Late Onset of Symptoms of Ebstein's Malformation
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INTRODUCTION

Ebstein's anomaly is an abnormality of the tricuspid valve in which the septal leaflets and often the posterior leaflets are displaced into the right ventricle and the anterior leaflet is usually malformed, excessively large, and abnormally attached or adherent to the right ventricular free wall. Thus, a portion of the right ventricle is "atrialized" in that it is located on the atrial side of the tricuspid valve, and the remaining functional right ventricle is small. The tricuspid valve is usually regurgitant but may be stenotic. Eighty percent of patients with Ebstein's anomaly have an interatrial communication (atrial septal defect or patent foramen ovale) through which right-to-left shunting of blood may occur. Depending on the severity of the hemodynamic derangements in patients with Ebstein's anomaly that is determined by the degree of displacement and the functional status of the tricuspid-leaflet leaflets, clinical presentation of Ebstein's anomaly varies from severe heart failure in a fetus or neonate to the absence of symptoms in an adult in whom it is discovered incidentally. Late onset of clinical symptoms despite severe tricuspid-leaflet displacement with valvular dysfunction and elevated right atrial pressure is rare. The article describes a 50 years old man with late onset of symptoms with at least moderate tricuspid-leaflet displacement and right valvular dysfunction.

CASE REPORT

A 50 year old man was admitted to hospital because of dyspnea, atypical chest pain and fatigue lasting for 3 months. On physical examination the heart rate was 80 per minute, and the first and second heart sounds were widely split. The blood pressure was 115 / 75 mm Hg. A systolic murmur was present at the left lower sternal border. Hepatomegaly was present. ECG revealed sinus rhythm and intermittent atrial flutter with 2:1 conduction and a ventricular rate of 136 beats per minute. Transthoracic echocardiography was inconclusive because of poor image quality. Transesophageal echocardiography (Figure; arrow indicates tricuspid valve) showed right ventricular dysfunction and severe tricuspid regurgitation. The tricuspid valve leaflets did not attach normally to the valve annulus, and the effective orifice was displaced downward into the right ventricular cavity at the junction of the inlet and trabecular components of the right ventricle consistent with the diagnosis of Ebstein's malformation. Clinical presentation of Ebstein's anomaly with symptoms occurring first in adulthood despite moderate to severe tricuspid-leaflet displacement with valvular dysfunction and elevated right atrial pressure is rare.

These findings were consistent with the diagnosis of Ebstein's malformation. Only the septal and posterior leaflets were displaced and divided the right ventricle into two portions. The inlet portion was integrated functionally with the right atrium ("atrialized portion"), while the other, including the trabecular and outlet portions, constitutes the functional right ventricle. Cardiac catheterization using an electrode catheter and monitoring the intracardiac electrogram and the intracardiac pressure simultaneously confirmed the diagnosis. The "atrialized" part of the right ventricle was depolarized like the right ventricle but showed right atrial pressure.

In the patient described, control of intermittent atrial flutter could be achieved pharmacologically and clinical symptoms significantly improved with diuretic therapy and digoxin.
The patient refused surgery. 18 months later the patient was still in stable condition with mild impairment of exercise tolerance.

**DISCUSSION**

The severity of the hemodynamic changes in patients with Ebstein's anomaly depends on the degree of displacement and the functional status of the tricuspid-valve leaflets. Patients with mild apical displacement of the tricuspid leaflets have normal valvular function, whereas those with severe tricuspid-leaflet displacement or abnormal anterior leaflet attachment, with valvular dysfunction, have elevated right atrial pressure and right-to-left interatrial shunting. Similarly, the clinical presentation of Ebstein's anomaly varies from severe heart failure in a fetus or neonate to the absence of symptoms in an adult in whom it is discovered incidentally. Clinical presentation of Ebstein's anomaly with symptoms occurring first in adulthood despite moderate to severe tricuspid-leaflet displacement with valvular dysfunction and elevated right atrial pressure, as found in the patient described here, is rare.

Echocardiography is used to assess right atrial dilatation, anatomical displacement and distortion of the tricuspid-valve leaflets, and the severity of tricuspid regurgitation or stenosis.

Patients with Ebstein's malformation usually have a reduced life expectancy and seldom reach the age of 50. Cardiac arrhythmia are common in patients with Ebstein's anomaly, in the vast majority, of the tachycardia type. In adults with Ebstein's anomaly, the most important predictors of outcome are the New York Heart Association (NYHA) functional class, the heart size, the presence or absence of cyanosis, and the presence or absence of paroxysmal atrial tachycardias. The management of Ebstein's anomaly centers on the prevention and treatment of complications. Prophylaxis against infective endocarditis is recommended. Patients with symptomatic heart failure, as the patient described in this case report, are given diuretic agents and digoxin. Those with atrial arrhythmias may be treated pharmacologically or with catheter ablation.

Repair or replacement of the tricuspid valve in conjunction with closure of the interatrial communication, if present, is recommended for older patients who have severe symptoms despite medical therapy. In addition, repair or replacement should be considered for patients with less severe symptoms who have cardiac enlargement, since this condition has a poor prognosis.

In the patient described here, control of intermittent atrial tachycardia could be achieved using digoxin and β-blockers and clinical symptoms significantly improved with diuretic therapy. The patient refused surgery and was still in stable condition.

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