Ebstein’s Anomaly with Coarctation of the Aorta. An Unusual Association

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Ebstein’s anomaly with coarctation of the aorta is an extremely unusual condition. In this report, the clinical and surgical features of 3 male patients, aged 7 months, 4 years and 14 years, are discussed. All patients were in situ solitus. The first 2 patients had atrioventricular and ventriculoarterial discordance and progressed to heart failure in the neonatal period. The third had atrioventricular and ventriculoarterial concordance, as well as Wolf-Parkinson-White syndrome, with frequent episodes of paroxysmal tachycardia.

The 3 patients underwent surgery for correction of the coarctation of the aorta. The patient with atrioventricular and ventriculoarterial concordance underwent tricuspid valvuloplasty using a DeVega-like technique. In addition, ablation of 2 anomalous pathways (Kent bundle), which were detected by the electrophysiologic study, was also subsequently performed.

The 3 patients showed a good postoperative outcome for 2 years, although, in those with discordance, the surgical procedure did not influence the dysplasia of the tricuspid valve, because this valve showed light to moderate dysfunction.

Congenital anomalies characterized by tricuspid valve stenotic lesions or regurgitation, alone or combined with other abnormalities, are unusual. The most frequent or involving this valve is the so-called Ebstein’s anomaly, first described by Wilhelm Ebstein in 1866.1

This anomaly has a prevalence of 0.5% among the different congenital heart diseases. Studies show that lesions such as pervious foramen ovale and atrial septal defect (ASD) are almost always present. Other types of associations, such as stenosis or atresia of the pulmonary valve, ventricular septal defect and transposition of the great arteries, are uncommon.2,3 However, the association of Ebstein’s anomaly with coarctation of the aorta is described as extremely rare, mainly when ventriculoarterial concordance is present.4

This report describes 3 individuals with Ebstein’s anomaly and coarctation of the aorta, 2 of them in discordance and the remaining 1 in atrioventricular and ventriculoarterial concordance (AV).

Report of the Cases

Patients in atrioventricular and ventriculoarterial discordance - Case 1 - A 7-month-old infant with a past medical history of dyspnea on strenuous effort and a heart murmur detected on the 10th day of life. The physical examination showed an infant in good health, with slight tachypnea and without cyanosis. His heart rate (HR) was 160bpm; his systemic blood pressure in the upper limbs was 110/70mmHg. His lungs showed no rales.

On physical examination, visible pulsations at the precordium were noted and his heart auscultation showed a split 2nd heart sound of increased intensity, as well as a moderate systolic murmur in the mitral area, radiating to the axilla. His liver was felt 4cm below the right costal margin. He had a strong pulse in the upper limbs and no pulse in the lower limbs.

His chest X-ray showed a moderately enlarged cardiac silhouette, as well as an increased pulmonar vasculature. The electrocardiogram (ECG) revealed sinus rhythm and a pattern of significant left ventricular hypertrophy. The vectorcardiogram, however, was consistent with biventricular hypertrophy with predominance of the left-sided chamber.

The Doppler echocardiography showed AV and ventriculoarterial discordance combined with Ebstein’s anomaly of the tricuspid valve, ASD and coarctation of the aorta, with a pressure gradient of 40 mmHg between the ascending and the descending aorta.

The angiocardiographic and hemodynamic evaluations confirmed the clinical diagnosis.

At that time, due to the lack of clinically significant consequences of the regurgitation of the left AV valve, only the correction of the coarctation of the aorta with terminal anastomosis was performed.

During the postoperative period, the infant showed systemic hypertension. This condition was controlled with captopril, which was discontinued 2 months later. Currently, the infant is in good health and, on his physical examination, a slight regurgitation of the left AV valve is noted. The infant remains on low-dose medications for treating congestive heart failure.

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Case 2 - A 4-year-old male patient with a past medical history of congestive heart failure that started in the 1st week of life. At that time, he was hospitalized for 12 days and was discharged with a prescription for digitalis. He remained asymptomatic until his current age, when a heart murmur was detected, and he was then referred for diagnostic tests. He was in good health, his breathing was normal, and he did not have cyanosis. His pulse was strong in the upper limbs but absent in the lower limbs. His blood pressure was 150/90 mmHg in the upper limbs. His lungs showed no rales. The auscultation of the precordium revealed a 1st heart sound more audible in the mitral area, a split 2nd heart sound, with increased intensity in the pulmonary area, and a moderately intense systolic heart murmur with diastolic vibrations in the mitral area.

The chest X-ray revealed an enlarged cardiac silhouette and an increased pulmonary vasculature.

The ECG performed when the patient was 10 days old showed a QRS complex of the R type and a positive T wave in lead V1 (fig. 1). At the age of 4 years, his ECG showed a pattern of left ventricular hypertrophy with QS morphology from lead V1 to lead V3 and R morphology in leads V5 and V6 (fig. 1).

Doppler echocardiography, combined with the hemodynamic study, revealed AV and ventriculoarterial discordance, coarctation of the aorta and tricuspid regurgitation with moderate repercussion. The pressure gradient between the ascending and the descending aorta was 55 mmHg. The patient underwent termino-terminal anastomosis to correct the aortic coarctation. Currently, he remains in good health.

Patient with atrioventricular and ventriculoarterial concordance - Case 3 - A 14-year-old male teenager with a heart murmur detected at the age of 2 months. At that time, he was diagnosed with coarctation of the aorta. The hemodynamic study revealed an increased end-diastolic pressure of the left ventricle (LV) and normal pressures in the right ventricle as follows: mean right atrial pressure (7), right ventricular pressure (26/6 mmHg), pulmonary artery pressure (28/14/20 mmHg), mean pulmonary capillary pressure (13 mmHg), left ventricular pressure (105/20 mmHg) and pressure of the ascending aorta (105/70/85 mmHg). The angiographic study showed a moderate constriction of the initial portion of the descending aorta and an enlarged right ventricular chamber (fig. 2). The patient underwent surgery at the age of 6 years for correction of the coarctation of the aorta; the technique of termino-terminal anastomosis was used. He remained asymptomatic until the age of 14, when he was admitted to the Emergency Department of the Clinic due to a syncopal episode.

On physical examination at admission, he was in good health, his respiration was normal, he was not pale and did not have cyanosis; his HR was 70 bpm, his systemic blood pressure was 120/70 mmHg. His lungs showed no rales. His precordium did not show any deformities and his heart auscultation revealed a regular heart rhythm, with a slight regurgitation murmur in the tricuspid area. His pulse was symmetric and of normal amplitude in the limbs.

The chest X-ray revealed a moderately enlarged cardiac silhouette and an anomalous pulmonary vasculature.

The ECG showed a pre-excitation pattern (Wolf-Parkinson-White) of the B type and the vectorcardiogram revealed a QRS shifted to the left, backwards and upwards, with counterclockwise rotation at the horizontal and frontal planes, as well as a clearly delayed conduction time in the initial and end portions. An electrophysiological study was performed in this patient and revealed 2 anomalous bundles (one anterosuperior and the other posterolateral).

The echocardiography revealed Ebstein's anomaly with moderate tricuspid regurgitation and effective aortic isthmoplasia, without pressure gradient at the site of the surgical correction of the coarctation. The hemodynamic and angiographic studies confirmed the diagnosis of Ebstein's anomaly.

![Fig. 1 - A) Case 2 - electrocardiogram showing a QR pattern in lead V1 due to atrioventricular and ventriculoarterial discordance and hypertrophy of the ventricle placed on the right (left ventricle) on the 10th day of life; B) electrocardiogram of the same patient when he was 4 years old, showing hypertrophy of the ventricle placed on the left as a result of the coarctation of the aorta combined with the malformation of the left atrioventricular valve (right ventricle), in addition to anterosuperior divisional heart block.](image)

![Fig. 2 - Angiography of the patient with atrioventricular and ventriculoarterial concordance showing coarctation of the aorta (arrow) below the subclavian artery (A) and signs of right ventricular enlargement through the trajectory of the right coronary artery (B).](image)
Ablation of the anomalous bundles and tricuspid valvuloplasty according to the De Vega technique were performed. The immediate electrocardiographic and vectorcardiographic studies showed morphology of right bundle-branch block (RBBB) with the features observed in Ebstein's anomaly (fig. 3). Two years later, the patient remains in good health and is not receiving any medication.

**Discussion**

Coarctation of the aorta is one of the most unusual abnormalities associated with Ebstein's anomaly. In patients with Ebstein's anomaly that show AV and ventriculoarterial discordance, the coarctation of the aorta is less unusual than in those with concordance; this may occur due to the lower anterograde flow in the ascending aorta during the intrauterine life.

The coarctation of the aorta aggravates the clinical findings, especially when it plays a direct role in the malformation of the tricuspid valve, as occurs with ventriculoarterial discordance. In fact, the 2 patients with discordance had heart failure during the neonatal period.

Arrhythmias are common in Ebstein's anomaly and occur in more than 50% of these patients. The most frequent are supraventricular paroxysmal tachyarrhythmias and atrial flutter or fibrillation, as a result of right atrial dysfunction or anomalous muscular bundles, as observed in the 3rd patient. In that case, the electrophysiologic study revealed 2 anomalous pathways that were responsible for the episodes of paroxysmal tachycardia. Of note in this case was the ECG with a morphology of RBBB after the ablation of the bundles; this is a common finding in Ebstein's anomaly.

The initial decision of correcting only the coarctation of the aorta in the 3 cases and of adopting a conservative approach in relation to the tricuspid valve regurgitation in the 2 patients in whom the transposition had been corrected took into account the grade of dysfunction. This dysfunction was mild compared with the moderate hemodynamic repercussion, and was expected to decrease with the elimination of the aortic obstruction; this actually occurred. Only the 3rd patient required surgical therapy for the AV valve because this valve would not be effectively improved with the surgical therapy of the coarctation alone.

In conclusion, Ebstein's anomaly and coarctation of the aorta are rarely associated, mainly when a patient presents with AV and ventriculoarterial concordance.

**References**