A Rare Form of Mesothelioma: Malignant Pleural Deciduoid Mesothelioma

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Abstract

Mesothelioma usually develops in elder male patients that are exposed to asbestos. Deciduoid mesothelioma is a rare variant of epithelioid mesothelioma and has a poorer prognosis than epithelioid mesothelioma. As this subtype was seen in the peritoneum of young females and its relation with asbestos was not definite, these lesions were formerly thought to be stimulated by endogenous hormones. In the subsequent years, a relation was established between this tumor and asbestos since it was seen in the pericardium and pleura of males. In this paper, we present a 64-year-old male patient with asbestos exposure, diagnosed as having malignant pleural deciduoid mesothelioma, in which a long survival was provided with chemotherapy after surgery.

Keywords: Mesothelioma, asbestosis, surgery

INTRODUCTION

Deciduoid mesothelioma is a rare variant of epithelioid mesothelioma. It was first defined in the peritoneum of a young female, and as asbestos exposure was not present in the history of that patient, asbestos mineral has not been considered to play a role in the aetiology of deciduoid mesothelioma (1). The fact that it was reported particularly in young females led to the consideration that it might be associated with endogenous hormones. However, as the tumour was identified in the pericardium and pleura of males in the subsequent years, this consideration is no longer valid (2). It is known that histological differentiation of mesothelioma from adenocarcinoma is difficult. Therefore, accurate diagnosis of deciduoid mesothelioma, which has a poorer prognosis than epithelioid mesothelioma, is important in guiding the treatment (3).

In this paper, we presented a male patient, who was diagnosed with malignant pleural deciduoid mesothelioma, was performed pleuropneumonectomy, and was followed up for one year after post-operative chemotherapy without any relapses.

CASE PRESENTATION

A 64-year-old male patient was admitted to our clinic with shortness of breath and right chest pain. His anamnesis revealed no additional complaints such as cough, fever or weight loss. His physical examination was unremarkable except for decreased breath sounds in the right middle and lower zones. On his postero-anterior chest x-ray, a homogenous density enhancement, which obscured the right costophrenic angle and rose to the level of the anterior end of the 3rd rib with a parabolic contour, was observed (Figure 1). No culture growth, including tuberculosis, was identified in the pleural fluid obtained by thoracentesis. However, cytological examination of the pleural fluid demonstrated atypical mesothelial cell groups in a lymphocyte-rich background and it was reported that a tissue sample was required for
definite diagnosis. Considering that high levels of pleural fluid would hinder a potential pulmonary mass, computed tomography (CT) of thorax was not performed. In April 2011, the patient underwent pleural biopsy via right videothoracoscopy (VATS). On exploration that was performed during videothoracoscopy after draining 2500 cc of fluid, patchy irregular thickening and various sizes of nodules changing between 2 and 6 mm were identified in the parietal pleura. Postoperative thoracic CT demonstrated nodular infiltrations in the pulmonary parenchyma (Figure 2). Histopathological examination of the specimen was reported as "malignant deciduoid mesothelioma". Accordingly, a detailed anamnesis revealed that he had lived in a village until the age of 20 years and had been exposed to asbestos mineral. Positron Emission Tomography (PET-CT) and Magnetic resonance Imaging (MRI) of brain were performed to explore the presence of potential metastasis and no distant metastasis was detected (Figure 3). As mediastinal lymph node involvement was not detected in PET-CT, preoperative mediastinoscopy was not performed. In positron emission tomography, nodular changes characterized by 18-Fluoro-deoxy-glucose (FDG) uptake were reported in the mediastinum, costo-diaphragmatic pleura and fissures and loculated pleural effusion was identified in the right hemithorax (primarily in favour of mesothelioma). Pulmonary function tests and cardiac assessment revealed that the patient could tolerate a major surgery such as pleuropneumonectomy. Moreover, it was concluded that this surgery would be more convenient as the histopathological type of the tumour was epithelioid and invasion of both fissures was detected in PET scan. Following necessary preparations, thoracic cavity was opened after resection of the right 6th rib. Observing the invasion of the fissure, as was mentioned in the report of PET-CT, pleuropneumonectomy with partial resection of the diaphragm and pericardium were performed and polytetrafluoroethylene (PTFE) grafts were used for reconstruction. After the patient was followed up in the intensive care unit for one day, he was transferred to the ward and was discharged from the hospital on the postoperative 6th day. Histopathological examination of the material was reported as follows: "The tumour involves entire pleura and infiltrates the lung. No tumour is seen on the rib. The resected 15 lymph nodes (2R=4, 4R=6 and around the main bronchus =5) are reactive. No tumour in bronchi and surgical border". Therefore, consultation was requested from the oncology clinic and the patient was started on chemotherapy 3 months after the surgery (Augusts 2011). The patient received 4 cycles of cisplatinum (75 mg/m²) and pemetrexed disodium (500 mg/m²), and then followed up. No relapse was determined on his examinations including Thoracic CT, Brain MRI, and PET-CT on the postoperative 1st year. Consent was obtained from the patient for writing this case report.

DISCUSSION
The pleura, which is involved by secondary tumours rather than primary tumours, is a membrane, in which lung and breast cancer metastases are detected most frequently. Mesothelioma arises from multipotent mesothelial or subserosal cells that are likely to produce epithelioid or sarcomatoid cancers. As is known, it is difficult to differentiate mesothelioma, which is more prevalent among advanced aged males with a history of asbestos exposure, from mesothelial adenocarcinoma (4). Diagnosis of mesothelioma is made by immunohistochemical examinations; deciduoid type, a variant of epithelioid type, is rarely seen. Less than 50% of this type detected in less than 50 cases has been reported in the pleura (3). Immunostaining techniques used in the differential diagnosis include pancytokeratin, cytokeratin (CK) 5/6, calretinin and Wilms tumor (WT1). While epithelioid type is strongly stained with pancytokeratin, CK 5/6 is beneficial in the differentiation of mesothelioma from pulmonary adenocarcinoma. Calretinin is the best known and the most specific marker of mesothelioma and positive nuclear staining is only seen in mesothelioma (5). The present case was differentiated from metastatic adenocarcinoma by detecting positive "anti-calretinin antibodies" in the staining of the specimens obtained by thoracoscopic biopsy. In addition, identification of deciduoid cells with large eosinophilic cytoplasm and strongly positive anti-nuclear calretinin antibodies made the diagnosis definite (Figure 3).

It is known that asbestos mineral plays a role in the aetiology of mesothelioma. However, initially, such a relation could not be established with the deciduoid subtype, but thereafter a probable relationship with asbestos has been considered based on reported papers (6). When the anamnesis of the present case was detailed, we as well learned that he was exposed to asbestos mineral, which is used in
the rural areas in Turkey for in-house whitewash instead of lime and
known as “white soil” and contains asbestos mineral, for his house
was in the Ovacık village of Biga county of Çanakkale, where he had
lived until the age of 20 years. The mean time-span for development
of asbestos-induced mesothelioma is reported between 14 and 72
years (7). In the present case, the disease appeared approximately 40
years later, which is consistent with the literature.

Mean survival reported for deciduoid mesothelioma, which is more
aggressive as compared to epithelioid mesothelioma, is 9.6 months (1,
2). However, there are authors reporting cases that lived for a mean of
23 months (17-39 months) only with chemotherapy or symptomatic
treatment without surgery (8). Nevertheless, authors could not explain
such a long survival and recommended investigation of new cases. In
the present case, we made the definite diagnosis by thoracoscopic bi-
opsy, which was performed due to the failure in making diagnosis via
thoracentesis, and performed systemic screening. The present patient,
who had no metastasis received chemotherapy after radical surgery,
and is being followed up for 12 months with no relapse.

Radical surgery in malignant mesothelioma is currently a debatable
issue. Because of the absence of randomized studies, the Mesothel-
ioma and Radical Surgery (MARS) study did not report a definite
judgment about the surgical method that is superior (pleurectomy or
extrapleural pneumonectomy) (9). We performed extrapleural pneu-
monectomy since we detected fissure on PET-CT and during surgery.
Moreover, preoperative evaluation of the patient revealed that he
could tolerate this radical surgery.

CONCLUSION
Malignant deciduoid mesothelioma, which could be detected in both
genders, could involve both the pleura and the peritoneum. It is a
variant of epithelioid mesothelioma, has a poorer prognosis and its
relation with asbestos is being increasingly detected. Along with the
case presented herein, we think that combination of radical surgery
and chemotherapy may contribute to longer survival and reduce re-
lapses in this subtype of mesothelioma, and that the etiopathogen-
essis of this tumour will be enlightened more along with immunohis-
tochemical studies.

Informed Consent: Written informed consent was obtained from patients
who participated in this study.

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