Genetic disorder or toxoplasma myocarditis: a case report of dilated cardiomyopathy with hypertrabeculation in a young asymptomatic woman

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Abstract

Isolated noncompaction of the left ventricle (LV) is a rare disorder, classified as a primary genetic cardiomyopathy by the American Heart Association. The European Society of Cardiology Working Group on Myocardial and Pericardial Diseases classified LV noncompaction as an unclassified cardiomyopathy. LV noncompaction cardiomyopathy characterized by the following features: 1) an altered myocardial wall with prominent trabeculae and deep intertrabecular recesses resulting in thickened myocardium with two layers, consisting of compacted and noncompacted myocardium and 2) continuity between the left ventricular cavity and the deep intertrabecular recesses, which are filled with blood from the ventricular cavity, without evidence of communication with the epicardial coronary artery system. Features of LV noncompaction can overlap with dilated cardiomyopathy, hypertrophic cardiomyopathy (especially the apical variant), and restrictive cardiomyopathy. The phenotypic expression can vary considerably within the same family. The LV noncompaction can rarely occur as a transient phenomenon during myocarditis.

We present the case of a 23-year-old patient, admitted to our Department for cardiac evaluation because of ECG changes and cardiac enlargement revealed at thoracic radiography. She had a history of chronic toxoplasmosis. An echocardiography was performed revealing left ventricular enlargement with severe systolic and diastolic dysfunction, diffuse hypokinesia and signs of isolated left ventricular non-compaction. Under these circumstances, we have considered the presence of isolated left ventricular non-compaction. A cardiac Magnetic Resonance Imaging was performed and it sustained the diagnosis. The alternative cause of isolated left ventricular noncompaction (prominent trabeculation due to myocardial toxoplasmosis) was considered improbable.

Key words: cardiomyopathy; noncompaction; trabeculae; toxoplasmosis

Introduction

The left ventricular (LV) isolated non-compaction is characterized by embryonic myocardium morphology due to an arrest in the compaction process during the first trimester, and it is a very rare cardiomyopathy. Clinical manifestations are non-specific, ranging from asymptomatic patients to acute congestive heart failure. The diagnosis is based on cardiac imaging using echocardiography and magnetic resonance imaging.

Case report

A healthy 23-year-old woman presented to hospital for cardiac workup after she had been referred to a cardiologist by a general practitioner, after detecting changes on electrocardiogram (ECG), consisting of negative T waves and cardiac enlargement on thoracic radiography. She was asymptomatic at presentation and without any familial cardiac history. She had a history of chronic toxoplasmosis (with fever and lymphadenopathy, 12 months prior to this hospitalization) for which she was taking antibiotic treatment. The clinical examination showed pallor of skin and mucous membrane, maculo-papular non-itching rash on the forearms and thighs. There were also mobile axillary enlarged (2 cm) lymph nodes not tender or painful with moderate hard consistency, hepatosplenomegaly, blood pressure (BP) of 110/60 mmHg, heart rate (HR) of 70 b/min, regularly and apexian systolic murmur grade III/VI radiating to the axilla. Laboratory tests showed a discrete inflammatory syndrome (elevated erythrocyte sedimentation rate and C-reactive protein level), IgG antibodies to Toxoplasma in serum (1497,3 UI/mL).
Initial ECG (see Figure 1) showed poor r wave progression in V1-V3 leads and negative T waves in DI, DII, aVF, and V4-V6 leads and thoracic radiography (see Figure 2) showed an increased cardio-thoracic index.

An echocardiogram (see Figure 3 A, B and C) was performed revealing left ventricular enlargement with severe systolic and diastolic dysfunction (LV ejection fraction 25%, filling pattern type III- restrictive pattern), diffuse hypokinesia and trabeculation suggesting isolated left ventricular non-compaction.

A cardiac Magnetic Resonance Imaging (MRI) (see Figure 4 A and B) was performed and it confirmed dilated cardiomyopathy with excessively prominent trabeculations in left lateral wall and ventricular apex, severe alteration of left ventricular function with diffuse hypokinesia.
Discussion

One of the most important questions in this case is the etiology of dilated cardiomyopathy: a non-compacted or a secondary dilated cardiomyopathy. The left ventricular isolated non-compaction is a genetic cardiomyopathy and can occur isolated or in association with congenital heart defects, genetic syndromes and neuromuscular disorders. It is characterized by prominent trabeculations and deep recesses within the ventricular myocardium resulting in apparently hypertrophied myocardium. The myocardium is actually thickened and has two layers consisting of compacted and non-compacted myocardium [1,2,3]. Normal myocardium can present trabeculae but always less than three, while in non-compaction, the trabeculae are increased in number and very prominent, the compact layer is thinner and between the trabeculae there are deep recesses in communications with the ventricular cavity, but not with the epicardial coronary system [3,4,5,6].

Most patients have systolic left ventricular dysfunction and they present with congestive heart failure. The diagnosis is given by the echocardiography. Current echocardiographic criteria include: a) presence of multiple trabeculae – especially in the apex and the left ventricular free wall; b) multiple deep recesses, between trabeculae, communicating with ventricular cavity (demonstrated by color Doppler imaging); c) a two-layer structure of the myocardium with non-compact increased layer (ratio between non-compact and compact layer suggested as >2.0) [5,6].

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