Acute Myocarditis Mimicking an Acute Coronary Syndrome: Case Report and Mini-Review of the Literature

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INTRODUCTION

Acute myocarditis is an inflammation of the heart muscle, confirmed histologically by a rate of inflammatory cells and signs of myocardial necrosis of non-ischemic origin [1]. Inaugural signs of heart failure, in the absence of coronary disease or progressive valvular disease stigma, should lead to the diagnosis of acute myocarditis, even more in the presence of flu symptoms during the previous days [2]. The diagnosis could be challenging, based on clinical presentation, confirmed mainly by cardiac magnetic resonance imaging (MRI) rarely by myocardial biopsy. We report the observation of a 19-year-old male admitted to the cardiology department of IBN ROCHD University Hospital of Casablanca, for acute myocarditis misdiagnosed initially as an acute myocardial infarction ST elevation and confirmed by cardiac magnetic resonance imaging.

CASE PRESENTATION

It is about a 19-year-old male, without any cardiovascular risk factor but with a history of a flu-like signs, who was admitted to the intensive unit care for acute onset of constrictive retrosternal chest pain, persisting for the last 10 hours before admission.

The initial clinical examination revealed a blood pressure of 115/65 mmHg, a Heart rate of 72 bpm and body temperature of 38.3 °C. The cardiac and pulmonary examination was normal—without signs of acute heart failure.

The electrocardiogram (ECG) showed an ST-segment elevation in leads in the inferior territory (Fig 1). A chest X-ray was ordered with no abnormalities. Initial blood analysis showed a significantly elevated level of Troponin Ic up to 10 ng/ml. The echocardiography revealed left ventricular systolic dysfunction with hypokinesia localized in the inferoseptal wall, with an ejection fraction mildly reduced of 50%, without any valvular abnormalities or pulmonary hypertension.

The patient received dual antiplatelet therapy, low molecular weight heparin and was admitted directly to the cath lab where an angiography showed the absence of coronary artery disease or spasm. Biology tests showed inflammatory syndrome with a sedimentation rate of 100 mm and C-Reactive Protein up to 130 mg/l with high Creatine PhosphoKinase levels of 512 UI/l. Viral serologies, however, were negatives.
Within 24h chest pain resolved, body temperature remained between 35.6 and 37.5; ECG showed a mild regression of the ST-segment elevation with T wave inversion on the same territory. Cardiac MRI (Fig 2) was performed showing T2 signal enhancement and late gadolinium enhancement in the subepicardial inferior wall which confirmed the diagnosis of acute myocarditis. The patient was discharged one week later with anti-inflammatory treatment, ramipril 5 mg and bisoprolol 2.5 mg with continuous pain sedation. A post-discharge one month follow-up visit showed that the patient was recovering well, with normalization of ECG, and echography parameters.

**DISCUSSION**

The diagnosis of myocarditis is challenging, due to the variability of symptoms that are often unspecific, ranging from asymptomatic to life-threatening. Clinical suspicion is based mainly on a medical history of flu-like signs as mild fever, myalgia, and nausea, one to three weeks before the other symptoms, and regressing spontaneously afterwards [3]. Few patients can develop retrosternal chest pain that can mimic acute coronary syndrome, shortness of breath, arrhythmia or even inaugural acute heart failure, which may require hospitalization. The difficulty of diagnosis in Myocarditis lies in the lack of established non-invasive gold standard confirmation test. As the endomyocardial biopsy (EMB) is the current invasive gold standard [4], it has a relatively low sensitivity due to the variability in interpretation and sampling error [5]. ECG often displays some abnormalities, T wave changes, ST elevation, atrial and ventricular arrhythmias, atrioventricular conduction defects, or blocks [6]. Nonetheless, these findings remains non-specific and may be present in various conditions. Cardiac troponin is often elevated in the acute phase of Myocarditis [7]. However, as electrocardiogram, Troponins levels are non-specific for myocarditis. Echocardiography might be useful to myocarditis diagnosis; it can show left ventricular dilation, decreased systolic and / or diastolic function, and localized wall motion abnormalities, even though these findings could be found during myocardial infarction [8]. The indication of coronary angiography remains justified, when the SCA chart is complete, so as not mislead an occlusion of epicardial coronary artery. Due to a lack of specificity of clinical signs, many diagnoses should be considered as differential diagnoses, including Acute Coronary Syndromes (coronary spasm, myocardial infarction or ischemia; unstable angina), Pericarditis and Pericardial effusion should be considered as a differential diagnoses. Recently, MRI has become the best non-invasive modality of diagnoses objectiving the presence of infiltrates characterized by areas of enhancement (late gadolinium enhancement) on approximately 10 minutes of gadolinium administration with a patchy or

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**Figure 1.** (Patient’s ECG): ST Elevation in Inferior Leads (II, III, AvF), ST Depression and Negative T Wave in V1 V2 V3 and I, AvL.

**Figure 2.** (Cardiac MRI): Enhanced Signal Intensity in T2-weighted Sequences which Translate Subepicardial edema; Late Gadolinium Enhancement LGE: High Signal Intensity (Late Enhancement) in Epicardial Regions with Respect of Endocardium.
multifocal pattern correlating well with regions of histologically proven myocarditis
more importantly, cardiac MRI can help differentiate myocarditis from ischemic cardiomyopathy. As ischemic cardiomyopathy tend to display endocardium damage with edema and enhancement localized the culprit vessel territory whereas a case of myocarditis would present as a characteristic epicardial and myocardial involvement sparing endocardium. cMri can also provide a prognosis clue. As it appears that contrast enhancement ratio at four weeks was predictive of the functional and clinical long-term outcome in myocarditis patients. Treatment of myocarditis is primarily supportive but should bear in mind to treat any identifiable cause.

Patients who experience heart failure should receive angiotensin-converting enzyme inhibitors or angiotensin II receptor blockers, beta-blockers, and diuretics if needed. Dysrythmias may be seen with or without left atrial or ventricular thrombus that may require anticoagulation. Even though the causal agent is often viral, antiviral therapy is not recommended, and non-steroid anti-inflammatory agents should be avoided in the acute setting as they may impair healing of the myocardium. Regular consultations as a follow-up should be planned with repeated echocardiography.

ConCLUSION
The pseudo-infarction forms of acute myocarditis are less known. The differential diagnosis remains difficult despite advances in exploration methods noninvasive, in particular, MRI which constitutes the first diagnostic means currently. The interest of early diagnosis and treatment adapted to the etiology on the acute phase of the myocarditis enhances prognosis.

Conflict of Interests
The authors declare that they have no conflict of interests.

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