Shone’s complex with dextrocardia and situs inversus totalis: a case report

Dekstrocardi ve situs inversus totalisle birlikte görülen Shone kompleksi: Olgu sunumu

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Summary– Parachute mitral valve complex is an unusual congenital anomaly that has been described by Shone et al. It is characterized by a parachute deformity of the mitral valve associated with additional forms of left heart anomalies, such as aortic valvular stenosis and coarctation of the aorta. A 21-year-old female who was referred to our department because of progressive dyspnea on effort and at rest and minimal cyanosis is presented in this case report. On cardiac auscultation, the patient had a grade III/VI pansystolic murmur best heard at the lower left sternal border. The chest X-ray demonstrated dextrocardia and mild cardiomegaly. Echocardiographic evaluation revealed Shone’s complex, including parachute mitral valve anomaly.


The finding of a parachute mitral valve (PMV) in an adult patient is extremely rare. Most patients are diagnosed in childhood with the pathologic complex known as Shone’s anomaly, which includes a PMV and aortic coarctation (68% of patients), atrial septal defect (54% of patients), ventricular septal defect (46% of patients), aortic valve stenosis (32% of patients), subaortic stenosis (20% of patients), and left ventricular hypoplasia (19% of patients).[1] Fewer than 50 cases of Shone’s complex have been reported in the literature, and it has previously only been observed in children.[2]

CASE REPORT

A 21-year-old female patient was referred to our hospital for evaluation of dyspnea, a systolic murmur, and minimal cyanosis. She experienced progressive dyspnea on effort and at rest. Cyanosis had been diagnosed when she was 2 years old but her family did not seek medical attention for socio-economic reasons. Cyanosis was not apparent at rest. She had no history of smoking or significant pulmonary disease history before this diagnosis. On cardiac auscultation, the patient had a grade III/VI pansystolic murmur best heard at the lower left sternal border, with radiation to the right lower sternal border. The electrocardiogram revealed sinus rhythm, right ventricular hypertrophy, and sinus tachycardia with marked right axis deviation. A chest X-ray showed dextrocardia and mild cardiomegaly with increased pulmonary vascularity. Ventilation/
perfusion imaging and spiral computed tomography excluded pulmonary embolism. Transthoracic echocardiography revealed dextrocardia. Moreover, the morphological right atrium was on the left side of the heart, and the morphological left atrium was on the right side of the heart. Concordance atrioventricular connection, PMV, supramitral membrane, insertion of all the mitral valve’s chordae tendineae into a single papillary muscle (arrow) (B) rudimentary left ventricle, (C) double outlet right ventricle, mild pulmonary stenosis, (D) and ventricular septal defect were observed. PMV: Parachute mitral valve; LA: Left atrium; RA: Right atrium; LV: Left ventricle; RV: Right ventricle; VSD: Ventricular septal defect; Ao: Aorta; PA: Pulmonary artery.

Figure 1. (A) Concordance atrioventricular connection, supramitral membrane, insertion of all the mitral valve’s chordae tendineae into a single papillary muscle (arrow) (B) rudimentary left ventricle, (C) double outlet right ventricle, mild pulmonary stenosis, (D) and ventricular septal defect were observed. PMV: Parachute mitral valve; LA: Left atrium; RA: Right atrium; LV: Left ventricle; RV: Right ventricle; VSD: Ventricular septal defect; Ao: Aorta; PA: Pulmonary artery.

In 1963, Shone et al.\cite{3} originally described the developmental complex that included PMV, supravalvular ring of the left atrium, subaortic stenosis, and coarctation of the aorta and reported that the degree of mitral valve involvement appeared to be the main factor determining outcome in these patients. This hypothesis has been confirmed in two reports of the long-term surgical outcome in patients with multiple left heart obstructions and mitral valve involvement.\cite{4,5}
This condition is characterized by insertion of all the chordae tendineae into a single papillary muscle group. The chordae are generally shortened and thickened, and the anatomy of the papillary muscles is highly variable. The anterolateral, or rarely both papillary muscles, may be completely absent; or two identifiable but partially fused papillary muscles may be present. Commonly associated conditions include isolated or combined supramitral ring, aortic subvalvar or valvular stenosis, coarctation of the aorta, or the complete ‘Shone complex’ of anomalies. Right ventricular outflow tract obstruction has also been described. Median longevity approaches 10 years of age, correlating best with left ventricular size and poorly with mitral valve orifice size.

Patients with a PMV who reach adulthood usually present with dyspnea. PMV patients have a wide spectrum of hemodynamic anomalies of transmitial flow. However, PMV may be incidentally diagnosed during echocardiography with normal hemodynamics across the mitral valve. Medical or surgical treatment is not indicated in such patients. Patients with hemodynamically significant stenosis or regurgitation need to undergo mitral valve replacement or repair. Only hemodynamically significant and/or symptomatic associated congenital cardiac lesions should be surgically corrected.

The most interesting aspect of this case is that the patient reached adulthood without receiving any medical therapy and was diagnosed in the third decade of her life. Clinicians must be alert for the other components of Shone’s complex when they encounter a mitral valve anatomy consistent with PMV.

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REFERENCES


Key words: Mitral valve; parachute mitral valve; Shone’s complex.

Anahtar sözcükler: Mitral kapak; paraşüt mitral kapak; Shone kompleksi.