Cardiac Mass and Hypertrophic Cardiomyopathy as Aggressive Presentation of Primary Cardiac Lymphoma: A Case Report*

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ABSTRACT

Primary cardiac lymphoma is very rare. This disease occurs mainly in immunocompromised patients and rarely in the immunocompetent. With the advent of echocardiogram, computed tomography and magnetic resonance imaging, more cases are being diagnosed ante-mortem although the prognosis still remains poor. These cardiac tumors provide unique diagnostic and therapeutic challenges. We present an interesting case of primary cardiac lymphoma diagnosed ante-mortem in an 81 year old immunocompetent female presenting as a large mass in the right atrium, right ventricle and encasing the right coronary artery; and asymmetric left ventricular hypertrophy, a variant of hypertrophic cardiomyopathy. Patient responded very well to chemotherapy with subsequent regression of the cardiac mass and ventricular hypertrophy.

Keywords: Primary Cardiac Lymphoma; Asymmetric Left Ventricular Hypertrophy; Chemotherapy

1. Introduction

Primary malignant cardiac tumors are rare, with the most common being sarcomas. Primary cardiac lymphoma (PCL) comprises about 0.5% of all lymphomas and occurs generally in the right atrium and ventricle [1]. PCL most commonly are diffuse B-cell lymphomas and are more common in males with a mean age of 67 years at presentation [2]. With the advent of echocardiogram, computed tomography and magnetic resonance imaging more cases are being diagnosed ante-mortem although the prognosis still remains poor [2]. The incidence has also increased in immunocompromised patients especially with Acquired Immunodeficiency Syndrome (AIDS) and have a poorer outcome. PCL patients can present with chest pain, shortness of breath, pulmonary edema, arrhythmias, atrioventricular blocks, left ventricular hypertrophy [3,4]. We present an interesting case of PCL diagnosed ante-mortem in an 81 year old immunocompetent female presenting as a large mass in the right atrium, right ventricle and encasing the right coronary artery; and asymmetric left ventricular hypertrophy, a variant of hypertrophic cardiomyopathy.

2. Case Report

An 81-year-old female with no significant past medical history presented with 2 week history of increasing dyspnea on exertion and fatigue. She denied any chest pain, palpitations, fever, cough or recent travel. She had lost about 8 pounds over the last 1 month. Human Immunodeficiency Virus (HIV) test was negative. An echocardiogram (Figure 1) revealed a lobulated mass 5 × 4 × 3 cm attached to the tricuspid valve, extending into the right atrium, right ventricle, asymmetric left ventricular hypertrophy, a variant of hypertrophic cardiomyopathy.

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1. Introduction

Primary cardiac lymphoma is classically defined as an extranodal non-Hodgkin’s lymphoma exclusively located in the heart and/or pericardium and is extremely rare. This disease occurs mainly in immunocompromised patients [3,6] and rarely in the immunocompetent [3,7,8]. It accounts for 1.3% - 2% of all cardiac tumours and 0.5% of all extranodal lymphomas [1,7,9], but it has been encountered with increasing frequency in patients with Acquired Immune Deficiency Syndrome (AIDS) or other severe immunodepressive syndromes. Most of the reported primary cardiac lymphoma cases are of B-cell origin, while T-cell cardiac lymphomas are much rarer.

Thirty-five cases of primary cardiac non-Hodgkin’s lymphoma were reviewed by Chalabreysse et al. [10], of which twenty-two of these cases were diffuse, large B-cell lymphoma (DLBCL) [10]. Similarly, Pertrich et al. [3] reviewed 197 cases of PCL, with half of all cases reported since 1995. The study is the largest analysis of PCL to date. Diffuse large B-cell lymphoma constituted a majority, and survival in PCL was affected mainly by 4 factors: immune status, left ventricular involvement, presence of extra-cardiac disease, and arrhythmia. Prognosis generally remains poor, however, lack of ventricular involvement and presence of arrhythmias were associated with improved survival. Mayayo Artal et al. [11] also described PCL among immunocompetent hosts, in which it was observed that Ebstein Barr Virus appears to play no role in the development of PCL, and this finding was also observed in a review by Lee et al. [12].

There have been other reports of immunocompetent patients with diffuse B-cell lymphoma presenting with SVC syndrome [8], cardiac tamponade, life threatening arrhythmias and intractable heart failure. Because of this rapid evolution, the situation represents an oncologic emergency, and therefore early diagnosis and treatment are crucial.

Most cases of cardiac lymphoma are solid, infiltrative nodule tumors in one or multiple chambers. Clinical series by Caresoli et al. [7], and Pertrich et al. and a clinicopathologic series by Chim et al. [13] revealed that primary cardiac lymphoma usually arise in the right chambers as a mass with or without pericardial effusion among >80% of the patients, and the most common presentation was unresponsive heart failure.

Lymphomatous infiltration of pericardium is also seen in a number of cases [2,13]. However the diffuse infiltration in all cardiac chambers along with intracardiac mass is rare and represents an aggressive presentation as seen in our patient. Massive infiltration of the myocardium results in irregular thickening of the walls mimicking hypertrophic cardiomyopathy and is generally associated with malignant lymphoma with spread to the myocardium [3,14]. A case of PCL presenting solely as hypertrophic obstructive cardiomyopathy was described by Lee et al. and diagnosed postmortem [12]. Similarly, Kadhim et al. [5] and Cabin et al. [15] in different case reports described extensive myocardial infiltration causing concentric left ventricular hypertrophy and mimicking hypertrophic cardiomyopathy. A substantial amount of the myocardium was replaced by infiltration of the tumor. The phenomenon of infiltration in these cases are similar to our case, in which there was extensive infiltra-

2. Case Report

A 68-year-old male with a history of hypertension, diabetes mellitus, and coronary artery disease presented with dyspnea on exertion. Transthoracic echocardiography revealed a large irregular inhomogeneous mass measuring 5 × 4 × 3 cm in right heart adjacent to tricuspid valve and infiltrating the walls of RA and RV. (RA—right atrium, RV—right ventricle, LA—left atrium, LV—left ventricle).

Figure 1. Transthoracic Echocardiography showing large irregular inhomogeneous mass measuring 5 × 4 × 3 cm in right heart adjacent to tricuspid valve and infiltrating the walls of RA and RV. (RA—right atrium, RV—right ventricle, LA—left atrium, LV—left ventricle).

Magnetic resonance imaging showed a large lobulated mass attached to the tricuspid valve. Mass projects into right atrium and right ventricle. (RA—right atrium, RV—right ventricle, LV—left ventricle.

Figure 2. Magnetic resonance imaging showing large lobulated mass attached to the tricuspid valve. Mass projects into right atrium and right ventricle. (RA—right atrium, RV—right ventricle, LV—left ventricle.

and asymmetric left ventricular hypertrophy which completely resolved after chemotherapy.

3. Discussion

Primary cardiac lymphoma is classically defined as an extranodal non-Hodgkin’s lymphoma exclusively located in the heart and/or pericardium and is extremely rare. This disease occurs mainly in immunocompromised patients [3,6] and rarely in the immunocompetent [3,7,8]. It accounts for 1.3% - 2% of all cardiac tumours and 0.5% of all extranodal lymphomas [1,7,9], but it has been encountered with increasing frequency in patients with Acquired Immune Deficiency Syndrome (AIDS) or other severe immunodepressive syndromes. Most of the reported primary cardiac lymphoma cases are of B-cell origin, while T-cell cardiac lymphomas are much rarer.

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tion involving the right chambers and he right coronary and asymmetric left ventricular hypertrophy however, there was a good response to treatment in our case. We are presenting the first case with right sided large mass and diffuse wall infiltration in an immunocompetent subject, mimicking hypertrophic cardiomyopathy and with good response to chemotherapy.

The clinical symptoms of PCL are nonspecific and related to the location of the tumor. Presentations could be acute including dyspnea, edema, arrhythmias and rarely cardiac wall rupture [7]. Other clinical manifestation includes conduction defects leading to tachy- or brady-arrhythmias [6], pericardial effusion and heart failure. Symptoms often include chest pain, exertional dyspnea, palpitations, syncope and other non-specific constitutional symptoms. Diagnosis can be made by echocardiogram, CT and MRI followed by tissue biopsy [7,9]. This case is also interesting as it emphasizes the advantage of echocardiogram for diagnosis and serial follow up.

Lymphomas should be considered in the differential diagnosis of cardiac mass and asymmetrical LV hypertrophy presenting as hypertrophic cardiomyopathy. Because the tumors are largely B-cell lymphomas, they are sensitive to chemotherapy with main regimen as CHOP (cyclophosphamide, doxorubicin, vincristine + prednisone) +/- rituximab [7,13]. The mortality is high due to delay in diagnosis and high relapse rate but survival up to 192 months also has been documented [16]. Extra care is required in the early stage of treatment as tumor lysis may lead to cardiac complications including rupture of the heart or asystole [17].

In conclusion, the diagnosis of primary cardiac lymphoma is difficult due to non-specific clinical manifestations and it should therefore be considered in patients with a cardiac mass with or without pericardial effusion. It is confirmed using imaging modalities such as transthoracic echocardiography and magnetic resonance imaging and certified using cytology or open biopsy. The most effective treatment is chemotherapy, but prognosis remains poor. Definitive approaches to obtain early histological diagnosis are important in order to initiate early treatment.

REFERENCES


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