoms late in life, are in fact congenital in origin.

Summary

A case of congenital bicuspid malformation of a normal pulmonary valve in the absence of any other abnormality of the septa or orientation of the great vessels but associated with gross calcific aortic stenosis is described. The aortic malformation is apparently superimposed on a bicuspid form of that valve and it is suggested that the present case is evidence that some of these distorted aortic valves are congenital in origin rather than acquired.

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References