ST-Elevation Myocardial Infarction in Duchenne Muscular Dystrophy

LUCIANA TEODORA ROTARU1, MARIA FORTOFOIU1, ADRIAN MITA2, ANCA BARAU ABU ALHIIJAI, OANA CRICIOTOIU3, VERONICA CALBOREAN4, CRISTINA GEORMANEANU5, LIVIU MARTIN6, MIHAI BANICIOIU3, LAVINIA MARIA GHEORMAN4, MIRCEA-CATALIN FORTOFOIU2

1University of Medicine and Pharmacy of Craiova, Department of Emergency Medicine and First Aid, 2-4 Petru Rares Str., 200349, Craiova, Romania
2University of Medicine and Pharmacy of Craiova, Department of Internal Medicine, 2-4 Petru Rares Str., 200349, Craiova, Romania.
3University of Medicine and Pharmacy of Craiova, Neurology Department, 2 Petru Rares Str., 200349, Craiova, Romania.
4University of Medicine and Pharmacy of Craiova, Cardiology Department, 2 Petru Rares Str., 200349, Craiova, Romania.
5Military Emergency Hospital Dr. Stefan Odobleja, Surgery Department, 150 Caracal Str., 200749, Craiova, Romania.
6Military Emergency Hospital Dr. Stefan Odobleja, Department of Diabetes & Metabolism, 150 Caracal Str., 200749, Craiova, Romania.

The paper presents the cases of a twelve years old boy brought in emergency department with the following symptoms: pain in the thorax and upper abdomen, nausea, vomiting with sudden onset 11 h ago.

Keywords: Duchenne muscular dystrophy, ST-elevation myocardial infarction

In Duchenne muscular dystrophy (DMD), the electrocardiogram (ECG) is abnormal at very early stages, before echocardiographic findings of cardiomyopathy or evidence of myocardial fibrosis (MF) on cardiac magnetic resonance (CMR). The appearance of ST elevation myocardial infarction (STEMI) are an exceptional event in DMD even in DMD-associated cardiomyopathy.

Experimental part

A 12-year old boy with confirmed DMD; without positive family history for this disease or personal history of cardiac diseases; under treatment with β-blockers and corticosteroids; is brought in emergency department (ED) for the following symptoms: pain in the thorax and upper abdomen, nausea, vomiting with sudden onset 11 h ago.

The electrocardiogram (ECG) and aspect of genetic analysis are illustrated in the figures below (fig. 1, 2).

Troponin I (cTnI) initial level was 3000 ng/mL and increasing up to 20000 ng/mL; N-terminal pro-B-type natriuretic peptide (NT-pro BNP) was 1799 pg/mL.

Echocardiography showed hypokinesia of posteroinferior wall and basal septum and left ventricular ejection fraction (LVEF) < 35%. Coronarography was normal. CMR with gadolinium has revealed MF in inferolateral wall.

The therapeutic decision was initially resorted to subcutaneous heparin administration, subsequently suppressed; adding enzyme conversion inhibitors (perindopril); doubling the corticosteroid dose to 60 mg/day and emergency air evacuation to hospital with the pediatric cardiology clinic. Administration of β-blockers was suppressed due downward trend of the LVEF.

Results and discussions

This case has some peculiarities: child with confirmed DMD cardiomyopathy under treatment, without personal history of ventricular dysfunction (VD) or arrhythmias with elements for STEMI sustained by those described above.

Starting from these prerequisites, several issues are raised related to the interpretation of clinical and biological signs, explaining the determinism of the phenomenon, differentiating aspects related to myocardial injury, the usefulness of current guidelines[1-6].

In this case, the most probable hypothesis can be accelerated apoptosis associated with maldistribution and the consecutive decrease of LVEF well supported by the stage of DMD, patient age and CMR aspects. Very rarely an acute myocardial infarction (MI) is a DMD-related cardiac event but is likely that is not caused by coronary artery disease and apoptosis is usually painless [7-10]. Previous arrhythmias was not documented and effects of long-term corticosteroid therapy or presence of vasculitis have not been confirmed by coronarography.

* email: catalin1972@hotmail.com; Phone +40722320438

Fig. 1. The ECG revealed the acute inferior myocardial infarction (ST-segment elevation in leads III and aVF)

Fig. 2. Genetic analysis features.
Chest pain is the most common clinical manifestation in DMD and is usually considered to be caused by skeletal muscle damage but many of the symptoms that belong to DMD are present in heart failure [11-17].

cTnI and BNP are serum biomarkers for ischemic coronary artery disease and there are no routine tests in patients with DMD [2].

The most common ECG findings in patients with DMD are left or right ventricular hypertrophy and sinus tachycardia. Q waves in the left precordial leads, tall R waves and increased R/S amplitude in lead V1, right axis deviation are another ECG findings. Atrial fibrillation and atrial flutter, can also occur. When LVEF<35%, may occur ventricular tachycardia, premature ventricular complexes and conduction abnormalities[11-14].

Sudden cardiac death occur occasionally when LVEF>35% and VD is predictive for rhythm arrhythmias [18-21].

In clinical guidelines, evaluations for cardiomyopathy are well established but for symptomatic patients with demonstrated cardiac disease assessment will be made whenever the cardiologist considers necessary [1, 22-25].

Current guidelines do not provide any recommendation on thrombolytic therapy in children and doesn’t stipulate the placement of implantable cardioverter-defibrillators in DMD-associated cardiomyopathy [1-2].

Conclusions

Acute chest pain in DMD patient requires, a detailed cardiac history, ECG, cTnI and reference to the cardiologist. The clinical features of VD and evidence of MF on CMR is usually considered to be caused by skeletal muscle damage but many of the symptoms that belong to DMD and is usually considered to be caused by skeletal muscle damage but many of the symptoms that belong to DMD are present in heart failure [11-17].

Acute chest pain in DMD patient requires, a detailed cardiac history, ECG, cTnI and reference to the cardiologist. The clinical features of VD and evidence of MF on CMR

References


Manuscript received: 19.07.2018