CASE REPORT

A RARE CASE OF CONGENITAL LEFT VENTRICULAR ANEURYSM WITH MITRAL REGURGITATION AND ABSENT LOWER LIMB PULSATIONS
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ABSTRACT: Left ventricular aneurysms are not uncommon and mostly occur as a sequel of acute myocardial infarction. Congenital left ventricular aneurysms (CLVA) especially submitral ones are very rare and only few cases are reported in literature. Here we report a unique case of congenital sub mitral aneurysm, in a 16 year old patient presenting as mitral regurgitation with absent lower limb pulsations.

KEYWORDS: Left ventricular aneurysm, Mitral regurgitation, absent pulsations.

INTRODUCTION: Left ventricular aneurysms can be congenital or acquired. They develop most frequently after myocardial infarction. Other causes include Hypertrophic cardiomyopathy, Arrhythmogenic right ventricular cardiomyopathy, myocarditis, chest trauma, sarcoidosis, Chaga's disease and without any identifiable cause.¹ A congenital left ventricular aneurysm is a rare condition involving protrusion or out pouching of a portion of the ventricular wall with a broad connection to the ventricular cavity which may or may not have paradoxical systolic expansion.² The combination of CLVA with mitral insufficiency is rare. The largest series of these kinds of aneurysms that were found in the literature is 6 cases over a period of 11years, described by Elio Marijon et al in 2006.³

We report one such rare and unique case of CLVA with mitral regurgitation and absent lower limb pulsations.

CASE REPORT: A 16 year old male patient admitted in the department of medicine, of our hospital on 18th July 2013 was apparently all right 2 years back, when he developed insidious on set pain in right calf muscles since 18 months which was increasing on walking and subsiding on taking rest.

One month before admission he developed left sided catchy type of chest pain which was increasing on exertion and relieved on taking rest. He also had progressive exertional dyspnea (NYHA Grade I-III). He did not give any history of Paroxysmal nocturnal dyspnea, orthopnea. There was no history of palpitation, syncope, giddiness. There was no h/o cough, fever and constitutional symptoms were not present.

No h/o alcohol intake, smoking or tobacco consumption.

There was no past history of chest pain, dyspnea, palpitation associated with fever. No past history of chronic cough, joint pains, skin lesions. No past h/o exposure to sexually transmitted disease.

No past history of trauma to the chest.

No family history of premature deaths or history suggestive of cardiac disease.

ON EXAMINATION: He was moderately built and moderately nourished. He had no pallor, icterus, cyanosis, pedal edema and was afebrile. His right radial pulse was 84/min, irregular, varying in
volume and character. Popliteal, dorsalis pedis, posterior tibial were not palpable bilaterally. His blood pressure was 120/84 mm of Hg in right arm. Supine position and JVP was not raised.

Examination of cardiovascular system revealed prominent S₃ and a high pitched pan systolic murmur of grade 3/6 heard at the apex, radiating to axilla. P₂ was loud.

Respiratory system and per abdominal examination were normal. No neurological deficit was present.

On investigating, the patient had normal complete blood counts. ASLO titers were 61.9IU/lit, CRP 1.87mg/dl, VDRL, HIV, HBs Ag, Hepatitis C antibody were negative. INR 1.34 with Ratio 1.33%. Prothrombin time 14.0 sec. Serum electrolytes were normal.

The patient’s ECG-showed heart rate of 102 /min, prolonged PR interval, ST depression in II,III, aVF, V₃ to V₆, and borderline LVH. Chest x- showed increased cardio thoracic ratio of 0.6 with normal lung fields.

Holter monitoring showed prolonged PR interval, Mobitz type II heart block and sinus pauses. Trans thoracic echocardiography (TTE) revealed large LV Aneurysm of sub mitral type. Below and posterior to aortic valve, of size 52x33mm, communicating with LV cavity (Fig. 1). Anterior mitral leaflet was distorted due to the aneurysm. Moderate eccentric mitral regurgitant jet was seen, directed posteriorly (Fig. 2). Left atrium, left ventricle mildly dilated. Good left ventricular and right ventricular function. No clot/vegetation. Aorta and arch were normal.

Patient was subjected to trans esophageal echocardiography (TEE) which confirmed large sub mitral LV aneurysm (Fig. 3).
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Patient underwent coronary angiogram (CAG), left ventricular and peripheral arterial angiogram (PAG) via right brachial artery as a part of evaluation. CAG revealed normal epicardial coronaries, aorta and sub clavian arteries. LV angiogram confirmed large sub mitral aneurysm communicating with LV cavity. (fig. 4).

PAG revealed normal iliac vessels. Renal arteries normal. 20-25% lesion at right common femoral artery. Both superficial femoral arteries showed long segment lesions in mid-thigh region. Reformation of both popliteal arteries seen by collaterals (fig. 5). Both anterior tibial arteries are not visualized; posterior tibial arteries are under filled.

With findings of normal coronaries in a young patient and after excluding infective and other acquired causes of LVA clinically and by ECHO, TEE and CAG, we made a diagnosis of CONGENITAL, SUB MITRAL, LEFT VENTRICULAR ANEURYSM WITH MITRAL REGURGITATION.

Patient was put on T. Ecosprin 150mg 0-1-0. Acitrom 2mg o-o-1 and was advised to undergo aneurysmectomy at the earliest.

DISCUSSION: Most cardiac LVAs are acquired due to coronary artery disease & myocardial infarction (MI) in adult hood. Uncommonly other causes of LVA are reported. LVAs could be of cardiac and non cardiac etiology. Cardiac causes apart from MI are coronary artery fistula, Hypertrophic obstructive
cardiomyopathy, Arrhythmogenic Right ventricular cardiomyopathy, and myocarditis. Non-cardiac causes include sarcoidosis, Chagas disease, Systemic lupus erythematosus, Behcet's disease, tuberculosis, HIV, Syphilis and Glycogen storage disease and blunt chest trauma. A congenital LVA is a rare cardiac manifestation, a study reported an incidence of 0.4% of 750 cardiac necropsies. 411 cases have been reported since its first description in 1816. Elio Marjoli et al reported 6 cases over a period of 11 yrs. Very few cases of congenital sub mitral LVAs are reported in literature. Our case of congenital sub mitral LVA is unique in presentation with claudication pain due to long segment blocks in both superficial femoral vessels. This could be due to multiple silent thrombo embolic episodes, however our case did not show any LV clot.

LV aneurysms must be distinguished from Left ventricular diverticula. The walls of a diverticulum are formed by all three layers and contracts normally, whereas an aneurysm is generally a fibrous saccular region with paradoxical contraction. The true incidence of sub–valvar aneurysms is certainly underestimated, especially in the early stages when there are no clinical symptoms. The etiology of sub valvar aneurysm is mostly thought to be a congenital weakness of the fibrous annulus of the valve, which is further supported by the predominantly race group as well as anatomy and embryology of the affected area.

Most LVAs are asymptomatic and are occasionally identified during routine diagnostic procedures. Sudden cardiac death could be the first manifestation. They may present with systemic embolism, valve regurgitation, ventricular rupture, or ventricular arrhythmias, sometimes life threatening.

Our patient did not have clinical features of arrhythmias, but ECG showed conduction disturbances.

Surgical repair is the gold standard; however controversy exists about surgical approach and technique. In view of their high rate of complications some authors advocate surgical resection even if they are asymptomatic. ICD and Radio frequency catheter ablation have been used successfully for the treatment of VT. Prognosis of LVAs depends upon size, location, degree of valve involvement, functional class and presence of arrhythmias.

CONCLUSION: Congenital left ventricular aneurysms though rare, should be considered in any young patient with unexplained symptoms of heart failure. They can cause life threatening complications like ventricular tachycardia and sudden death, which are preventable by surgery, intra cardiac defibrillator, or radio ablation. CLVAs can be easily picked up by transthoracic echocardiography.

REFERENCES:


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