Case Report

An unusual ECG pattern in restrictive cardiomyopathy

M. Selvaganesh a,*, A.S. Arul c, S. Balasubramanian c, N. Ganesan b, S. Naina Mohammed b, G.S. Sivakumar b, S.R. Veeramani b, P. Jeyasinh a, S. Sathishkumar a, S. Selvaraju a

a Resident, Dept of Cardiology, Madurai Medical College, India
b Assistant Professor, Dept of Cardiology, Madurai Medical College, India
c Professor, Dept of Cardiology, Madurai Medical College, India

ARTICLE INFO

Article history:
Received 8 January 2015
Accepted 19 May 2015
Available online 8 July 2015

Keywords:
Restrictive cardiomyopathy
Electrocardiogram
Diffuse ST depression
aVR ST elevation

ABSTRACT

Restrictive cardiomyopathy is the least common type of primary cardiomyopathies. Electrocardiographic recording is abnormal in 99% of patients with RCM. Biastral enlargement, obliquely elevated ST segment with notched or biphasic late peaking T waves are considered characteristic ECG finding. Significant ST depression with T inversion mimicking subendocardial ischemia has also been reported in patients with RCM and is even suggested as a predictor of sudden cardiac death. We noted a similar ECG pattern in a 16 yr girl with Idiopathic restrictive cardiomyopathy. Coronaries were normal, stress perfusion imaging did not show any perfusion defect. This diffuse resting ST depression with T inversion in precordial & inferior leads along with ST elevation in aVR was persistent for more than six months.

Copyright © 2015, Cardiological Society of India. All rights reserved.

1. Introduction

Restrictive cardiomyopathy (RCM) is the least common type of primary cardiomyopathies. Electrocardiographic recording is abnormal in 99% of patients with RCM. ST segment, T wave changes are observed in nearly 75% of RCM patients. Significant ST depression mimicking ischemia has also been reported especially in idiopathic RCM. Classically in a patient with anginal pain, ST elevation in lead aVR with diffuse ST segment depression was considered as a sign of obstructive Left Main Coronary Artery disease. But similar ECG pattern can be seen in other conditions like left ventricular hypertrophy (LVH) as well. Here we report such an ECG pattern mimicking ischemia observed in young girl with RCM.

2. Case report

SA 16 yr old girl was admitted with history breathlessness class II–III since 3 months, history of atypical anginal type of chest
pain since 3 weeks. Her pulse, blood pressure were normal and she had a prominent a wave in JVP with a loud P2.

ECG was showing sinus rhythm with QRS axis of 100°, bifid P wave in lead II, 5–7 mm ST horizontal depression with inverted T waves in precordial leads, 3–5 mm ST depression in lead I, II, III aVF, 3 mm ST elevation in lead aVR (ST elevation in aVR with ST depression in >8 leads) (Fig. 1A).

Echocardiography revealed bialtral enlargement normal sized ventricles (Figs 3A, B & 4). Interventricular septal, LV free wall thickness was normal and LV mass was about 110 gm (Fig. 7A). She had no regional wall motion abnormalities and a normal systolic function with EF (68%). Doppler examination was suggestive of restrictive pattern (diastolic dysfunction Grade IV) Figs. 5A, B & 6A, B.

Simultaneous LV and RV pressure tracing on cardiac catheterization showed characteristic dip and plateau pattern (square root sign) with elevated LV and RV end diastolic pressures (Fig. 9) typical of restrictive hemodynamics.

She had normal blood cell count and morphology, serum ferritin and transferrin were within normal range. Liver function test, bone marrow examination and electrophoretic pattern of Immunoglobulins were also normal.

Coronary angiogram revealed normal epicardial coronaries (Figs. 8A, B). Myocardial perfusion study (MPI) using 99 mTc-Sestamibi did not show any evidence of resting or stress induced hypoperfusion or transient ischemic dilatation (TID) (Fig. 10). Cardiac MR revealed bialtral enlargement normal LV mass, pericardium (Fig. 7B).

### 3. Discussion

Restrictive cardiomyopathy has a nonspecific clinical manifestation. It is usually diagnosed by echocardiographic, catheterization findings of restrictive physiology along with normal or near normal sized ventricle and normal systolic LV function. A prominent a wave in jugular venous pulsation (JVP) with a loud P2 on the examination is typical. ECG may show sinus rhythm with QRS axis of 100°, bifid P wave in lead II, 5–7 mm ST horizontal depression with inverted T waves in precordial leads, 3–5 mm ST depression in lead I, II, III aVF, 3 mm ST elevation in lead aVR (ST elevation in aVR with ST depression in >8 leads). Echocardiography will reveal bialtral enlargement normal sized ventricles, normal systolic function with EF (68%), and no regional wall motion abnormalities. Doppler examination will be suggestive of restrictive pattern (diastolic dysfunction Grade IV). Simultaneous LV and RV pressure tracing on cardiac catheterization will show characteristic dip and plateau pattern (square root sign) with elevated LV and RV end diastolic pressures.

![Fig. 1](image.png)

**Fig. 1** – A: ECG recording in our patient diffuse ST depression (>7 leads), T inversion with ST elevation in lead aVR (3 mm). B: ECG after 6 months of initial admission – ST depression persisting.
function. Endomyocardial fibrosis is the commonest cause for RCM worldwide. Others include storage diseases, infiltrative disorders, hypereosinophilia and idiopathic RCM (IRCM). IRCM is considered as a spectrum of hypertrophic cardiomyopathy and associated with mutation in the sarcomeric proteins. Restrictive hemodynamic pattern without evidence for secondary causes suggested idiopathic RCM as the possible diagnosis in our patient. The crux of discussion is unique ECG pattern with normal coronaries without any perfusion defect.

In the setting of coronary artery disease widespread ST depression in >7 leads with ST segment elevation in aVR is an indicator of left main coronary obstruction, multivessel disease or proximal Left anterior descending artery occlusion. However, similar ECG pattern may be seen in common conditions like LVH characterized by ST depression, T wave inversion in left sided leads (I, aVL, V5, V6) and ST elevation in right sided leads mainly in V2 V3, rarely in aVR also. The absence of LVH by QRS voltage criteria does not exclude this possibility. Cases of RCM showing such ECG pattern

Fig. 2 — ECG reported by Rivenes et al, demonstrating biatrial enlargement ST depression in inferior lateral, lateral precordial leads, also ST elevation in aVR but not mentioned.

Fig. 3 — A: Biatrial enlargement with normal ventricles. B: Enlarged left atrium.

Fig. 4 — LA volume index = 30.7 ml/m2 (Three orthogonal D method), 36.9 ml/m2 (2 areas & length method RA area 19.8 cm2).

Fig. 5 — A: E/A >2 Grade IV diastolic dysfunction (Restrictive pattern) B: Deceleration Time 106 ms (<130).
mimicking severe subendocardial ischemia have been rarely reported.3 (Fig. 2).

ECG is a useful screening tool in RCM patients and is abnormal in nearly 90–99% of patients especially in idiopathic RCM. Bialtrial enlargement evidenced by biphasic P waves in precordial leads was reported in 91%. 74% of patients demonstrated atrial fibrillation. ST, T changes observed in nearly 80% of individuals was considered nonspecific by Naser M. Ammash et al.,2 But Rivenes et al demonstrated 3–12 mm ST depression suggestive of ischemia preceding sudden cardiac death (SCD) and Torsade’s de pontes in patients with RCM. Pathological changes of acute ischemia but without coronary obstruction were seen in autopsied heart. Hence they co-related ST depression (a sign of ischemia) to SCD in RCM.

Hayashi et al, suggested obliquely elevated ST segment with notched or biphasic late peaking T waves (67%), and ST depression with T inversion (25%) were characteristic of idiopathic RCM. By demonstrating normal epicardial coronaries in angiogram and absence of any perfusion defect on MPI, they concluded ECG changes were a reflection of

![Fig. 6](image1)  
**Fig. 6** – A: Tissue doppler imaging of lateral annulus velocity 6.9 cm/S (<8) B: Medial septal annulus velocity 7 cm/S.

![Fig. 7](image2)  
**Fig. 7** – A: Normal LV systolic function EF — 62%, normal sized LV with normal mass. B: Cardiac MR enlarged LA, RA, normal sized ventricles with normal LV mass.

![Fig. 8](image3)  
**Fig. 8** – Coronary angiogram. A: AP caudal view of LCA—showing normal left main coronary, left anterior descending and left circumflex artery. B: LAO view of RCA – Normal.
repolarization abnormalities, and SCD was due to arrhythmias rather than ischemia.6

We hypothesize that ST depression observed in precordial, inferior and lateral leads may be due to high end diastolic pressure (EDP). High EDP can produce repolarization abnormalities by impairing perfusion in the subendocardial region inducing ischemic repolarization abnormality, or by stretching the myocardium and activating stretch sensitive channels. In our patients resting ST depression lasted for more than 6 months (Fig. 1B).

Elevated EDP can induce diastolic stretching of the myocardial fibers, and this stretch may increase calcium leak at the sarcoplasmic reticulum level (Iribe et al, 2009).7 Calcium leak activates BKca (excessive in endocardium), or ClCswell channels and may alter the repolarization and induce ST T changes and arrhythmias.8 But Restrictive cardiomyopathy (RCM) myocardium too stiff to be stretched, hence stretch sensitive channel activation is unlikely to be the culprit for the observed repolarization abnormality in our restrictive cardiomyopathy patients.

High EDP will reduce the coronary perfusion pressure and induce subendocardial hypoperfusion in the ventricle globally. Diffusely impaired subendocardial blood flow in all the 3 vascular territories may not be diagnosed by SPECT Technetium 99 m MPI (False Negative).9 Hence normal coronaries and negative myocardial perfusion study by SPECT cannot exclude diffuse subendocardial hypoperfusion. ECG is a sensitive tool to diagnose subendocardial ischemia and perfusion MRI10 might be the ideal investigation for quantitative estimation of blood flow but could not be done in our patient due to non-availability.

We speculate that in our patient ischemia induced repolarization abnormality might be the most likely mechanism for ST depression, ClCswell channel activation may also be (albeit less likely) contributor for this exaggerated ST depression in our case.

4. Conclusion

Diffuse ST depression in the precordial, lateral and inferior leads with ST elevation in aVR may be seen in RCM. Severe subendocardial ischemia involving whole of the ventricle due to high EDP is a well-known phenomenon and may be most
likely mechanism of ST T changes observed in RCM. CICswell channel activation could be a less likely contributor for this exaggerated ST depression.

Conflicts of interest

The authors have none to declare.

REFERENCES

9. Denfield Susan W, Moss, Adams. Heart Disease in Infant, Children, Adolescents Including the Foetus and Young Adult. 8th ed. Lippincott Williams&Wilkins; 2013:p1267–1276 [Chapter 57 Restrictive cardiomyopathy].