Ewing’s Sarcoma/Primitive Neuroectodermal Tumor (ES/PNET) in the lung- A Rare Case Report

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ABSTRACT

Ewing’s sarcoma/Primitive Neuroectodermal Tumor is an uncommon malignant bone neoplasm that usually occurs in children and young adults and involves major long bones, pelvis and ribs. However, Ewing’s sarcoma/Primitive neuroectodermal tumor in the lung is very rare. We herein describe a rare case of ES/PNET in the right lung of 21 year old man, who presented with shortness of breath of rapid progression. X-ray chest and CT Scan of the chest showed a right lower lobe mass. Microscopic examination of the mass lesion was suggestive of Ewing’s sarcoma/Primitive Neuroectodermal Tumor and was confirmed by immunohistochemical analysis.

Keywords: Ewing’s sarcoma/Primitive neuroectodermal tumor, Lung, CD99.

INTRODUCTION

Primitive neuroectodermal tumor is an uncommon tumor, classically described under small blue round cell tumors with neural differentiation and developing from the nervous or sympathetic system. PNETs described originally by Askin and colleagues are infrequent thoracic tumors found in infants and children. The condition is exceedingly rare in adults. PNET is classified into 2 types based on location in
the body: Peripheral PNET (ES) and CNS PNET. Peripheral PNET is now thought to be identical to Ewing’s Sarcoma, based on evidence that indicates that both ES and PNET have a similar neural phenotype and, because they share an identical chromosome translocation.\(^1,2\) Primitive neuroectodermal tumors are tumors that demonstrate neural differentiation by light microscopy, IHC, or electron microscopy, where as Ewing’s sarcomas are undifferentiated by these analysis. PNET of the CNS generally refers to supratentorial PNETs.\(^2\) To the best of our knowledge, only eight cases of primary pulmonary ES/PNET have been reported in the literature. We present here such a rare case of Ewing’s Sarcoma/Primitive Neuroectodermal Tumor in the lung of young adult male.

CASE REPORT

A young adult of 21 years complained of shortness of breath, cough and pain in chest since 20 days. He was a non-smoker with no history of exposure to any occupational or inorganic dusts. There was no clubbing and lymphadenopathy on general examination. The respiratory system evaluation revealed signs of mass in the right hemithorax. Complete blood counts and biochemical investigations were within normal limits. X-ray chest showed a mass in the right lower lobe (figure-1). CECT of the thorax showed a large heterogenous lower lobe mass, in the right hemithorax (figure-2). There was no involvement of mediastinal lymph nodes. Right lower lobectomy was done and sent for histopathological examination. Grossly, cut section of the tumor was gray white with large areas of hemorrhage and necrosis (figure-3). Histologically, the tumor showed closely packed uniform round cell sheets with intervening delicate fibrovascular stroma. The cells show small round to ovoid nucleus with fine powdery chromatin and scant pale eosinophilic cytoplasm and indistinct cell borders (figure-4, 5). Some of the cells had vacuolated cytoplasm. At this point, small blue round cell tumor was suspected. The sections were then subjected to immunohistochemistry for CK, LCA, TTF-1 and CD99. IHC staining for CD99 (figure-6) showed positive result and there was negative staining for CK, LCA and TTF-1. Based on histomorphology and IHC the case was diagnosed as Ewing’s Sarcoma/ Primitive Neuroectodermal Tumor in the lung.

DISCUSSION

Ewing’s Sarcoma and extraskeletal Ewing’s sarcoma were first described by James Ewing and Tefft.\(^3,4\) Since then these two conditions have been confused histologically with other small blue round cell tumors, including PNET,\(^3,4\) but recent immunoperoxidase and cytogenic studies showed that PNET and ES belongs to the same group with varying degrees of neural differentiation, and they were categorized into a group known as the Ewing family of tumors.\(^1,4\) The ES/PNET family is a rare malignant tumor that shares common histological features of small primitive round cells. It most often arises in the soft tissues and bones, but has rarely been reported on other sites, such as the ovary, testis, uterus, kidney, pancreas, palate, myocardium, and lung.\(^5-8,14\) The morphological features of intrapulmonary tumors are similar to those of the ES/PNET in a variety of other locations.

The tumors are diagnosed using several modalities such as light microscopy, immunohistochemistry, and cytogenetics.\(^9\) Immunohistochemical staining is positive for neuron-specific enolase (60%), S-100 protein (50%), and CD99 marker (90%), as well as negative findings for leukocyte common antigen, epithelial membrane antigen, cytokerin.\(^9\) In this case, positive
staining for CD99 was observed confirming the diagnosis.

The CT-scan finding of ES/PNET reported is a heterogeneously enhancing mass. In our case also heterogeneously enhancing opacity was observed.

The treatment for ES/PNET is aggressive with the most effective treatment being a surgical resection with combination chemotherapy and high-dose radiation therapy. When resectable, the surgical excision must be wide. Right lower lobectomy was performed and combination chemotherapy was subsequently administered. As a part of a multidisciplinary team approach, physiotherapy and rehabilitation were given to achieve optimal survival and better quality of life.

CONCLUSION

Although, the ES/PNET in the lung is very rare soft tissue tumor, it should be considered in the differential diagnosis of a lung mass. The diagnosis is difficult on light microscopy alone and relies on immunohistochemistry for confirmation. These tumors present at a younger age, and are very aggressive. They usually have a poor prognosis despite treatment.

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**Figure 1.** Chest radiograph showing mass lesion in the right lower lobe with right pleural effusion

**Figure 2.** CT Scan thorax showing heterogenously enhancing mass lesion in the right lower lobe
Figure 3. Gross picture showing friable and haemorrhagic lesion involving entire lower lobe.

Figure 4. Microscopy showing closely packed tumor cells in sheets separated by thin fibrovascular septae. H&E, x100.
Figure 5. High power showing uniform round to oval cells with powdery chromatin, H&E, x400

Figure 6. Immunohistochemically tumor cells showing membrane positivity for CD99, x400