Clear Cell Sarcoma of the Kidney: Correlational Study

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Abstract
Clear cell sarcoma of kidney is a rare malignant renal neoplasm of childhood, known for its aggressiveness, its tendency for recurrence and metastasis to bone. Three cases of clear cell sarcoma are being reported.

Case 1: A two year old male child having complaint of pain in right lumbar region. USG abdomen revealed right hypoechoic lesion. CECT abdomen revealed 13x11x10cm heterogenous mass involving right kidney. Microscopy revealed clear cell sarcoma of kidney (classic pattern). Immunohistochemistry showed strong vimentin positivity and negative for cytokeratin and WT-1.

Case 2: A thirty months old male child having complaint of pain in left abdominal region since 8 months. There was history of fever and increased micturition. There was no history of weight loss. His vitals stable. Biochemical and hematological profiles were normal. USG revealed left heterogenous mass lesion ms 6x4x2cm involving left kidney. CT revealed hypoechoic heterogenous mass ms 6x4x2cm involving left kidney. Microscopy revealed clear cell sarcoma.

Case 3: A 2 year old male child presented with right abdominal pain since 1 year. There was no history of fever, weight loss, hematuria. On examination patient had pallor only. Hematological finding normal. Urine examination normal. Chest X-ray normal. USG revealed right hypoechoic heterogenous mass ms13x10x10cm involving right kidney. CT revealed hypoechoic heterogenous mass ms 13x10x10cm involving right kidney.

The purpose of this paper is to review the published series and case reports of CCSK and to create an up-to-date overview of clinical and histological features and IHC.

Keywords: Tumor, Malignant, Bone metastasizing, child.

Introduction-
Clear cell sarcoma of kidney (CCSK) is a rare malignant neoplasm of childhood, known for its aggressiveness, its tendency for recurrence and metastasis to bone1,2. Its peak incidence is in 3-5 years with slight male preponderance3. In this study, we report three cases of clear cell sarcoma.
kidney, which were diagnosed in Department of Pathology, SMS medical college, Jaipur.

**Case 1**
A two year old male child presented with complaint of pain in right lumbar region since 6 months. There was no history of hematuria, vomiting, loss of weight. On examination patient had pallor, Vitals were stable. Investigations revealed: Hemoglobin of 11gm%, normal total and differential cell count, ESR 15mm/hr. Urine examination was normal. USG whole abdomen revealed a 13x10x10cm hypoechoic heterogenous mass uniformly involving right kidney. CECT abdomen revealed 13x8x6cm heterogenous mass involving right kidney. Provisional diagnosis of Wilms tumor was made on radiology. Right nephrectomy was done and specimen was sent for histopathological examination. On gross examination nephrectomy specimen measured 15x12x12cm. External surface grey brown bosselated, covered by Gerotas fascia with no capsular breach. On cutting tumor ms 13x10x10cm, and grey white, glistening and showed focal whorled appearance. At one end normal kidney parenchyma identified (Fig 1). Microscopically revealed presence of monomorphic cells arranged in sheets and nests separated by delicate arborizing fibro vascular network. The cells were polygonal to spindle shaped with indistinct cell borders and vesicular nucleus showing grooves at many places and variable mitosis. Diagnosis of clear cell sarcoma of kidney (classic pattern) was made (Fig 2a and 2b). Immunohisto chemistry showed strong positivity for vimentin and negativity for cytokeratin and WT-1 (Fig 3a, b, c).

**Case 2**
An thirty months old male child presented with complaint of pain in left abdominal region since 8 months. There was past history of fever and increased micturition. His vitals were stable. Biochemical and Hematological profiles were normal. USG revealed left heterogenous mass lesion ms 6x4x2cm involving left kidney. CT revealed hypoechoic heterogenous mass ms 6x4x2cm involving left kidney. Patient underwent surgery and specimen was sent for histopathological examination. Gross examination showed unoriented grey white soft tissue mass ms 6x4x2cm. External surface was grey white. Cut surface fleshy grey white. Microscopically showed round to oval cells having vesicular nuclei, inconspicuous nucleoli, clear to eosinophilic cytoplasm arranged diffusely and in nest, surrounded by delicate arborizing vasculature. Occasional nuclear grooving also noted. Mitosis are brisk. Feature favoured diagnosis of clear cell sarcoma kidney (Classic pattern).

**Case 3**
A 2 year old male child presented with right abdominal pain since 1 year. There was no history of fever, weight loss, hematuria. On examination patient had pallor only. Hematological profile was normal. Urine examination normal. Chest X-ray normal. USG revealed right hypoechoic heterogenous mass ms13x10x10cm involving right kidney. CT revealed hypoechoic heterogenous mass ms 13x10x10cm involving right kidney. Right nephrectomy was done and specimen was sent for histopathological examination. On gross examination specimen measured 15x12x10cm. External surface bosselated in appearance. No breaching of capsule seen. Cut surface showed a tumor ms 13x10x10cm and grey white, showed whorling appearance. At one end normal kidney parenchyma identified. Microscopically revealed presence of monomorphic tumor cells arranged in cords, sheets, nests and alveolar pattern separated by thin capillary network. The cells were spindle shaped to oval with distinct cell borders and vesicular nucleus showing nuclear grooves at many places and variable mitosis surrounded by compressed normal renal tissue. Overall morphology favoured diagnosis of clear cell sarcoma of kidney (Classic pattern).
Fig 1. Right nephrectomy specimen of clear cell sarcoma kidney. Cut surface grey white, transluscent.

2a

Fig 2. Microscopic view a). Tumor cells arranged in nest pattern. (10x) b). Tumor cells having vacuolated cytoplasm and surrounds by delicate vasculature. (40x)

2b

Fig 3a). Tumor cells show vimentin positivity. b). Cytokeratin negative. c) WT1 negative.

3a

3b

3c

Clinical Features
The mean age of presentation in my study is about 24 months. Argani et al found that 50% of their 351 cases were diagnosed between 2 and 3 years of age. CCSK is extremely rare in the first 6 months of life and in adults. The youngest
CCSK patient reported in literature was a foetus (at 31 week gestation) and the oldest patient was 58 years old. Contrary to WT’s slight propensity to the female gender, a male predominance has been noted in all large CCSK series (average male to female ratio of about 2:1). In my study all three cases male predominance noted. All three patients in my study series show common clinical presentation of abdominal pain. One case has fever and increased micturition also noted. Other symptoms that have been described are hematuria, vomiting, decreased oral intake, constipation and hypertension in other studies which were absent in my study series. Some study suggested a predilection for involvement of the right kidney, in my study case 2 out 3 showed right predilection.

Pathology
All CCSK typically presents as a large, unicentric, well circumscribed and sharply demarcated mass. The tumor arises from the central region and replaces normal kidney tissue or is located in the medulla of the kidney. The diameter of the tumor ranges in my study from 6 to 13 cm. CCSK is soft, tan grey in color. External surface bosselated and cut surface glistened due to mucinous material in all cases. Microscopically classic pattern predominant on other histologic patterns. In other studies also classic pattern is more common. The classic subtype of CCSK is characterized by round /oval cells with clear cytoplasm, fairly uniform round/ oval and often vesicular bland nuclei with finely dispersed chromatin, inconspicuous nucleoli and infrequent mitotic figures. In the classic pattern, tumor cells are arranged in nests or cords, separated by fibrovascular septa. Other pattern that have been described include myxoid, sclerosing, epithelioid, myxoid, pallisading, storiform and anaplastic clear cell sarcoma. The histological heterogeneity emphasizes the importance of discriminating the disease from other entities. Considering the low incidence of CCSK from other renal tumors, it is important that suspected CCSK cases are reviewed by pathologists who were experts in paediatric renal tumors.

Immunohistochemistry
Immunohistochemistry can help to distinguish CCSK from other renal tumors of childhood. IHC done in one of the 3 cases shows non specific vimentin positivity and negativity for WT-1 and cytokeratin.

Discussion and Conclusion
Clear cell sarcoma also known as “Bone metastasizing tumor of Childhood” comprise 4% of all primary childhood renal tumors. Histological variants of CCSK include spindle cell, sclerosing, epithelioid, myxoid, pallisading, storiform and anaplastic clear cell sarcoma. Accurate and early diagnosis of clear cell sarcoma as a separate entity from Wilms tumor is of extreme importance. Careful study of morphology, the characteristic delicate arborizing vascular septae, monomorphous population and invasive property will help to distinguish it from more common and less aggressive counterparts of childhood renal neoplasm like Wilms tumor. Points helpful in this matter are: Foci of blastema are not seen in clear cell sarcoma; nonrenal elements such as cartilage or muscle are not found in clear cell sarcomas. Clear cell sarcomas are unilateral and unicentric, and sclerotic stroma is uncommon in Wilms tumor before therapy. The vascular pattern typical of clear cell sarcoma is often often helpful in distinguishing it from Wilms tumor. The border with the kidney is usually infiltrative in CCSK, whereas the border of Wilms tumor is typically pushing. Immunohistochemistry for Clear cell sarcoma which is positive for vimentin negative for WT1, and Wilms tumor positive for WT 1. Treatment of CCSK generally involve surgical intervention coupled with radiation and chemotherapy with cyclophosphamide, etoposide and vincristin and doxorubicin for 24 weeks. It is of considerable...
therapeutic importance that CCSK be not only correctly diagnosed but early too.

References