A Case Report of the Recurrence of a Rare Cardiac Low-Grade Fibromyxoid Sarcoma and Mortality

Nadir bir Düşük-grade Fibromiksoyid Sarkom Rekurrensi ve Mortalitesine Ait Vaka Sunumu

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ABSTRACT

As the majority of cardiac tumors are benign, malignant tumors may be easily missed. The case presented here was a female with complaints of dyspnea who was diagnosed with a left atrial mass of regular shape. After surgical excision with a pre-diagnosis of hemangioendothelioma in the frozen-section, the operation finished with mitral valve replacement. The pathological diagnosis from the immunohistochemical examination was of low-grade fibromyxoid-sarcoma. At three months postoperatively, the patient was re-hospitalized with dyspnea. Echocardiography detected a giant mass in the left atrium, 45 x 24 mm in size. The hemodynamic collapse resulted in mortality. The idea that cardiac tumors are more frequently benign causes malignant masses to be overlooked. With the perspective of an internal medicine specialist, the characteristics of each cardiac tumor should be evaluated with both clinical and imaging methods before the cardiac surgery.

Keywords: Cardiac tumor; fibromyxoid sarcoma; myxofibrosarcoma.

ÖZET

Kardiyak tümörlerin çoğunun iyi huylu olması nedenli, malign tümörler gözden kaçabilmektedir. Bu yazida, dispne yakınıması ve sol atrofımda düzgün şekilli kitle tanısı olan, bir bayan hastanın sunumu yapılmıştır. Cerrahi eksizyondan sonra, frozen-kesit bakıda, ön tanı olarak hemangiyöendotelyoma tanısı olan hastanın, operasyonu mitral kapak replasmanı ile sonlandırılmıştır. Takibinde patolojik tanı olarak immunohistokimyasal bakı sonrası düşük-grade fibromiksoyid sarkom, operasyonandan üç ay sonra tekrar dispне ile hastaneye sevk edilmiştir. Ekokardiyografide aynı bölgede, tüm atrofımda dolduran, 45 x 24 mm boyutlu dev kitle saptanmıştır. Hasta takibinde hıvomiksoyid karsa ve operasyonandan üç ay sonra tekrar dispне ile hastaneye sevk edilmiştir. Veriler normalde büyük hücreli bir hücreli kitle olan sarkom için, vaka sunumu bazık hücreli malign olma lehine olarak değerlendirilmiştir. Kardiyak tümörlerin daha sık iyi huylu olduğu fikri, malign kitlelerin gözden kaçmasına neden olabilir. Vaka sunumu ve vaka sunumunda plak hücreli bir kimyasal, dev hücreli bir formu olan sarkom için, malign olma lehine olarak değerlendirilmiştir.

Anahtar sözcükler: Kardiyak tümör; fibromyxoid sarkom; miksofibrosarkom.

The incidence of cardiac tumors has been reported as 0.02% in a large autopsy series. [1] The incidence of primary cardiac tumors is 1/1000 at autopsy and 1.5/1000 on echocardiography, but general population data are not well-known. [1] It has been stated that 75% of cardiac tumors are benign, and 25% are malignant forms. Metastatic tumors of the heart are 30 times more common than primary tumors. [1] The case presented here is a patient with a rare sarcoma of the heart, a low-grade fibromyxoid sarcoma (LGFMS), which resulted in mortality four months after the pathological diagnosis.
Case Report

A 72-year-old female presented at the cardiology with dyspnea. A systolic murmur was detected in the mitral focus on physical examination. Transthoracic-echocardiography (TTE) revealed a 4.5 x 2.4 cm (Fig. 1a) regularly-shaped tumoral mass, on the left atrial side of the posterior mitral valve (Fig. 1b). This tumoral mass caused a moderate mitral stenosis with a 1.4 cm² valve area (Fig. 1c). Clinical pre-diagnosis was myxoma and surgical excision was decided after cardiovascular surgery, cardiology and anesthesia (heart team) consultation. The tumoral mass was extracted together with the mitral valve because of valve invasion of the tumor. With the pre-diagnosis of hemangioendothelioma in the frozen-section, the operation was successfully finished with mitral valve replacement (Fig. 1d). Coumadin and 100 mg acetylsalicylic acid treatment were started with effective INR value. All operation material was sent to the pathology department. The tumor was seen to continue through the surgical margins on the paraffin sections. Immunohistochemical examination showed that the tumor cells were immunopositive for vimentin, focal positive for smooth-muscle actin, CD34, CD31, and immune negative for epithelial membrane antigen, CD68, myogenic-differentiation-1, desmin. The pathological diagnosis was made of LGFMS (Fig. 2). Computed tomography (CT) was applied to the patient, but no other tumor was observed. The patient was referred to oncology after this diagnosis. Chemotherapy or radiotherapy could not be applied due to the refusal of treatment of the patient and relatives. At third-months postoperatively, the patient presented at the Emergency Department, again suffering from dyspnea. The patient was admitted to the intensive care unit, and a diuretic infusion was started. The TTE and trans-esophageal-echocardiography (TEE) were performed, and a giant mass with cystic areas was observed to fill the left atrium completely and was disrupting valve movements (Fig. 3). After consultation with the cardiac team on the basis of the new findings, it was decided to re-operate when hemodynamic stabilization was obtained. On the 10th day of hospitalization, the patient was exitus due to hypotension despite inotrope infusion.

Figure 1. Dimensions are measured as 45 x 24mm (a); regularly-shaped tumoral mass is on the left atrial side of the posterior mitral valve (b); there is a moderate mitral stenosis with a 1.4 cm² valve are due to tumor (c); the echocardiographic view of a bioprosthetic mitral valve (d).

Figure 2. Histopathological and immunohistological preparations.
Discussion

Primary cardiac tumors are rare, with 0.3%-0.7% incidence of all recorded cardiac tumors. Metastatic tumors are 30 times more common.[2] Of the primary tumors, 2/3 is benign, and myxoma is the most common type. The remaining third of primary tumors are malignant, of which 2/3 are sarcomas.[1] The most common sarcoma of the heart is angiosarcoma, and primary cardiac myxofibrosarcoma is much rarer. In 2012, Wang et al.[3] reported a case of a 42-year-old male, and only 17 cases were found since 1963 in their literature research. This tumor, which has been described in recent years, is mostly seen in young adults and is mostly located in deep soft tissues.[4] In contrast, the current case was middle-aged, and the mass was located in the heart, thereby making this case one of the known rare patients.

Macroscopically, the tumor is well-circumscribed and has a cross-sectional face that can vary between a fibrous and myxoid appearance. Microscopically the tumor surrounds a pseudo-capsule. However, focal infiltration can be detected in surrounding soft tissues. The tumor consists of smooth, fibroblastic spindle cells that show a fibromyxoid growth-pattern.[5] It may be confused with many other soft tissue tumors.[6] In accordance with these data, the pre-diagnosis in the current case was hemangioendothelioma on the frozen-section, and the surgical section borders were not sufficient for total tumoral resection.

Low-grade fibromyxoid sarcoma is detected predominantly in the left atrium, and less frequently, the tumor can invade the mitral valve.[3] In the current case, the tumor was also on the left atrial side of the posterior mitral valve. Dyspnea is the most common symptom on presentation, as was seen in the current patient.

The main treatment is as much surgical resection of the tumor as possible, obtaining a negative limit of the surgical margin. Curative resection and adjuvant radiotherapy are recommended, but the long-term efficacy is not well known.[7] Also, Reardon et al.[8] reported a 30-day mortality rate of 9%, with survival on the patient that had negative surgical margins.

As LGFMS is known to have high mortality and the late diagnosis of tumoral malignancy in the current case, rendered total resection of the tumor on re-operation technically challenging due to the functional importance of the heart and the difficulty of surgical access.

Early recurrence of LGFMS is not expected. According to this data, Wang et al.[9] presented an LGFMS with a patient who developed a bone metastasis two years later and is still alive 26 months after the cardiac surgery. In our patient, recurrence in the same region was probably due to incomplete excision due to perioperative benign diagnosis and refusal of chemo-radiotherapy of the patient.

Conclusion

Due to the rare frequency of malignant cardiac tumors, cardiologists and cardiovascular surgeons have little experience in the clinical evaluation of cardiac malignancies. The idea that cardiac tumors are more frequently benign may cause malignant masses to be missed.

With the perspective of an internal medicine specialist, the characteristics of each cardiac tumor should be evaluated with both clinical and imaging methods, also if possible with transcatheter biopsy (especially for right heart originated tumors), before cardiac surgery. Furthermore, perioperative frozen-section evaluation by experienced pathologists may change the prognosis with a true diagnosis.

Disclosures

Informed consent: Written informed consent was obtained from the patient for the publication of the case report and the accompanying images.

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Conflict of Interest: None declared.

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