Surgical Management for a Rare Pedunculated Left Ventricular Apical Lipoma: A Case Report and Review of Literature

Long Song†, Chukwuemeka Daniel Iroegbu†, Jinfu Yang and Chengming Fan*

Department of Cardiovascular Surgery, The Second Xiangya Hospital, Central South University, Changsha, China

Cardiac lipomas, though extremely rare, are encapsulated tumors composed primarily of mature fat cells. Despite their benign character, cardiac lipomas can cause life-threatening complications by rapid growth. Cardiac lipomas, which are frequently located in the left ventricle (LV) or right atrium, can originate either from the subendocardium, subpericardium, or the myocardium. They are usually asymptomatic and carry a good prognosis during long-term follow-up; however, published reports show that untreated cardiac lipomas may be fatal when they cause arrhythmic or obstructive symptoms. In addition, several surgical options have been reported to obtain an appropriate operative view following poor visualization, primarily when tumors are located in the LV. Herein, we present a case of a pedunculated LV apical lipoma in a symptomatic patient successfully managed by surgical resection. We also discuss diagnostic modalities in surgical planning and the choice of surgical approach.

Keywords: cardiac surgery, computed tomography, left ventricular lipoma, primary cardiac tumors, transthoracic echocardiography

INTRODUCTION

A round 75% of primary cardiac tumors are benign, with an incidence of 0.2–0.4% in autopsy series (1). Cardiac lipomas, though extremely rare with a reported incidence ranging from 2.9 to 8% among all benign cardiac tumors (1) and 10% of all neoplasms of the heart (2), are encapsulated tumors composed primarily of mature fat cells. Cardiac lipomas are 50 times less common than myxomas and usually present in combination with lipomas of other organs. These tumors are often asymptomatic and usually detected incidentally, mostly during autopsies. In symptomatic patients, the diagnosis can easily be made by echocardiography, computed tomography (CT), or magnetic resonance imaging (MRI).

The etiology of lipomas remains uncertain; however, an association with chromosome 12 gene rearrangements has been established in solitary lipoma cases with an abnormality in the HMGA2-LPP fusion gene (3). These tumors are typically found in adult patients in their fifth or sixth decade of life but can affect patients of all ages, including both sexes (4). Lipomas of the heart can be intracavitary, epicardial, or intramyocardial and may be located in the atria or ventricles. Although lipomas are benign, these tumors can cause cardiac chamber(s) compression, chest tightness, arrhythmias, obstructive symptoms, and other life-threatening complications when the tumor increases in size (5). Conservative management includes clinical observation and, in some cases, anticoagulation. Surgical management includes resection of the mass, which is considered curative. Whether one approach is superior to the other is unknown given the lack of data and the tumor’s rarity.
Herein, we present a case of a pedunculated left ventricular (LV) apical lipoma in a symptomatic patient successfully managed by surgical resection. We also discuss diagnostic modalities in surgical planning and the choice of surgical approach.

**CASE PRESENTATION**

A 52-year-old male was referred to our institution complaining of chest tightness and dyspnea on exertion the last few years, which only progressively worsened the last few months (NYHA III) with palpitations and edema of the lower limbs. The patient had neither previous cardiovascular antecedents and denied familial history. He also denied medication use with only occasional smoking and alcohol intake. The patient's body temperature was normal on admission, 36.8°C with blood pressure 126/82 mmHg, and a radial pulse rate of 83 bpm. The patient body mass index was 24.2.

Complete blood count, basic metabolic profile, B-type natriuretic peptide, and troponins were also within normal limits. The biochemical investigation showed no alteration in the following serum levels: thyrotropic hormone, potassium, alkaline phosphatase, glycemia, sodium, hyaluronic acid, urea, total bilirubin, and creatinine. Hemoglobin level was 103 g/l, hematocrit level was 31.4%, while serum leukocyte and platelet levels were 6.61 × 10^9/L and 214 × 10^9/L, respectively.

Physical examination and vital signs were unremarkable. The patient had good psychomotor development and no neurological signs or nystagmus. On cardiovascular examination, the patient had poor peripheral perfusion, asymmetric pulses, postural hypotension, peripheral lower limb edema, and a mild ejective cardiac murmur at the mitral region. ECG showed non-specific T-wave abnormalities in the lateral leads. The lungs were clear to auscultation, while other body systems showed no abnormalities on physical examination.

**Radiological Investigations**

Chest radiograph showed a moderately enlarged cardiac silhouette with increased cardiothoracic ratio and no signs of pulmonary congestion. Transthoracic echocardiogram (TTE) examination showed the following characteristics: aorta 34 mm, left atrium 35 mm, LV cavity 51 mm (end-diastolic diameter), and a hyperechoic mass 30 × 27 mm located on the left side of the heart (middle-apical region), occupying almost 2/3 of the LV with blood supply from the left anterior descending (LAD) coronary artery (Figure 1). The vast mass caused a marked dilatation of the LV with mild to moderate LV outflow tract (LVOT) obstruction and decreased ventricular ejection. In addition, the apical and diaphragmatic part of the tumor was mildly calcified. CT scan of the chest confirmed a mass measuring 28 × 32 mm (Figure 2). The hyperechoic mass was highly mobile, with a peduncle measuring ~4 cm in diameter, adhering to the apical diaphragmatic surface of the LV free wall and papillary muscle.

**Differential Diagnoses**

The characteristics of the mass following clinical investigations were suggestive of a fat-containing lesion at the LV apex. Hence, the differential diagnoses were relatively limited to subendocardial fat collection, LV thrombus, lipomatous hypertrophy, fibroma, intracardiac varix, and lipoma. However, lipomatous hypertrophy primarily occurred in the interatrial septum and was therefore judged less likely given the location of the patient’s mass. Nonetheless, other intracardiac masses such as teratomas, fibroma, and myxoma could not be initially excluded.

**Management**

The patient was anticoagulated with apixaban as a “bridge” to definitive surgical therapy. The case and radiological reports were discussed in a multi-disciplinary heart team meeting, and the final consensus for a surgical resection was reached. Likewise,
the case was also discussed with the patient. A unanimous decision was made to proceed with the surgical resection. Surgery was performed via median sternotomy, and cardiopulmonary bypass was established by aortic and two-stage single venous cannulation (mild hypothermia at 30°C). An aortic cross-clamp was applied, and myocardial protection was established with an antegrade infusion of cold Custodiol® HTK solution via the aortic root.

The left heart and the LV mass were approached via a left atriotomy approach. However, given the extensive and deeply located position of the mass at the LV apex with mild adherence to the LV free wall and papillary muscles, we inserted a mediastinoscope into the LV. The mediastinoscope provided a more detailed view than the usual valve retractors, providing a safe resection of the mass (Figure 3). The tumor looked like a lobulated fatty mass, which was well-encapsulated with a yellowish appearance. Under direct vision, the lipoma was resected using scissors, and the LV was rinsed and cleared with long laparoscopy forceps and continuous suction.

The operation was uneventful. Hemostasis was achieved, and the chest was closed after placing one chest tube in the pericardium. The patient was discharged from the anesthetic recovery room on the 2nd post-operative day. Apixaban was discontinued; however, whether or not post-operative anticoagulation is needed is debatable. Intraoperative histopathologic examination of the excised specimen revealed fatty structure (mature adipose tissue) with areas of increased vascularity (Figure 4). The results were consistent with benign encapsulated lipoma. The post-operative course was also uneventful, and the patient was discharged from the hospital on the 6th post-operative day.

**DISCUSSION**

Cardiac lipomas of the heart are sporadic tumors, accounting for 2–8% of all benign cardiac tumors (6, 7) and incidentally discovered during routine chest X-rays or cardiac investigations. Lipomas occur at an equal frequency in both genders (8), with the typical “age at presentation” between 40 and 60-years of age (9). Cardiac lipomas, particularly in the early stages, are usually indolent and asymptomatic and can originate either from the subendocardium, subpericardium, or myocardium and may be located more frequently in the right atrium, LV (7, 10), or both ventricular cavities (11). Symptoms are not characteristic. Generally, most cardiac lipomas are silent, with only a few situations where clinical symptoms manifest, depending on their location and size. Thus, angina, arrhythmia, and syncope are symptoms a patient may complain of on routine clinical visits.

In the case herein, the left-sided heart failure symptoms were precipitated by the (fat-attenuating) pedunculated intracardiac mass obstructing the LVOT (Sun et al.). Also, the rare issue with lipoma does not reside in its histological characteristics but its intracavitary component. Thus, the presence of cardiac lipoma becomes lethal when it occupies the LV cavity, resulting in obstruction and compression, which eventually causes peripheral embolism, LV failure, and rhythm and conduction disorders (12–14). The patient's symptoms in the presented case were those of heart failure with dyspnea and peripheral edemas. Given that lipomas are well-encapsulated,
lipomatous hypertrophy of the interatrial septum due to their lipomas are distinctively differentiated from adiposarcomas and characterize the mass. Furthermore, on cardiac imaging, cardiac is challenging. Hence, more dedicated imaging is warranted to resolution (views also distant from the transducer, thus limiting its cardiac CT and cardiac magnetic resonance imaging (CMR) can be pursued to provide vital information concerning the extent of myocardial infiltration and tissue characterization and help differentiate a fat-containing lesion from a thrombus (7, 10). However, in the case herein, only cardiac CT was used. Notably, low-attenuation features with a density similar to fat on CT are pathognomonic for lipoma, as with the presented case. In addition, coronary arteriography though not used in the case herein, can provide crucial information by outlining the arterial supply and defining the coronary anatomy, which will be essential for the operating team.

The total surgical resection of lipomas with its pedicle is advocated (as with the presented case herein) to prevent a recurrence, whether the lipoma is intracardiac or subcutaneous in origin. However, there are significant surgical challenges and considerations when treating intracavitary lesions, such as (i) the maintenance of adequate atrioventricular valvular function, (ii) preserving an adequate portion of the ventricular myocardium, (iii) the satisfactory excision of the lesion from the LV without injuring it, and (iv) preserving the conduction system. Currently, given the rarity of the lesion, there are no evidence-based guidelines concerning their optimal treatment. Thus, conservative vs. surgical management decision solely relies on factors such as the patient’s preference, symptomology, the lesion size and location, and surgical know-how.

Given the location of the lesion and the stalk deeply attached to the cardiac apex in the presented case, a left atriotomy approach was opted for, which, however, proved to be a surgical “pitfall.” Though the usual surgical approach for lipoma resection is through a median sternotomy, other surgical methods have been used with excellent results, particularly for apical and ventricular lesions. The reported surgical approaches include: (i) the LV tip approach; a similar incision used for LV aneurysms, which provides adequate exposure (preventing partial resection) and decreases the risk of embolization; (ii) video-assisted removal of coronary lesions, which improves lighting inside the surgical field and avoids left ventriculotomy and the possible damage of surrounding structures (7, 23).

Though prompt and careful pre-operative qualitative diagnosis is necessary (7), definitive diagnoses for cardiac lipomas are made on post-operative pathological examination irrespective of the pre-operative diagnostic modalities used. In the presented case, cardiac lipoma following post-operative pathological examination was confirmed given the presence of mature, differentiated fatty tissue, with small amounts of fibrous connective tissue and blood vessels, which were surrounded by a fibrous membrane. Given that post-operative prognoses for lipomas are favorable, surgical managements are without complications such as LV systolic dysfunction and prolonged follow-up (as with the presented case), ventricular septal defects, and even death (8), primarily when performed late or with the use of a “non-suitable” surgical approach.

CONCLUSION

Though there are no evidence-based guidelines concerning the optimal treatment of the disease given its rarity, the case herein exemplifies how early diagnosis, appropriate cardiac investigations, and surgical management of patients with cardiac lipoma should be individualized for favorable long-term outcomes. Also, with cases of apical cardiac lipoma, cardiac investigations should include coronary arteriography to define coronary anatomy and outline arterial supplies.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.
ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Ethics Committee of the Second Xiangya Hospital of Central South University. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

CI drafted the manuscript. CF and LS designed the study. JY and CF revised the final manuscript. LS was responsible for the collection of data or analysis. All authors have read and approved the final manuscript.

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