Primary Pulmonary Fibrosarcoma in Giant Mass Appearance: A Rare Case Report

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INTRODUCTION

The primary sarcomas of the lung constitute 0.5% of all pulmonary malignancies.[1] Fibrosarcomas, which are a subtype of sarcomas and originate from the mesenchymal cells, may emerge in any part of the body and may be encountered in any age group.[2] Because they are reported as rare case presentations in the literature, there is no definitive treatment protocol.

Primary pulmonary fibrosarcomas are usually low-grade tumors, but have a high recurrence rate.[3] The survival rate is better in tumors ≤5 cm compared with tumors >5 cm.[4] Early diagnosis is essential to improve survival, and a complete resection with sufficient surgical margins should be carried out.[5]

CASE REPORT

Respiratory sounds could not be detected in the upper and middle zones of the right lung during the physical examination of the 56-year-old female patient, applied with the complaint of dyspnea, due to the suspected malignancy. We carried out intrapericardial bilobectomy superior and the final pathological examination showed primary pulmonary fibrosarcoma. Primary pulmonary fibrosarcoma is a very rare type of tumor and surgery is the main choice of treatment. We present this case to contribute to the literature.

ABSTRACT

We planned surgery a fifty six year-old female patient, who applied with the complaint of dyspnea, due to the suspected malignancy. We carried out intrapericardial bilobectomy superior and the final pathological examination showed primary pulmonary fibrosarcoma. Primary pulmonary fibrosarcoma is a very rare type of tumor and surgery is the main choice of treatment. We present this case to contribute to the literature.
hanced cranial magnetic resonance imaging, which was carried out during the preoperative period, did not show any pathological findings.

We performed a right posterolateral thoracotomy and observed a soft-structured mass lesion, which completely filled the upper and middle lobes. We collected samples from the lesion, and a frozen-section analysis was performed. The result of the frozen-section examination was interpreted in favor of malignant mesenchymal tumor consisting of fusiform and epithelioid cells. As the mass lesion involved the vein of the upper lobe, intrapericardial bilobectomy superior and mediastinal lymph node dissection was carried out.

In the final pathology report, the macroscopic dimension of the tumor was 14×7×6 cm. The histopathological examination showed a mesenchymal tumor consisting of monomorphic spindle cells. The stroma contained variable collagen ranging from a sensitive intercellular network to regions with few cells. Immunohistochemically, tumor cells displayed positive staining with vimentin and CD34. No tumor tissue was determined in the surgical margins of bronchus-vessel-parenchyma. Tumor tissue was determined in the pericardial surgical margin. The lymph nodes 3A-7-8-9-10 were considered anthracotic.

No morbidity and mortality were encountered in the postoperative period. The drain was removed on the 8th day, and the patient was discharged with recovery on the 10th day. Radiotherapy was initiated in the postoperative period (Figs. 1–3).

**Figure 1.** Preoperative radiological evaluation of the patient: (a) Chest X-ray, (b) axial images in computed tomography, and (c) axial, coronal, and sagittal images in positron emission tomography.
**Figure 2.** Perioperative images of the patient and pictures of the extracted piece and postoperative PA - lung X-rays. Single arrow: giant mass located in the upper lobe and middle lobe; double arrow: lower lobe.

**Figure 3.** Sections from the patient’s pathological specimen. (a) Histological examination revealed spindle-shaped cells in the fibrous matrix. (b) The stroma has a delicate intercellular network. (c) The stroma has particular areas with diffuse sclerosis or hyalinization. (d) Diffuse CD34 positivity was observed in tumor cells.
Radiotherapy in the postoperative period. Our case is cur-
pericardial surgical margin was positive, our case received
the largest dimension of the tumor was also 14 cm. As the
recently in the 3rd month of the follow-up, and we have not
worse in the soft tissue sarcomas of the extremities.[11] The survival rate of primary pulmonary sarcomas is usually
matoid sarcoma was excluded as epithelial differentiation
ovial sarcoma with the negative TLE1 staining, and sarco-
pulmonary fibrosarcoma in our case, the solitary fibrous
margins are positive and complete resection is not possi-
to improve survival, and a complete resection with suffi-
cases did not originate from bronchi. In the study by Mal-
ancy. As only a limited number of cases are reported in
neuroblastoma, rhabdomyosarcoma. When diagnosing primary
the lung constitute 0.5% of all pulmonary malignancies.[3] Primary bronchopulmonary
fibrosarcoma is the second most common intrathoracic sarcoma after leiomyosarcoma. Despite this fact, less than
100 cases have been reported worldwide.[9]

The intrathoracic fibrosarcomas tend to emerge as endo-
bronchial masses. Lobar bronchi are involved in children and adolescents, while solitary and multinodular or mass
lesions are usually seen in the lungs of adults.[9] When the intrapulmonary fibrosarcomas develop as small endobronchial
lesions, they cause cough and hemoptysis. On the other
hand, intraparenchymal lesions usually do not cause symptoms.[2] In our case, there was dyspnea depending on
the tumor compression, and this complaint was relieved
immediately after surgery. As the tumor was located in the intra-
parenchymal area in our case, we believe that it remained
symptomatic until its size grew to cause compression.

The most common histological subtypes of lung sarcomas
are synovial sarcoma, epithelioid hemangioendothelioma, leiomyosarcoma, and malignant peripheral nerve sheath,
which are followed by pleomorphic sarcoma, liposarco-
a, and rhabdomyosarcoma.[10] When diagnosing primary
pulmonary fibrosarcoma in our case, the solitary fibrous
tumor was excluded with negative STAT6 staining; and syn-
ovo sarcoma with the negative TLE1 staining, and sarco-
matoid sarcoma was excluded as epithelial differentiation
was not determined with multiple cytokeratins.

The survival rate of primary pulmonary sarcomas is usually
worse in the soft tissue sarcomas of the extremities.[11] Endobronchial tumors can be detected earlier, and thus they have a better prognosis.[11] Primary pulmonary fibro-
sarcomas are usually low-grade tumors but have a high
recurrence rate.[9] Early diagnosis of a tumor is essential
to improve survival, and a complete resection with suffi-
cient surgical margins should be performed. If the surgical
margins are positive and complete resection is not possible,
postoperative radiotherapy should be implemented.

The largest primary pulmonary fibrosarcoma reported in
the literature had a size of 14 cm.[13] In our case, the
largest dimension of the tumor was also 14 cm. As the pericardial surgical margin was positive, our case received
radiotherapy in the postoperative period. Our case is cur-
rently in the 3rd month of the follow-up, and we have not
yet observed recurrence or mortality.

CONCLUSION

Primary pulmonary fibrosarcoma is a rare pulmonary ma-
lignancy. As only a limited number of cases are reported in
the literature, a definitive treatment protocol is not avail-
able, but surgery is considered the main treatment choice.
We presented this case to contribute to the literature.

Informed Consent

Written informed consent was obtained from the patient
for the publication of this case report and the accompany-
ing images.

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CONCLUSION

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able, but surgery is considered the main treatment choice.
We presented this case to contribute to the literature.
Nefes darlığı şikayeti ile başvuru alan elli altı yaşında kadın hastayı malignite şüphesi ile ameliyat etmeyi planladık. İntraperikardiyal bilobektomi superior yaptığımız hastanın nihai patolojisi primer akciğer fibrosarkomu olarak raporlandı. Primer akciğer fibrosarkomu oldukça nadir görülen bir tümör olup tedavisinde cerrahi esas rolü oynamaktadır. Olgumuz literatüre katkı amacıyla sunulmaktadır.

Anahtar Sözcükler: Akciğer fibrosarkomu; primer fibrosarkom; sarkom.