Asymptomatic Left Atrial Myxoma in Elderly Patient: A Case Report

Yaşlı Hastada Asemptomatik Sol Atrial Miksoma: Olgu Sunumu

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Abstract

Approximately 75% of cardiac tumors are benign and approximately 50% are constituted of myxomas. Myxomas are more common among women, and generally appear between the third and sixth decade of life. The most common symptom of myxomas is respiratory distress associated with obstruction. The main goal in the surgery of cardiac myxomas is the full excision of the tumor with minimal manipulation and disruption/fragmentation together with the aid of a cardiopulmonary bypass. This case report is reported a seventy years old patient with asymptomatic cardiac myxoma whom undergoing surgery with general anesthesia twice at different times and undiagnosed cardiac myxoma in past seven years.

Keywords

Cardiac Tumor; Myxoma; Atrial Septal Defect; Cyanosis; Angiography

Özet

Kardiyak tümörlerin yaklaşık %75’i benign olup yaklaşık %50’sini miksomalar oluşturmaktadır. Miksomalar kadınlar daha sıktır ve genellike hayatın üçüncü ve altınınci dekaden arasında görülenirler. En sık görülen semptomu obstruksiyona bağlı solunum zorludur. Kardiyak miksomaların cerrahisinde ana prensipler kardiyopulmoner bypass eşliğinde tümörün minimal manipülasyonla ve parçalanma olmadan tam eksizyonudur. Bu olgu sunumu, son yedi yıl içerisinde farklı zamanlarda general anestezide iki kez cerrahi geçirmiş ve kardiak miksoma tanısı konulamamış, yetmiş bir yaşında, asemptomatik kardiak miksomalı yaşlı bir hastayı bildirmektedir.

Anahtar Kelimeler

Kalp Tümörleri; Miksoma; Atrial Septal Defekt; Siyanoz; Anjiografi

DOI: 10.4328/JCAM.2410 Received: 17.03.2014 Accepted: 31.03.2014 Published Online: 01.04.2014
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Introduction
The incidence of primary cardiac tumors, which are uncommon in all age groups, is 1.7-10 in 100,000 within the population. Approximately 75% of these tumors are benign, and approximately 50% are constituted of myxomas.[1] Myxomas are more common among women, and generally appear between the third and sixth decade of life.[2] Myxomas are pathologies that may have fatal outcomes due to their associated complications, and most patients with this condition experience embolization, cardiac symptoms associated with obstruction, and structural symptoms.[1,3] The most common symptom is respiratory distress associated with obstruction. The clinical features of myxomas varies according to tumor localization, size and motility, and tumors are generally observed on the left side of the heart (75% in the left atrium).[3,4] Transthoracic and transesophageal echocardiography tests are the most accurate, reliable and commonly used diagnostic tools, and provide important information regarding the imaging and characteristics of the tumor. The first surgical excision of a myxoma was reported by Bahnson in 1952. The main goal in the surgery of cardiac myxomas is the full excision of the tumor with minimal manipulation and disruption/fragmentation together with the aid of a cardiopulmonary bypass.

Case Report
71-year-old female patient admitted to the hospital with complaints one time fainting temporarily and dizziness at the last two weeks of sometimes. Connective tissue disease, cerebrovascular disease, malignancy, rheumatic fever, radiation exposure, renal failure, metabolic disorders, peripheral embolism and chest trauma haven't be at resume questioning of patient and there isn't palpitation, chest pain, weight loss and respiratory distress story. In the last seven years, patient has a history of two operations because of the bile duct stone and ileus that made under general anesthesia and examinations and preparation tests conducted before these operations had not received the diagnosis of cardiac myxoma. At physical examination, blood pressure was 110/70 mmHg, pulse was 78/sec, fever was 36.6 degree and respiratory rate 20/min respectively. On inspection there were no examination findings other than hemorrhagic rash outside of the chest wall and abdominal operation scars. Evident on palpation of peripheral pulses. There was no sound except for the pathological murmur apical 1-2/6 presistolik at auscultation. Chest X-ray showed an increase in vascularity in the lungs. Electrocardiography were in sinus rhythm. There was not the result of abnormal hematology and biochemical tests (WBC: 5300, CRP: 0.6 mg / dl and Sedimentation: 22/ hour). Echocardiography were used in preoperative diagnosis. In the left atrium a mass was found with transesophageal echocardiography after transthoracic echocardiography which depending on the size 31x22mm, connected interatrial septum with a handle about 1 cm length and showing calcifications (Figure 1). Ejection fraction of patient was 67% with echocardiography. Due to the advanced aged patient coronary angiography was built and there was no significant coronary artery disease. The patient was operated under general anesthesia with a midline sternotomy, cardiopulmonary bypass has been entered following ascending aorta and bicaval cannulation. Moderate hypothermia and cardiac arrest was achieved with blood cardioplegia. The left and right atria were opened and followed yellowish gelatinous mass in the left atrium, stem and base of origin with the interatrial septum was excised. Interaltrial septal defect was closed using a PTFE patch. After excision of the tumor mass the cardiac chambers were checked and due to absence of any mass, cardiac chambers were irrigated with cold saline over and over again. The mitral valve was found to be normal. The patient was not made an additional attempt. The histopathological examination of the excised mass which sized 50X40X30mm was built and is compatible with myxoma (Figure 2, 3). Postoperative period was uneventful and the patient was discharged on the fifth postoperative day. Patients were followed without recurrence for a year without any problems.
Discussion
Cardiac myxomas are more common among women and between the ages of 30-60, and are frequently localized to the left atrium.[1,3] Our patient was a 70 years-old women with a left atrial myxoma rooted in the fossa ovalis of the interatrial septum. Histologically, myxomas are formed of polygonal myxoma cells and small capillary canals within a mucopolysaccharide-rich matrix.[3] However, the cellular structure of myxomas is still a matter of debate.[3] Histopathological tests of the tumor mass excised from our case demonstrated that the tumor had myxoma characteristics. Cardiac myxomas are the primary cardiac tumors with the most variable clinical symptoms; their major symptoms include complications associated with embolization, and hemodynamic complications secondary to the obstruction of blood flow into the chambers of the heart. Non-cardiac symptoms that are not associated with the tumor localization include constitutional symptoms such fever, weight loss, myalgia, erythematous eruptions and lethargy.[1,5] No embolic lesions were identified in our patient, and complaints of tachycardia and dyspnea were also absent. However, the patient had complaints of short-duration syncope attacks, myalgia, and erythematous eruptions. Auscultation of left atrial myxoma generally reveals auscultatory findings of mitral stenosis, which includes hardened primary heart sounds and diastolic murmur in the apical region.[3] Auscultatory findings indicative of mitral stenosis were also identified in our patient. The electrocardiography findings of myxomas are nonspecific, while telecardiography can identify and show generalized cardiomegaly, enlargement in the relevant heart chambers, and pulmonary congestion findings associated with pulmonary venous hypertension in the lungs.[3] Our patient had a sinus rhythm, as well as increased vascularity in the lungs. Nowadays, echocardiography represents the most commonly used and accurate diagnostic tool for the diagnosis of cardiac myxomas. Transesophageal echocardiography provides very useful information, and is superior to transthoracic echocardiography in fully demonstrating the relationship between the tumor and the cavity wall, and also for the planning of surgery. Our patient underwent two surgeries under general anesthesia on two different occasions within the past seven years; however, her cardiac myxoma was not identified during the preoperative physical examinations and preparatory tests that were performed for these surgeries. Definite diagnosis in our patient was established with the aid of transesophageal echocardiography performed after transthoracic echocardiography. In addition, as our patient was of advanced age, coronary angiography was also performed to evaluate the coronary arteries. Myxomas are generally benign and rarely metastasize. No metastasis was identified in any of the tests performed on our case.[5] Myxomas that are not treated with surgery generally have poor mid and long-term prognosis. They must hence be removed after diagnosis, even if they are asymptomatic. Surgical excision is the most effective treatment method, and provides fairly good outcomes.[5,6] In our case with left atrial myxoma, the tumor was excised as a whole and without fragmenting, along with the endocardial portion to which the tumor was attached, by using biatrial incision. The resulting atrial septal defect was closed with a patch. No mortality developed in our case. Our case was followed for a period of 12 months with periodic echocardiographies, and no recurrence of myxoma was identified during this period.

Conclusion
Cardiac myxomas are generally benign tumors, with surgical excision being the most effective method for their treatment. The majority of patients who are operated following diagnosis with echocardiography continue their lives asymptotically. Against the possibility of recurrence, patients should undergo periodic echocardiography controls at certain time intervals.

Competing interests
The authors declare that they have no competing interests.

References