Multimodality imaging of a recurrent case of right-sided cardiac leiomyosarcoma with an unusual clinical course

Sıradışı klinik seyirli tekrarlayan sağ tarafı kardiyak leomiyosarkomlu olgunun çok yöntemli görüntülenmesi

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Summary—Primary cardiac sarcoma is an uncommon entity and is related with poor long-term survival. Presented here is an unusual case of right atrial leiomyosarcoma which almost completely occluded both the right atrium and ventricle and destroyed the tricuspid valve. Multimodality imaging was used to characterize the mass and guide surgical management. A combination of two radical surgical excisions and both radiation and chemotherapy resulted in a better long-term outcome. The patient was one of the longest survivors.

Primary cardiac sarcomas are extremely rare, and the few reported cases have been treated surgically.[1] Leiomyosarcoma is a highly aggressive and locally invasive tumor with a mean survival time of 6 months in untreated patients.[2]

We presented a case of a pathologically proven right-sided leiomyosarcoma associated with subtotal occlusion of both right atrial and ventricular cavities and complete destruction of the tricuspid valve. The longest ever period of remission was observed after aggressive surgical resection and subsequent adjuvant radiochemotherapy.

CASE REPORT

A 40-year-old woman presented to our clinic with 1-month history of progressively worsening dyspnea at rest. Physical examination revealed a rough diastolic murmur in the third left intercostal space at the left sternal border on auscultation. Her electrocardiography and chest radiography were unremarkable. Two-dimensional transthoracic echocardiography (2D TTE) with Doppler imaging revealed a mass causing significant right ventricular inflow tract obstruction. Two-dimensional transesophageal echocardiography confirmed the presence of a multilobular mass arising from the interatrial septum. The mass filled up the right atrial cavity, extended through the right ventricular cavity and was causing obstruction of the tricuspid valve (Figure 1a). Real-time 3-dimensional TEE view provided exact spatial information about the shape and contour of the intracardiac mass (Figure 1b, Video*). Computed tomography (CT) scan did not show any mass in the extracardiac area. Cardiac magnetic resonance imaging (MRI) with gadolinium contrast revealed a mass measuring 9x6x3 cm exhib-
iting a broad base of attachment and intramural location (Figure 1c and d).

A cauliflower-like mass within the right atrium (RA) and extending to the right ventricle (RV) was observed during surgery and completely removed (Figure 2a). The histopathological study revealed a spindle cell neoplasm, consistent with leiomyosarcoma, invading the underlying myocardium (Figure 2b). The patient was discharged home on the 17th postoperative day in a stable condition with planning for anthracycline-based chemotherapy and radiation. Follow-up serial TEE studies at 6, 12, 18, 24, 30 and 36 months showed no residual RA and RV mass, but there was severe tricuspid regurgitation.

After a 42-month period of uneventful clinical and laboratory follow-up, the patient was readmitted with dyspnea. 2D TTE and MRI revealed a relapsed tumor in the RA, while CT scanning did not show any

Figure 1. (A) Transesophageal echocardiography revealed a right atrial tumor extending from the septum through the tricuspid valve into the right ventricle. (B) Real-time 3D transesophageal echocardiographic view of the right atrium provided exact spatial information about the shape and contour of the intracardiac mass. Cardiac MRI showed the intramural extension of the tumour by bright blood (C) and black blood MRI technique (D). Imaging of the relapsed tumor with TTE (E) and MRI (F).
extracardiac mass (Figure 1e and f). The tumor was subtotally resected and the patient underwent chemotherapy. Histopathological examination of the tumor showed a leiomyosarcoma. The patient eventually died due to RV heart failure.

**DISCUSSION**

Cardiac leiomyosarcoma is a rare entity, with 76% of tumors detected in the left atrium, 16% in the right atrium, and 8% diffusely in the ventricles.[3] Primary leiomyosarcomas are highly aggressive and infiltrate adjacent structures. Although most patients die within 1 year of diagnosis, Glaoui et al. have reported an 8-year survival duration after combined surgical and adjuvant radiochemotherapy.[4]

While echocardiography is currently a widely-used method for diagnosing cardiac masses, differential diagnosis of cardiac tumors often presents a challenge for the sonographer. When malignancy is suspected, MRI and CT remain essential for operative planning and identification of tumor extension. As demonstrated in our case, the use of gadolinium increases the conspicuity of the tumor by showing different contrast enhancement with respect to the surrounding normal myocardium. Furthermore, it permits differentiation of thrombus from tumor, as thrombus would not be enhanced.[5]

In our case report, the site of origin of the tumor was the right atrium, which is rare. The presentation was consistent with thromboembolic events frequently associated with this tumor site. Other possible presentations include progressive or sudden right-sided cardiac failure on the account of tricuspid valve blockage by the tumor, or paroxysmal atrial arrhythmias. Echocardiographic imaging is the most sensitive imaging technique, with the ability to identify tumors as small as 3 mm. The spatial relationship between the mass and adjacent structures could be seen clearly in preoperative 2D and 3D echocardiography, but tissue characterization could not be demonstrated. The primary contribution of MRI in our case was to delineate the anatomical extension of tumor invasion. Inadequately evaluated by echocardiography, the extension of the tumor underneath the myocardium and the broad base of attachment were and were completely assessed by MRI.[6]

A malignant neoplasm was suspected on the basis of MRI findings and a radical excision was planned. The combination of two radical surgical excisions and both radiation and chemotherapy resulted in a relatively better long-term outcome. To our knowledge, our patient was one of the longest survivors in the literature.

We conclude that the use of echocardiography in conjunction with MRI can provide very valuable data in evaluation of intracardiac masses. MRI may differentiate between malign and benign tumors and provide better anatomical information for preoperative planning. We suggest that cardiac MRI should be performed in all cases of suspected cardiac tumor unless contraindicated.
This report suggests that multimodality imaging and treatment protocol can improve survival in patients with leiomyosarcoma.

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*Supplementary video file associated with this article can be found in the online version of the journal.

REFERENCES


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