Giant Interatrial Septal Lipoma Filling the Right Atrium Causing Slight Symptoms: A Case Report

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ABSTRACT

Cardiac lipoma is extremely rare. Here, we present a unique illustrative case of an interatrial septal lipoma protruding into the right atrium, causing symptoms in a 54-year-old male. Echocardiogram and computed tomography showed a well-shaped, giant, and fixed mass located in the interatrial septum and right atrium. The only manifestation was palpitation, though the mass filled almost all of the atrium and compressed the superior vena cava. The patient received resection of the large-sized lipoma, which was 87mm in diameter and weighed 300-400g. Pathological exam demonstrated mature lipocytes and substantiated the diagnosis of lipoma. The patient did well postoperatively, and symptoms were resolved.

CASE PRESENTATION

A 54-year-old East Asian male patient was admitted to the local hospital with a chief complaint of recurrent spontaneous paroxysmal palpitation for one year. This symptom became more frequent for one month. No aberrant signs were found on physical examination. The patient's NT-proBNP was 493.30 pg/ml (normal range: 0-125 pg/ml). The cardiac function of the patient was class I on NYHA classification. The rhythm was sinus on electrocardiogram (ECG). Echocardiogram presented an ovoid mass located in the right atrium and tightly attached to the interatrial septum. The border of the mass was clear, except the adherent part (Figure 1A, 1B). Left ventricular ejection fraction was 58%. Coronary computed tomography angiography (CTA) revealed a large mass within the interatrial septum and right atrium that took up nearly the entire roof of the right atrium, constricting the superior vena cava (SVC). The upper part of the mass surrounded the aortic root. The size of the mass was 58mm×87mm×52mm, and the density was distinctly lower than the peripheral tissues (Figure 1C).

The patient was diagnosed with cardiac lipoma and received surgery through a median sternotomy under cardiopulmonary bypass. An ovoid, yellowish and brownish mass was exposed between the right auricle and SVC after opening the patient's pericardium (Figure 2A). The mass was well-capsulated, vessel free, and connected with the interatrial septum. Through right atrial atriotomy, a firm, lipid change was exposed (Figure 1B). The patient did well postoperatively, and symptoms were obviously relieved. Postoperative CT revealed no anomaly (Figure 3B). Frequent ventricular premature beats were found on postoperative ECG when the patient was suffering palpitation (Figure 3C). The patient complained of no other discomfort during the next eight days of in-hospital follow up, and he eventually was discharged. The patient remained asymptomatic on follow up over the next 15 months.

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DISCUSSION AND CONCLUSION

Cardiac lipomas mostly occur in the age group of 40-60 years [Ismail 2015]. Patients with cardiac lipoma usually are asymptomatic, thus the tumor often is found incidentally during imaging check or autopsy [Bois 2014]. Symptoms caused by cardiac lipoma depends on its size and location. In this reported case, the only clinical manifestation of the patient was palpitation. Adipocyte infiltration in the myocardium attributes to perturbation of myocardial conductivity and leads to arrhythmia [Wong 2017]. Although the preoperative ECG of this patient was normal, postoperative ECG revealed frequent ventricular premature beats, alongside the HE staining result that mature fat cells and myocardial cells are crossing, indicating that the patient's preoperative palpitation probably was caused by adipocytes causing related arrhythmia. The mechanisms of adipocytes disturb cardiac conductivity is so far poorly elucidated and still in progress of research. Mitochondrial dysfunction, autonomic dysfunction, autophagy, oxidative stress, mitophagy, and myocardial death may all play vital roles in inducing aberrant signals [Pabon 2018].

Cardiac lipoma accounts for 5% of primary cardiac tumors, it develops from the mesoderm and arises subendocardially, subepicardially, and myocardially [Puvaneswary 2000]. Among these origins, lipomas of subendocardial and subepicardial origin are encapsulated and partially encapsulated as for myocardial origin. Cardiac lipoma can occur anywhere in the heart, of which interatrial septum is the most common position [Monti 2015]. In this case, lipoma at the right atrium was well-enclosed, while in the interatrial septal area, no capsule was presented. It baffled the surgeons, however, whether the origin of the cardiac lipoma in this case arose from the interatrial septum and protruded to the atrium or arose from the epicardium and infiltrated to the interatrial septum.

Because of the location and the traits of lipoma, interatrial septal lipoma usually is differentiated with lipomatous hypertrophy in interatrial septum (LHIAS). LHIAS and interatrial septal lipoma are both benign lipomatous changes in the heart, consisting of mature fat cells. LHIAS, occurring in 2% of the population, involves interspersed lipocytes growing within the interatrial septum and making it thicker than 2cm, and the fossa ovalis is usually spared [Poterucha 2019]. Of note, differing from lipoma, LHIAS is unencapsulated and presents a typical dumbbell shape on CT scan [Hudzik 2010].

Though the etiology of cardiac lipoma is under investigation, genetical change can strongly be related. The rearrangements of two gene members of the high-motility group family, HMGA1 (formerly HMGIY, 6p21) and HMGA2 (formerly HMGI-C, 12q14) have demonstrated involvement in the pathogenesis of 65% soft tissue lipomas [Rodriguez 2008]. Moreover, amplification of MDM2 and/or CPM, derived from chromosome 12q13-15, also are implicated in lipomatous change [Erickson-Johnson 2009]. Nevertheless, in one study, HMGA2 rearrangement was found in 42% cardiac lipomas and in 43% LHIAS cases, while no HMGA1 rearrangement and MDM2/CPM amplification were demonstrated in either lipomas or LHIAS [Bois 2014]. Therefore, genetic screening can be performed for strengthening the diagnosis and also may be an effective prevention and treatment of cardiac lipoma in the future.
The scarcity of cardiac lipomas presently brings no treatment guideline. In this reported case, resection was inevitable. SVC was deformed by the tumor, though no symptoms of obstruction were shown, further growth of the mass could have led to potential SVC obstruction syndrome and venal thrombus formation. Even if a lipoma is benign, it can expand in size and oppress the heart, shrinking the volume of the cardiac chambers and impairing heart function, which can be fatal. Furthermore, due to part of this lipoma not being encapsulated, fat liquefaction can occur and subsequently cause fat embolism. The surgical option can decide the outcome of patients with cardiac tumors. Naseerullah et al [Naseerullah 2018] reported a large cardiac lipoma case with surgery removal and reconstruction of the atrial septum and roof of the right atrium using a patch. No patch, however, was used in our case to avoid risk of thrombogenesis and other unpredictable symptoms like arrhythmias. Surgery provides 95% cure of cardiac benign tumor [Naseerullah 2018], however, surgeons should take the surgery option into consideration so that potential hazards get evitable on patients.

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