CONGENITAL MITRAL VALVE DYSPLASIA IN A FEMALE CHILD: INFECTIVE ENDOCARDITIS OR HERITABLE DISORDER OF CONNECTIVE TISSUE?

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KONGENITALNA DISPLAZIJA MITRALNE VALVULE U ŽENSKOG DETETA: INFEKTIVNI ENDOKARDITIS ILI NASLEDNI POREMEĆAJ VEZIVNOG TKIVA?

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INTRODUCTION

The multitude of names for mitral valve disorders - the floppy mitral valve (FMV), mitral valve prolapse (MVP), or mitral valvular regurgitation (MVR) - and the variability of diagnostic criteria related to children have emerged (1-6). The fusion of FMV/MVP/MVR dates back to the 1960s (2). Prevalence of FMV/MVP is about 1-5% and it increases with age (1, 2). Three autosomal dominant and one X-linked genetic loci for FMV/MVP have been found over the last few years (2, 7-10).

The basis for diagnosis is (1, 2, 6): a history or detailed family history of the development of connective tissue or MV disorders; general inspection, anthropometrics (a patient may have asthenia, hypomastia or skeletal abnormalities: pectus excavatum/carinatum, scoliosis, straight back, marfanoid appearance); symptoms of FMV/MVP/MVR dates back to the 1960s (2). Prevalence of FMV/MVP is about 1-5% and it increases with age (1, 2). Three autosomal dominant and one X-linked genetic loci for FMV/MVP have been found over the last few years (2, 7-10).

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(palpitations, fatigue, exercise intolerance, dyspnea, chest pain, postural phenomena, syncope/presyncope, neuroendocrine or autonomic dysfunction); auscultation (the mid-systolic click accompanying a systolic murmur, mild or progressive MV regurgitation); electrocardiogram (the vast majority of patients have a normal resting electrocardiogram; ST and T-wave changes may improve with exercise); chest radiograph; echocardiography (it is a sensitive tool in the differentiation of degenerative MV disease); cardiac magnetic resonance.

Congenital MV dysplasia can become symptomatic at any age and may sometimes require surgical treatment (3, 4, 11). Clinical expression (phenotype) in FMV/MVP may vary among family members with heritable connective tissue disorders (Marfan syndrome, Stickler syndrome, Ehlers-Danlos syndrome types I, II or IV, Osteogenesis imperfecta syndromes, Fragile X syndrome- Martin-Bell syndrome, Polycystic kidney disease in adults, Pseudoxanthoma elasticum etc.) posing additional diagnostic challenges (10-14).

Possible complications are MR progression, atrial or ventricular arrhythmias, congestive heart failure, infective endocarditis, embolic phenomena or sudden death (1, 2).

**THE CASE**

Three years and seven months old female child weighing 14.4 kg, body height 98.9 cm was referred to our clinic with clinical signs of upper respiratory tract infections (sore throat and lower legs, nasal secretion, fever up to 39.9 °C). During the previous month the child had suffered from frequent respiratory infections. Complete biochemical analyses were within normal ranges except for slightly elevated erythrocyte sedimentation (SE 35 mm/h, repeated was 5 mm/h), and leukocytosis 12.2 x 10⁹/L (neutrophils 85%), repeated was 7.4 x 10⁹/L.

She has systolic murmur, the intensity 2-3/6 left parasternal and at the apex. Electrocardiogram and chest radiograph were normal. Echocardiography was performed (29/09/2011, Figures 1 and 2). Because of the suspected infective endocarditis, 5 blood cultures were taken. In the second one, Staphylococcus haemolyticus was isolated.

Echocardiographic examination (13/10/2011) showed that both mitral leaflets significantly protruded into the left atrium during systole, with MR grade +1.5/4. The anterior

![Figures 1 and 2. Two dimensional echocardiography-left parasternal four-chamber view, showing mitral valve leaflets with spherical apical thickening diameter around 8-10 mm, inhomogeneous structure, (central hypoechoogenic).](image1)

![Figures 3 and 4. Two dimensional echocardiography-long axis view, showing altered morphology of both leaflets, especially anterior mitral valve leaflet with systolic regurgitate jet semi-quantified as +1 - 1.5 / 4](image2)
mitral leaflet was dysplastic with registered suspected verrucous formations of 8 x 4-5 mm diameter. The left ventricle was circular in cross-section and of good global contractility: end-diastolic diameter (EDD) was 32 mm, end-systolic diameter (ESD) was 20 mm and ejection fraction (EF) was 70%, fractional shortening (FS) was 38%. Left atrial (LA) was 17 mm; aorta (Ao) was 13 mm. Repeated complete biochemical analysis (parameters of inflammation) and five blood cultures (sterile) were within normal limits.

Control echocardiography was performed (7/12/2011): altered morphology of both MV leaflets, especially the anterior one. In the left ventricle (LV) there were two papillary muscles, degenerative leaflet by normally developed chordae tendineae. Posterior MV was normal appearance and movement. Anterior cusp was very elongated with spherical apical thickening of about 9 mm diameter (inhomogeneous structure with central hypoechogenic zone) which was not supportive of bacterial verrucae; there was also no typical flotation. Doppler on the MV recorded systolic regurgitate jet semi quantified as +1-1.5 / 4 directed to the left and rear. (EDD 34 mm, EDS 22 mm, FS 35%). Other findings were normal. Echocardiography, which was made on 14/03/2012 (6 months after the initial ultrasound examination of the heart), was identical to the first one (Figures 3 and 4).

DISCUSSION

The aim of this paper is to review our experience with a rare form of MV dysplasia. Three years and seven months old female child was strongly suspected of having infective endocarditis (15) according to differential diagnostic procedure based on clinical signs (sore throat and sore lower legs in the evening, long-lasting fever with pauses in the period of one and a half months, one positive blood culture, first time detected heart murmur projected by the mitral valve apparatus and echocardiographic finding - suspected verrucae on mitral valves). According to the Duke criteria, the patient met 1 major criterion (positive ultrasound finding on the heart - existence of verrucae on mitral valve apparatus) and 3 minor criteria: high temperature - fever, one positive blood culture and arthralgia. Based on phenotype characteristics of the little girl and her father, repeated echocardiographic finding (apical thickening of front mitral cusp is of inhomogeneous structure and without characteristic flotation) and good general condition of the child during hospitalization, the case was most probably the congenital mitral valve dysplasia within broader clinical condition belonging to the group of heritable connective tissue diseases (1, 2, 13-15).

Father had a chest deformity, skin hyper-elasticity of smaller extent (most prominent on abdomen) and thinner skin, especially on the feet. Based on the medical history, clinical signs, phenotype characteristics and performed investigations, the Ehlers-Danlos syndrome was suspected. According to differential diagnostic procedure some other connective tissue diseases might be considered as well (1, 2, 13-15), such as Cutis laxa, Marfan syndrome and homocystinuria. Molecular genetic analyses of isolated DNA or skin fibroblast (1, 2) are performed for some forms, which was not done here due to technical and financial reasons (such analyses are not performed in Serbia).

Three years and seven months old girl with dysplasia of both mitral leaflets (particularly anterior one) with MR graduate 1, 5/4 was strongly suspected of having infective endocarditis according to differential diagnostic procedure. According to the Duke criteria, the patient met 1 major criterion (positive ultrasound finding on the heart - existence of verrucae on mitral valve apparatus) and 3 minor criteria: high temperature - fever, one positive blood culture and arthralgia. Based on phenotype characteristics of the little girl and her father, repeated echocardiographic finding (apical thickening of front mitral cusp is of inhomogeneous structure and without characteristic flotation) and good general condition of the child during hospitalization, the case was most probably the congenital mitral valve dysplasia within broader clinical condition belonging to the group of heritable connective tissue diseases. This was supported by the fact that the results of repeated echocardiographic examination remained unchanged 6 months after the first examination.

REFERENCES


