ASKIN'S TUMOR
– A RARE TUMOR OF THE THORACOPULMONARY REGION

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SUMMARY – Askin’s tumor is a malignant small round cell tumor affecting the thoracopulmonary region. Because of its neuroectodermal origin it is also known as primitive neuroectodermal tumor. This newly seen and aggressive malignant tumor most frequently occurs in children and young adults. The symptoms and clinical presentation as well as radiomorphologic abnormalities are nonspecific. Therefore, establishing an accurate diagnosis of Askin’s tumor is difficult, thus histologic and immunohistologic findings are important. Immunohistologic staining of the specific marker neuron-specific enolase is essential. In addition, Askin’s tumor shows histologic features similar to Ewing’s tumor located in the bones, which suggests it to belong to the group of Ewing’s tumors. Therapeutic approach is multimodal, e.g., surgical resection, chemothera-py, and radiation. Overall prognosis is poor. The objective of this report is to point to the unusual tumor of the thoracopulmonary region. A 25-year-old female with a giant right chest tumor mass and immunohistologically characteristic Askin’s tumor is described.

Key words: Lung neoplasms—pathology; Lung neoplasms—histology; Lung neoplasms—diagnosis; Case report

Introduction

Askin’s tumor is a very rare, highly malignant tumor of the thoracopulmonary region1-3. It was first described by Askin et al. in 1979. They suggested its neuroectodermal origin based on the morphology and ultrastructural changes. Later it was confirmed by Linsinella et al. using immunohistochemical staining for neuron-specific enolase (NSE). Histologic features of Askin’s tumor include small round cells, which are often NSE-positive1,3,4. The term primitive neuroectodermal tumor (PNET) is synonymous, however, it is usually used for a localization outside the thoracopulmonary region. The usual and unusual PNET localizations have been identified2,5,6. The clinical presentation and radiologic appearance of the lesions are nonspecific, which adds to the importance of the histologic and immunohistochemical diagnosis. Due to its histologic similarities to classic Ewing’s sarcoma, Askin’s tumor is also referred to as extraskeletal Ewing’s sarcoma1,7, and it belongs to the Ewing’s family of tumors1,8,9. In contrast to Askin’s tumors, Ewing’s sarcoma is immunohistochemically characterized by negativity for NSE immunostaining. Genetic analysis, electron microscopy and immunohistochemical studies have allowed for more reliable identification of the origin and classification of this tumor10, which is usually observed in children and young adults.

Case report

A 23-year-old female was admitted to the hospital for generalized weakness, sweating, recurrent fever to 38 °C, occasional chills and shaking, dry cough, intermittent attacks of suffocation and dyspnea. The patient had experienced pain on and off in the right side of her chest as well as in her thoracic and lumbar spinal bone but the pain was transitory. The onset of the discomforts was three months prior to hospital admission. Antimicrobial therapy failed to
provide any improvement. Apart from childhood pneumonia and surgery for treatment of appendicitis, the patient’s medical history included no other diseases. She occasionally took nonsteroidal antirheumatics for pain.

The patient was of average osteomuscular constitution and nutritional status, with somewhat pale skin, lucid, oriented, febrile (up to 38 °C), with no peripheral lymphadenopathy. In her head and neck status no abnormality was detected, and the chest wall was normal. On auscultation normal respiration was heard, except above the upper half of the right pulmonary lobe, where it was found to be weakened. Heart action was rhythmic, sounds were clear, and no murmur was detected. Blood pressure was 110/70 mm Hg. The abdomen was tender and painless with no organomegaly, and no extremity abnormalities were detected.

Biochemical findings: erythrocyte sedimentation rate 84; WBC 8.8x10^9/L (neutrophils 66.8%, lymphocytes 17%, monocytes 15.2%, eosinophils 0.6%); RBC 3.8x10^12/L; hemoglobin 10.9 g/L; hematocrit 32%; platelets 278x10^3/L; total protein 59.00 g/L (albumins 45.4%, globulins: alpha 1 8.7%, alpha 2 19.3%, beta 12.1%, gamma 14.5%); lactate dehydrogenase 1803 U/L (reference range 0-460 U/L), and fibrinogen 16.6 g/L (reference range 1.7-3.9 g/L). The results of all other biochemical blood tests were within the normal values. The urinalysis findings complied with the reference values.

Summation chest radiography and native tomography showed in the upper half of the right hemithorax laterally and dorsally along the thoracic wall a giant tumor mass of polymorphic characteristics, 14 cm in size, reaching the hilus and even compressing and dislocating the bronchi. The other tumefaction in the region measuring approximately 4 cm in diameter was localized medially in the retroclaveicular area. No dislocation of the mediastinum was found. On the left side, no abnormalities were detected. The heart was found to be of an appropriate shape and size (Fig. 1). Chest CT scans revealed two tumor masses: one of a polymorphic structure, 10x12 cm in size, in the upper half of the right hemithorax (Fig. 2), penetrating into the intercostal space (Fig. 3), and one smaller mass, 3 cm in diameter, that was located paravertebrally. Upper abdominal CT scans revealed no focal pathologic lesions of either the spleen or the liver. Abdominal ultrasonography showed

![Figure 1. A giant right chest tumor mass spreading from hilus to chest wall](image1)

![Figure 2. Polymorphic structure of the tumor](image2)

![Figure 3. Extension of the tumor into intercostal spaces](image3)
no metastatic changes. On the radiographs of the lumbar spine in two projections and of the pelvis and hips no metastatic changes were detected.

Electrocardiography revealed sinus rhythm of 97/min, semivertical electrical axis, and nonspecific repolarization abnormalities. Spirometry identified minor restrictive ventilatory defects. Diffusion capacity for carbon monoxide was moderately impaired. Arterial blood gas analysis determined partial respiratory insufficiency with hyperventilation.

Cytology of the transthoracic biopsy specimen obtained from the right intrathoracic mass revealed blood, lymphatic cells, and several clusters of poorly preserved cells suspect of malignancy.

The patient was operated on ten days following hospital admission. At surgery the tumor was found to infiltrate the major part of the right upper lung lobe, posteriorly paravertebral intercostal spaces, and laterally the 5th rib. Involvement of the diaphragm and parietal pleura from the 7th to the 12th rib by numerous metastatic masses was also evident. The tumor was posteriorly paravertebrally separated from the tumor infiltrated thoracic wall. In addition to right upper lung lobe and resection of a segment of the 5th rib, numerous metastatic masses were excised from the parietal pleura and diaphragm, and the mass excised from the diaphragm was referred for histopathologic analysis.

Gross pathomorphologic examination of the resected right upper lung lobe revealed that the lobe was for the largest part permeated with a grayish-white soft tumor, measuring 14.5x11x7 cm with focci of intramembraneous hemorrhage. Histologic examination showed a small cell tumor, and immunohistochemical findings positive for NSE and CD 99 revealed an Askin’s tumor. The tumor tissue identical to that of the primary tumor involved the resected section of the 5th rib as well as the adjacent connective tissue and thoracic wall musculature along with an infiltrate resected from the diaphragm. No tumor tissue was found in the mediastinal lymph node.

The postoperative course was satisfactory. Postoperatively, two masses developed in the soft tissues of the ocipital right and parietal left region of the head. Skull radiography showed no pathologic changes. Fine needle aspiration biopsy of the masses revealed poorly preserved cells of a malignant metastatic neoplasm with low-grade differentiation, and metastases of PNET were inferred. The patient was transferred to oncology ward for further chemotherapy and radiotherapy. Magnetic resonance tomography performed at that point supported osteolytic changes in the thoracic and lumbar spine with metastatic infiltration of the spinal canal. The patient received telecobalt irradiation therapy to the spine, and later to the thorax for metastatic lesions. She was also given cytostatic polychemotherapy consisting of three cycles of vincristine, adriamycin and cyclophosphamide. After the third cycle of chemotherapy the patient exhibited disease progression and development of multiple metastases, and chemotheraphy was withdrawn for a markedly deteriorated general condition. The patient died seven months after the onset of the disease symptoms.

Discussion

Askin’s tumor typically involves soft tissues of the thoracic wall and paravertebral structures. It rarely affects the pulmonary parenchyma, and it seems to most commonly arise from the intercostal nerves[1-2]. The differential diagnosis of Askin’s tumor includes Ewing’s sarcoma, thymidonyosarcoma, neuroblastoma and lymphoma, whereas some other primary or secondary tumors of the thoracic wall are less frequently considered. Askin’s tumor is occasionally observed as a secondary neoplasm, after the treatment for acute lymphoblastic leukemia and complete remission of Hodgkin’s disease[3]. For the diagnosis of Askin’s tumor, certain criteria should be met: localization in the thoracic region, typical histomorphologic characteristics, and specific immunohistochemical markers[4]. In our patient, all these preconditions were met.

Clinical manifestations are relatively nonspecific. Pain in the thoracic wall, which may additionally be deformed by the tumor mass, seems to be the most common symptom associated with Askin’s tumor. Further features would include subfebrile or high temperature, dyspnea, proproductive cough, weight loss, pleural effusion, therapy resistant pneumonia, hemoptysis, hoarseness, and Horner’s syndrome[4,11,12]. The clinical manifestations presented in our patient were also nonspecific. Incidentally discovered Askin’s tumors have also been reported in the literature[13].

Conventional radiography usually shows a mass of various size accompanied by calcifications, ossions of the ribs, pleural effusions, and even pneumothoraces[11,12]. On CT scans heterogeneity is noticed particularly in large tumors due to the areas of necrosis and hemorrhage as well as infiltrations of the thoracic wall and rib destruction[11]. In addition, small pulmonary metastases from this tumor are also detectable on CT. Magnetic resonance tomography is not necessary, and should be reserved to when an infiltration of the thoracic wall is suspected. It has proven useful
in the visualization of an extrathoracic disease. Polymorphisms of the tumor and infiltration of the thoracic wall were also observed in our patient. Histologically, Askin’s tumor is characterized by invasive growth, small round cells with a hyperchromatic nucleus, and sometimes a weak PAS-positive cytoplasm. Our patient also exhibited a cytoplasm weakly positive for PAS. The cytoplasm was scanty; the nucleus–plasma quotient was high, and multiple prominent nuclei were as well as spindled pseudosarcomatous cells were revealed. An accurate histologic diagnosis requires immunohistochemistry.NSE appears to be an important marker. Although immunohistochemistry tests may not necessarily be always positive for NSE, other neural markers such as chromogranin and synaptophysin may also confirm a neural phenotype. Some authors have observed NSE in 95%–100% of these patients. Immunohistochemically, our patient showed tumor cells diffusely positive for NSE, focally positive for vimentin, and membranous positivity for CD 99.

Late dissemination despite a local finding was described by Askin as being characteristic of this tumor. Some authors found distant metastases at the time of diagnosis in more than a half of patients. The finding of distant metastases at initial diagnosis is an exception rather than a rule. The most usual sites of dissemination include the bones, bone marrow, lungs, liver, lymph nodes and spleen, and multiple metastases are present in one third of patients. A multimodal approach, which consists of radical surgical resection, chemotherapy and irradiation is advocated for the treatment of this tumor, however, there has been no unique scheme so far. The majority of authors support resection because this tumor shows a significant propensity towards local recurrence. Complete resection appears to be superior to other modalities of surgical treatment. Multiple resections for recurring Askin’s tumors have been reported in the literature. Neoadjuvant therapy, which seems to improve resectability, is proposed. In all instances postoperative chemother-apy and radiotherapy are mandatory due to micrometastases and tendency to recurrences. There is no unique scheme for chemotherapy. Because of resemblance to Ewing’s sarcoma, the analogous therapeutic scheme is followed. More intensive therapy is advocated. In our patient, the multimodal approach to the treatment was also chosen. The major part of the primary tumor and metastases were surgically removed, and after postoperative recovery palliative irradiation telecobalt therapy in combination with cytostatic therapy was administered. The prognosis of Askin’s tumor is very poor in spite of improvements that have been achieved in therapy over the last twenty years. Variable data on the average survival rates have been reported. The 2-year survival rate is reported to be 20%–30%, and the 5-year rate is 14% to 17%. An average survival of 8 to 18 months has been reported by some authors. Lethal outcome within one year of the diagnosis has been documented in numerous reports and small studies. Our patient also died seven months of the first manifestation of symptoms. The presence of metastases at the time of diagnosis is considered to be a highly unfavorable prognostic factor.

References

Akin’s tumor – a rare tumor of the thoracopulmonary region.


Sažetak
ASKINOV TUMOR – RIJEĐAK TUMOR PRSIŠTA I PLUĆA

A Bekić, M Mehdić, M Gorščan, S. Kabul, D. Kompan i M. Hör


Ključne riječi: Plućne neoplazme – histologija; Plućne neoplazme – klinička; Plućne neoplazme – dijagnostika; Prikaz slučaja

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