

# Sarcomatoid Carcinoma of the Lung – a Case Report

## Sarkomatoidní karcinom plic – kazuistika

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### Summary

**Background:** Sarcomatoid carcinoma (SARC) of the lung is a very rare and aggressive type of non-small cell lung cancer. It belongs to a group of poorly differentiated carcinomas with partial sarcomatoid differentiation or with a direct sarcoma component. Characteristic findings include a large tumor with an invasive tendency, early recurrence and systemic metastases. **Case:** The authors present a case of SARC in the 77-year-old patient. Preoperative staging confirmed sarcomatoid carcinoma of the lower lobe of the left lung without generalization on PET/CT. However, an infiltration of more than 2/3 of the diaphragm was ascertained. A resection was performed – a left lower lobectomy with resection of the diaphragm and its replacement by a muscle flap made from the latissimus dorsi muscle with vascular pedicle. Histological findings confirmed the diagnosis of sarcomatoid (pleomorphic) carcinoma pT3N0M0. The patient underwent adjuvant chemotherapy; recurrence and systemic dissemination of the disease occurred after 20 months; the patient died 21 months after the surgery.

### Key words

lung cancer – neoplasm invasion – surgical flaps – diaphragm – sarcomatoid carcinoma

### Souhrn

**Východiska:** Sarkomatoidní karcinom (sarcomatoid carcinoma – SARC) plic patří mezi velmi vzácné a agresivní typy nemalobuněčného karcinomu plic. Jedná se o skupinu nízce diferencovaných karcinomů s částečnou sarkomatoidní diferenciací nebo přímo s komponentou sarkomu. Charakteristickým nálezem bývá objemný nádor se sklonem k invazi, brzká rekurence a systémové metastazování. **Případ:** Autoři prezentují případ SARC u 77leté pacientky. Předoperační staging prokázal sarkomatoidní karcinom dolníhó laloku levé plíce bez generalizace na PET CT, ale s infiltrací více než 2/3 bránice. Byla provedena resekce – dolní lobektomie s resekci bránice a náhradou svalovým lalokem z m. latissimus dorsi na cévní stopce. Histologický nález potvrdil diagnózu sarkomatoidního (pleomorfního) karcinomu pT3N0M0. Pacientka prodělala adjuvantní chemoterapii, za 20 měsíců došlo k relapsu a systémové diseminaci nádoru, pacientka zemřela 21 měsíců od operace.

### Klíčová slova

karcinom plic – nádorová invaze – svalový lalok – bránice – sarkomatoidní karcinom

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### Introduction

The diagnosis of sarcomatoid carcinoma (SARC) of the lung includes a group of poorly differentiated non-small cell lung tumors, which include a component with sarcomatoid differentiation (spindle-cell and/or giant cell population) or a sarcoma component. The incidence is only 0.1–1.3% of all malignant lung tumors [1]. They affect more frequently males than females (2–10 : 1), primarily smokers; the average age at diagnosis is 60 years [2]. SARC is characterized by rapid growth, invasion, disease recurrence and metastases.

### Case presentation

The 77-year-old patient was examined due to shortness of breath, chest pain on the left side of the chest and dry cough. X-ray and CT findings showed a left-sided pleural effusion and a large heterogeneous tumor measuring 110 × 95 × 95 mm. Endobronchial findings were with no direct signs of

the tumor; cytology findings from brush biopsy were suspected to be malignant. A transparietal biopsy under CT was performed and diagnosed sarcomatoid granuloma. PET CT confirmed an extensive tumor arising from the lower lobe of the left lung with infiltration of approximately 2/3 of the diaphragm, with no signs of invasion into the abdominal cavity or pericardium and with no signs of dissemination of the disease (Fig. 1). Laboratory findings showed only slight elevation of CA125, while other markers were with no pathological values. Primary staging was concluded to be cT3N0M0. The patient was indicated for surgical resection, which, due to the extent of the tumor, was performed through a double thoracotomy in the 5<sup>th</sup> and 8<sup>th</sup> intercostal space. The resection included a left-sided lower lobectomy and resection of 3/4 of the surface of the diaphragm. The extensive defect resulting from the diaphragm resection was reconstructed using a muscle flap made

from the latissimus dorsi muscle with vascular pedicle (Fig. 2). No serious complications were observed in the postoperative course (Fig. 3). The patient was discharged on the 14<sup>th</sup> postoperative day.

Histological findings confirmed sarcomatoid cancer – pleomorphic subtype pT3N0MX, immunohistochemically positive vimentin 100%, TTF1 5–20%, AE1/AE3 80%, only sporadic positivity of CK7. The patient subsequently underwent adjuvant chemotherapy with carboplatin and gemcitabine. PET CT performed 11 months after the resection showed the disease recurrence. Nine months later the patient complained of back pain, weight loss and weakness. Follow-up PET CT revealed disease recurrence with generalization. The patient died 21 months after the operation.

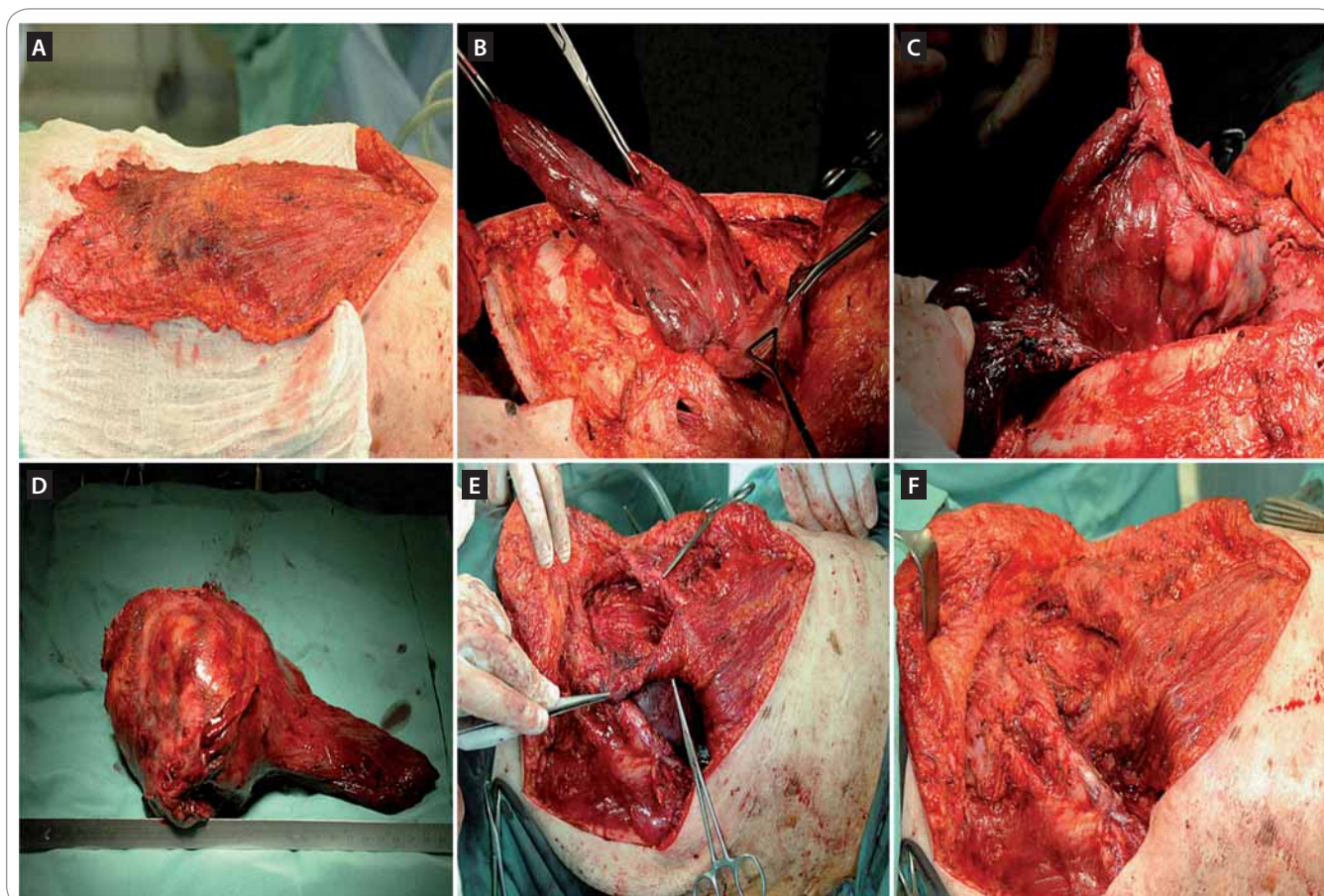
### Discussion

The histogenesis of SARC remained unclear for a long time due to its histo-



Fig. 1. CT images of the tumor before the surgery. The tumor arises from the lower left lung lobe and invades the diaphragm.





**Fig. 2. Intraoperative pictures. A. preparation of the muscular flap; B, C. resection of the tumor with the diaphragm; D. the resected tumor; E, F. the diaphragm repair with the muscular flap.**

logical heterogeneity. Years ago led this to the proposal of numerous theories. Only with the help of modern molecular methods could be the origin of sarcomatoid differentiation partially revealed. Currently, support for the so-called totipotent hypothesis prevails, which assumes the origin of the tumor cell to be from stem cells, which can differentiate into an individual epithelial or mesenchymal component [3]. This theory was also supported by the convincing determination of monoclonality and clonal identity of cell lines of carcinoma and sarcoma [4].

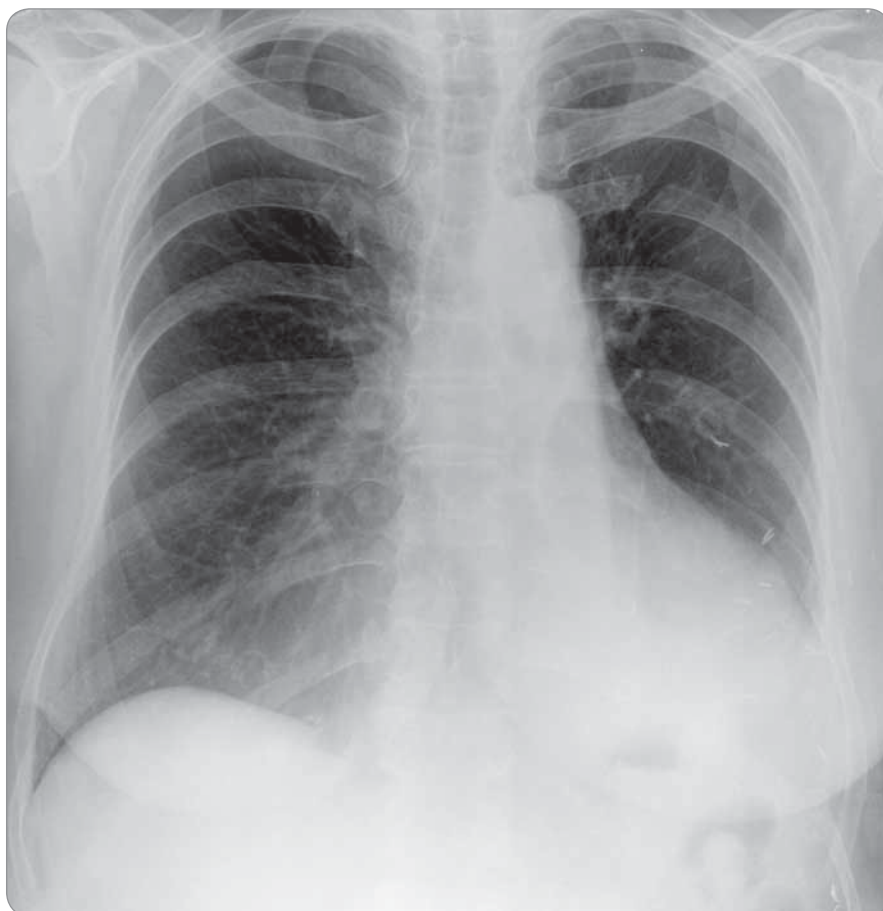
In an earlier WHO classification, individual types of SARC were included among conventional non-small cell cancers. Since 2004, the WHO classification has remained stable and the SARC group contains five subgroups – pleomorphic carcinoma, spindle cell carcinoma, giant cell carcinoma, carcinosarcoma and pulmonary blastoma [5].

Histopathological diagnosis of SARC and its subtypes may cause certain problems. Due to difficulties with capturing all the possible components, which may be within a tumor represented only in a minority, the tumor should be processed for histological analysis extensively. Certain subtypes may be identified by means of basic staining with hematoxylin-eosine; however, usually other methods must be used for specification. Immunohistochemical analysis must show cytokeratin positivity (most often AE1/AE3, CAM5.2, CK18, CK7). Vimentin, CEA and smooth-muscle markers are also positive. Positivity of epithelial markers is not essential if components of conventional cancer are morphologically present [5].

To date, no prospective studies regarding SARC are available. This is primarily due to the low incidence of this type of carcinoma, and also due to more complicated diagnostics, when in

a high number of cases it is recognized as late as according to the postoperative histological findings. Furthermore, due to tumor aggressiveness, majority of patients are SARC diagnosed in advanced stages or even upon autopsy. To date, only retrospective studies with mainly small numbers of patients with SARC have been documented, and these patients are usually compared to the groups of patients with conventional non-small cell cancer. These studies differ somewhat in their results. In most of these studies, patients with SARC had significantly worse results in terms of median of survival, disease progression as well as overall survival [6,7]. On the other hand, some studies did not find any difference in survival between the two aforementioned groups of patients [8,9].

Nonetheless, SARC diagnosis is generally considered to be a disease with a significantly poorer prognosis in terms



**Fig. 3. Postoperative X-ray three months after the surgery.**

of survival and early disease relapse. A complete resection in the early stages of disease brings the best results; however, five-year survival in stage I is achieved by only 37% of patients [10].

SARC has a low sensitivity to radiochemotherapy and improved responses to this modality have not been recorded, nonetheless it is recommended especially for bulky disease tumors, tumors invad-

ing the thoracic wall or in cases of metastatic lymph node involvement. This is due to the higher incidence of loco-regional and distant disease recurrence. In advanced stages, where surgical treatment is not indicated, palliative oncological therapy is recommended [7].

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