Primitive Neuroectodermal Tumor of Kidney: A Rare Case Report in an Elderly Lady

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Abstract

Primitive Neuroectodermal Tumors (PNET) are malignant, small round cell tumors (RCTs), which commonly develop in bone and soft-tissue of the body. In this article we report a case of Renal PNET that presented in an elderly lady of 85 years of age.

Introduction

Primitive Neuroectodermal Tumors (PNET) are malignant, small round cell tumors (RCTs), which commonly develop in bone and soft-tissue of the body [1,2]. Rarely have they been found in other parts of the body, such as the kidney, bladder, prostate, testis, ovary and uterus, etc [3-6]. PNET of the kidney are very rare malignant neoplasms. They were first recognized by Stout in 1918 [7]. Because of the similarity of PNET to Ewing’s tumor, it is difficult to estimate their exact number [8]. Until date, about 50 cases have been reported in the literature. These tumors frequently arise in childhood and adolescence with an aggressive clinical course [9]. In this article we report a case of Renal PNET that presented in an elderly lady of 85 years of age.

Case Presentation

An 85 year old female patient presented with vague left loin discomfort of a few days duration. On routine Ultrasonography of Abdomen and pelvis she was found to have a mass lesion in the left Kidney involving the mid and upper pole, with dimensions – 20 cm × 15 cm × 7 cm. The mass was reported to be confined to the kidney with no perinephric extension. The diaphragm and the retroperitoneum were found to be free from any infiltration. A subsequent Contrast enhanced CT abdomen and pelvis revealed a similar finding with a doubtful lesion in the lower lobe of left lung. There was no renal vein or IVC involvement or thrombus. Few Lymph nodes were identified in the renal Hilum. Patient was offered exploratory laparotomy and Left Radical Nephrectomy. Patient underwent the same on 31/03/2017. Operative findings – A highly vascular mass lesion with dimensions as per the CT findings, involving the mid and upper pole of left kidney with vascular and tissue infiltration into the Psoas sheath posteriorly and into the diaphragm was noted. Radical nephrectomy was performed with few hilar lymph nodes. Grossly, the nephrectomy specimen measured 17 cm × 16 cm × 7.5 cms with large tumor at the upper pole which measured 15 cm × 14 cm × 6.5 cms. The tumor occupied the entire kidney with a very thin rim of parenchyma seen at the periphery, at the lower pole. The tumor was seen infiltrating the renal sinus, capsule and perirenal fat. Cut section of the tumor was variegated, grey white with hemorrhagic and necrotic areas. The attached ureter appeared unremarkable. Microscopically, it showed an infiltrating malignant tumor with tumor cells arranged diffusely in sheets with a perivascular preservation of cells and intervening necrotic areas. Tumor cells demonstrated rounded nuclei with monomorphic appearance. Mitotic figures were noted with few atypical mitoses. Tumor was seen invading the renal sinus, renal capsule with extension into the perirenal fat. Ureretic resection margin was free of tumor.

Immunohistochemically (IHC), the tumor cells were positive for Vimentin, CD 99(Mic-2), NSE and CD 56. They were negative for Wilm’s tumor protein 1, pancytokeratin, synaptophysin and chromogranin. The Ki-67 index was 10-15%. Based on the morphological features and IHC, a diagnosis of Renal PNET was made.

Discussion

Renal PNET appears to be a unique clinical entity that behaves more aggressively than PNET arising at other sites [2]. First described by Stout in 1918, the concept of PNET has evolved to
include a group of small round cell malignancies of ubiquitous location and presumed neur ectodermal origin. They are defined by the expression of the same protooncogene [10] and the presence of a balanced t (11;22) (q24;q12) chromosomal translocation or a (21;22) rearrangement that results in the fusion of the gene EWS with the FLI1 or ERG gene, respectively [11]. PNET manifests a continuum of neurogenic differentiation, with Ewing’s sarcoma representing the least differentiated and peripheral neuroepithelioma the most differentiated forms [12]. In general, PNET is a very aggressive neoplasm, with 25-50% of patients presenting with metastatic disease, and a 5-year disease free survival rate of 45-55% [13-15]. Renal location of PNET is extremely rare, with only three previously reported cases [16-18] and a small series from the National Wilms Tumor Study (NWTS) published only in abstract form [19]. The largest series of renal PNET is a series from India published by Thyvihally et al. This series included 16 patients who were treated in Tata Memorial hospital. However, the first report came in the year 1994 when Mor et al. [17] described a characteristic renal neoplasia consistent with
the diagnosis of malignant peripheral PNET. The male: female ratio is about 3:1. The presenting symptoms and clinical signs are non-specific and similar to those of other renal tumors. The tumor has been mainly reported in the younger age group with age ranging from 4 years to 61 years. However, our patient was much older than this. Of all reported metastatic cases, 44% present with lung metastases only and 51% have a bone or bone marrow involvement with or without lung metastases and 5% present with metastases in other organs.

**Conclusion**

In conclusion it can be said that renal PNET is an extremely rare peripheral PNET having neural crest origin. They have aggressive clinical course and poor prognosis. It usually presents in adolescents and young adults, but our case was different as it presented in an elderly woman.

**References**


**Figure 8:** Wilm’s tumor protein 1 Negative.