Biventricular Endomyocardiofibrosis Associated with Renal Amyloidosis

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Endomyocardiofibrosis is a restrictive cardiomyopathy characterized by fibrotic involvement of the endocardium and adjacent myocardium, and by diastolic dysfunction caused by changes in distensibility making ventricular filling inadequate while preserving the systolic function. Clinically, it appears as heart failure, but etiological symptomatic discernment, suspicion and a clinical examination would be necessary in order to make a correct etiological diagnosis. The case of a patient with biventricular endomyocardial fibrosis associated with renal amyloidosis is presented.

CASE REPORT

A forty-nine year old black woman, native of Ilhéus-Bahia (Brazil), sought medical help at Emergency with a history of fatigue, anasarca, dyspnea with slight exertion and night paroxistics. These symptoms had appeared about a year previously and there had been a remarked worsening in the last six months. At first, the patient had a discrete edema in her lower members in the evening. This had gradually developed into anasarca two months earlier. There had been a considerable deterioration in her general clinical condition (lack of appetite, weight loss, greater tiredness) and dyspnea now even while resting. She had a history of chronic smoking since adolescence and hypertension controlled by captopril. During the physical examination she was pale, anxious, and had tachypnea (her breathing was more than 30 per minute). An examination of the lungs showed bilateral crepitant rales. She had a raised heart beat, of approximately 120 bpm and low blood pressure: 60 x 40 mmHg. Her cardiac rhythm was regular, showing a presence of a mitral and tricuspid systolic murmur and B4. An examination of her throat showed evident arterialization of the jugular venous pulse. An abdominal examination showed a painful hepatomegaly and moderate ascitis. Having made a syndromic diagnosis of heart failure, clinical treatment with intensive hemodynamic support and complementary tests were undertaken. The relevant findings were: creatinine, 2.37 mg/dl; urea, 98 mg/dl; albumin, 1.66 g/dl; total proteins, 4.2 g/al. Urina analysis with proteinuria and hemoglobinuria. The chest X-ray showed cardiomegaly, signs of pulmonary venocapilar hypertension, and the presence of protruding hilums, perihelium infiltrations, Kerley’s septal lines, laminar and costo-diaphragmatic effusions. The electrocardiogram showed an alternation between the sinusal and junctional rhythms, biatrial overloading and ventricular repolarization alterations in the side wall, with QRS complexes of low amplitude. A Doppler echocardiogram was taken, which showed evidence of a reduction in the dimensions of the left ventricular cavity, a slight enlargement in the right ventricle and in the myocardial thickness and a discrete pericardial effusion on the right (figures 1 and 2). The transvalvar flows showed systolic regurgitation in both atrioventricular valves (moderate/considerable mitral and tricuspid regurgitation). Systolic ventricular functioning was normal, with clear diastolic restriction when biventricular filling took place. Based on the initial tests, a diagnosis of...
restrictive cardiac insufficiency was reached, with the possibility of establishing three etiologies: cardiac amyloidosis, endomyocardial fibrosis or endomyocardial fibroelastosis. A myocardial scintigram with MIBI-Tc99 was performed evidencing normal perfusion and biventricular hypertrophy.

In order to make the definitive diagnosis, the patient underwent a myocardial biopsy and the histological examination proved endomyocardial fibrosis.

When using a polarized light technique, the Congo red stain was negative for amyloidosis, and Orcein for elastic fiber. With Masson’s Trichroma stain, the fibrotic conjunctive tissue showed positiveness (figures 3 and 4). Because of renal function loss a biopsy was carried out and associated amyloidosis was diagnosed; 24-hour proteinuria (4.9g) was investigated, as was the absence of erithrocitarian dismorphism. Hepatitis B, C and HIV, serical complements C3 and C4 were investigated and all negative. The patient has heart failure, functioning class III according to NYHA, and is waiting for cardiac surgical procedure.

**DISCUSSION**

Endomyocardial fibrosis is characterized by the deposit of fibrotic tissue covering the endocardium initially at the apex, and then, proceeding towards the atrioventricular valvar plane, leading to an increase in end diastolic pressure of the affected ventricle. It consequently leads to enlargement of the chamber and an increase in either systemic or pulmonary venal pressure. The final result is a hardening of the endocavity causing consequently ventricular dysfunction because of restriction of its filling.

In the case of the preferential affecting of the right ventricle the following was observed: an obstruction of the pointed end (associated or not with a thrombus), dilation in the outflow tract and paradoxical movement of the interventricular septum; The right chambers may be enlarged and the systemic veins dilated; Tricuspid regurgitation in varying degrees and an increase in the end diastolic pressure of the right ventricle usually without the association of pulmonary hypertension.

If the left ventricle is predominantly affected the alterations are repeated, in relation to the left ventricular cavities. The increase in pressures is transmitted to the left atrium and, consequently, to the pulmonary venal bed, thereby causing pulmonary hypertension. This increase in pulmonary pressure could, in turn, be transmitted over the tricuspid valve causing regurgitation, even without affecting the right cavities. Mitral or tricuspid valve regurgitation may or may not be expressed through cardiac murmurs, especially when the end diastolic pressure of the ventricular cavity is very similar to the average pressure of the chamber.

In relation to the electrocardiogram, the affecting of the right ventricle shows a decrease in the extent of the QRS complexes, due to the almost-constant pericardial effusion. Other signs found are the changes in the ST segment and T wave and right chamber overload criteria with the classic qR pattern in V1. In the affected left ventricle, it is already possible to find changes similar to those of the right ventricle, including atrial fibrillation and overloading of the left chambers. The EKG associated with cardiac amyloidosis disease may present low QRS voltage and evidence of sinusal nodule disease and/or a blockage of the atrioventricular tract with abnormal Q waves suggesting chronic myocardial infarct.

When the thorax was X-rayed, an enlargement of the heart area in different degrees was found, associated with signs of pulmonary venocapilar congestion.

The bidimensional echocardiogram greatly facilitated the diagnosis of endomyocardial fibrosis, because the point of the ventricles could be examined. The presence of fibrous

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**Fig. 1** - Longitudinal paraesternal transthoracic echocardiogram showing acoustic texture with thick granulations and apical obliteration due to a deposit of fibrotic tissue. AO: aorta; AE: left atrium; VE: left ventricle; VM: mitral valve
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Fig. 2 - A modified apical echocardiogram of the right cavities during ventricular diastole and ventricular systole showing apical obliteration of the right ventricle by fibrotic tissue. DERR - pericardial effusion; AD - right atrium; VO - right ventricle; VT - tricuspid valve.

Fig. 3 - Panoramic view of the biopsy demonstrating the cardiac muscle with extensive areas of fibrosis intermingling with myocardial fibers (arrows). HE stain increased 40 times.

Fig. 4 - Thickened endocardium by fibrosis, presenting tonality clearer (black arrows) in relation to normal fibers (blue arrows). HE stain 100 times. Endocardio = Endocardium.
tissue showed obliteration. Fibrosis is more ecogenic than the myocardial walls and this makes different from other forms of restrictive cardiomyopathies1-2.

The Doppler shows an aspect which is quite characteristic of restrictive myocardialopathologies, especially of endomyocardial fibrosis. The flow through the mitral or tricuspid valves, depending on which ventricle is damaged, is presented as practically forming only one velocity peak in the protodiastole (E wave) with an acute reduction in deceleration time, because the A wave is absent or greatly reduced. This aspect indicates a pattern of rapid flow immediately after the valvar opening, then followed by an abrupt disruption due to the existence of cavity obliteration and, therefore, a smaller cavity volume, or a restriction to the entry of blood due to the reduction in distensibility.

Biopsy endomyocardial helps in the differential diagnosis with other restricted cardiopathies that, with the clinical and complementary examination make the diagnosis.

The mitral flow is of great prognostic value. So, patients who present a mitral flow with restrictive characteristics (decelerating time less than 150 ms) have only a 49% chance of survival for one year against 92% in those whose flow indicates only a relaxation abnormality. The reduction in relaxation can be seen by means of the tricuspid and/or mitral valves1,2.

During the cardiac catheterization, the angiogram shows areas of obliteration in the ventricular apexes. The size of the cavities can be reduced and the myocardial wall thicker than normal. The pressure curves may be consistent with other restricted cardiopathies that, with the clinical and complementary examination make the diagnosis.

The complimentary test to finalize the diagnosis is, inexorably, the endomyocardial biopsy, which makes it possible to differentiate between the various modalities of restrictive myocardiopathy1,3,5.

In the studied case, the myocardial and renal biopsies both confirmed the diagnosis of endomyocardial fibrosis and renal amyloidosis, respectively.

Patients suffering from restrictive myocardialopathies, in particular those with endomyocardial fibrosis could first be labeled as carriers of congestive cardiac insufficiency without establishing a more definitive etiological diagnosis. Clinical treatment is frustrating and unsatisfactory1,6. Surgical treatment is the principal therapeutic option7. Diuretics are necessary to reduce both pre and after loading. However, it is unsafe to increase the diuresis to the point of reducing the pressures of cardiac filling8. Therapy with corticosteroids did not prove beneficial.

The prognosis varies from two to ten years depending on the gravity of the clinical condition, after the first signs8. The alternations which negatively influence the prognosis are: the intensity of the apical fibrosis, signs of diastolic restriction, damaging of the systolic ventricular functioning, involvement of the subvalvar system, presence of pulmonary hypertension and intercavity thrombi3,5. It is, however, the duty of both the physician and the cardiologist to undertake a careful examination, complemented by subsidiary exams such as an electrocardiogram, thorax X-ray, a Doppler echocardiogram, angiogram, myocardial ventriculography and biopsy, in order to make a correct diagnosis so as to be able to indicate the necessary treatment as quickly as possible. Surgical treatment has been prematurely indicated immediately after diagnosis1,6. This has led to endocardial resection, but however, with a high morbidity-mortality rate, even with experienced surgeons10.

In some patients the tricuspid and mitral valves are substituted with prothetic6,7 valves and a heart transplant could be necessary in certain cases8.

**REFERENCES**