Arrhythmogenic Right Ventricular Cardiomyopathy/Dysplasia (ARVC/D), Case Presentation and Literature Review

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Case Report

Abstract

A case of Arrhythmogenic Right Ventricular Cardiomyopathy/Dysplasia (ARVC/D) has been presented. ARVC/D is a disorder of RV myocardium which involves replacement of the myocardium with fibrofatty tissue. The diagnosis was based on the findings of enlarged RV by echocardiography, ECG findings of: a) TWI (T-wave inversion) in V1, V2 & V3. b) Epsilon wave in V1, V2, and more than 500 VEs/24 hour by Holter monitoring. Positive family history of sudden death in first degree relative (mother) was found.

This case supports the first report of 34 Cases of ARVC/D reported in Iraq and attracts the attention of the physicians, echocardiographers and cardiologists to the diagnosis of this fatal disorder where SCD (sudden cardiac death) can be prevented by ICD (implantable cardiac defibrillator) implantation.

Introduction

Arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D) is predominantly a genetically determined heart muscle disorder that is characterized pathologically by fibrofatty replacement of the right ventricular (RV) myocardium [1]. It starts in the inflow and outflow tract of the RV then become progressively diffuse and may involve the posterolateral part of the left ventricle (LV) [2,5]. It generally presents in adolescents and young adults, mostly in men [3,4]. The most accepted prevalence rate is 6/10 000 [2,3,5]. Clinically patients commonly presents with symptomatologies of palpitation, dizzy spells or syncope due to ventricular arrhythmias mostly sustained ventricular tachycardia (sVT) of LBB pattern (LBB VT) or frequent LBB shape ventricular ectopic beats. Sudden cardiac death (SCD) due to ventricular fibrillation
(VF) can occur and may be drastically the presenting feature [5-9,14]. It may be discovered during Echocardiographic examination for assessing patients for different cardiac symptoms [5,10].

ARVC/D is a worldwide disease, but in middle east Arab countries it was first reported from Iraq in 2010 [11]. It is commonly missed disease and needs a high clinical suspicion to fulfill the diagnosis based on the International Task Force Criteria (TFC) [12].

We are reporting this case of ARVC/D in a 32 years female from Mosul region in the north of Iraq.

**Case Report**

KK is a 30 years old lady from Singar /Mosul presented to Dr Falah Al-Azawi in Baghdad , Iraqi centre for heart diseases, with two months history of recurrent dizzy spells ,pre syncope and palpitation. She is not diabetic or hypertensive and always enjoyed a good health . On physical examination: BP 135/80 , PR=75 bpm regular, chest is clear and the heart showed normal double rhythm heart sounds. ECG : partial RBBB.TWI V1,V2 &V3.Epsilon wave at the end of the QRS in V1&V2.Figure 1 & 2. CXR was normal. Echocardiography showed enlarged RV of 4.8 cm at the RVOT region with segmental dyskinetic RV wall and RVEF 35%. Figure 3. Family history revealed three relatives died suddenly; mother at age of 35 years and two cousins at age of 25 and 26 years. Holter: very frequent unifocal VE reaching to 500/day.

Diagnosis of ARVC/D is made according to the new task force criteria published in February 2010 in the European Heart Journal [12]. In this patient two major criteria were found . 1) Epsilon wave at V1,V2 . 2) Enlarged dyskinetic RV of more than 3.2cm. Additional criteria also seen: a) T wave inversion (TWI) in V1,V2 and V3 with no complete RBBB (major criteria).b) ventricular ectopics (VE) of more than 500/24 hours in Holter monitor (minor criteria). C) premature sudden death in mother (minor criteria).

According to the fatal family history of SCD in the mother , ICD implantation was advised.

**Discussion**

ARVC/D is genetically and pathologically a disease of the cardiomyocyte junction and plakophylline-2 is the most frequently targeted gene [10,13]. It presents clinically in many ways; SCD is one of the ways of presentation often in athletes [8,14]. The finding of sVT of LBBB pattern , non sustained VT or frequent VEs, incidental finding of RV dilatation by Echocardiography, and TWI in V1,V2 & V3 or beyond may attract the attention to look for other criteria of ARVC/D to reach a diagnosis based on the TFC [12]. Family survey of patients with the disease may discover other affected family members [7].

Patient may be diagnosed in stable period but may progress slowly or may have bursts of VT /VF and/or deterioration of RV function to show features of right sided heart failure [8,10,14,15].

ARVC/D is a world wide disease , it’s prevalence is more in certain localities like Padua in Italy [2,15].

In the Middle East Arab countries the only report of this disease was from Iraq, where 34 patients has been reported in 2010 ,all those patients presented with LBB pattern VT and fulfilled the TFC for the diagnosis of ARVC/D [11].

This presented case high lights the presence of this disease in Iraq but it needs careful consideration of the
diagnosis when patient presents with sustained or non sustained VT of LBB morphology, frequent LBB pattern VE, enlarged RV by Echocardiography, ECG findings of TWI in the precordial leads with or without partial or complete RBBB and Epsilon wave in V1, V2.

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References
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Figure 1 Twelve lead electrocardiogram showing partial RBBB, T wave inversion in V1, V2 and Epsilon wave in V1 (arrow head).

Figure 2 Magnified ECG strips of V1, V2 and V3 showing Epsilon wave (red arrow) at V1 & V2.
Figure 3. II D and M-Mode echocardiogram showing enlarged dilated RV (double head white arrow). RV=right ventricle. LV=left ventricle.