A kidney tumor to know: mucinous tubular and spindle cell carcinoma of the kidney (mtscck)

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Abstract

Mucinous tubular and spindle cell carcinoma of the kidney (MTSCC-K) is an unusual renal tumor. It is important to increase the recognition of MTSCC-K and improve the level of clinical diagnosis. We report a case of mucinous tubular and spindle cell carcinoma in a 50-year-old woman. The tumor, located on the left kidney, was well circumscribed and a partial nephrectomy was applied. In microscopically, the tumor was composed of cylindrical cells arranged in tubules and was making abrupt transitions to the spindle cell morphology in a myxoid stroma. Due to its favourable prognosis, mucinous tubular and spindle cell carcinoma must be differentiated from papillary renal cell carcinoma, especially that with sarcomatoid change.

Introduction

Mucinous tubular and spindle cell carcinoma of the kidney (MTSCC-K) is a rare renal epithelial tumor, believed to be a type of low-grade malignant tumor. The precise origin is unclear, certain researchers have hypothesised that it originates from the loop of Henle or the distal tubule. However, the majority of researchers hypothesise that its origin is in the distal tubule [1].

MTSCC-K is difficult to pathologically differentiate from high-grade malignant renal tumors, including collecting duct carcinoma and sarcomatoid carcinoma. It was first reported in 1997 by MacLennan et al and was known as a ‘low-grade collecting duct carcinoma’ [2].

It was introduced in the latest WHO classification of renal tumors in 2004 as a separate entity [3]. Through a new case and literature review, we discuss the histological features of the tumor as well as the differential diagnosis.

Case report

We report a new case in a 50-year-old woman presented with a 3-month history of left flank pain without gross hematuria and fever. The Renal function test results were normal, and without history of stone disease.

Hypovascular renal masse were noted on ultrasonography, and the imaging examination by abdominal computed tomography (CT) scan revealed a 7x5-cm well-circumscribed solid mass, protruding outside the renal contour in the superior pole of the left kidney. The CT was performed initially to obtain baseline attenuation values of lesions and to identify calcification with no clear enhancement identified in the corticomedullary phase. However, marginal uneven enhancement was observed in the nephrographic phase (Figure 1).

No metastasis was identified to the retroperitoneal lymph node, abdominal organs or lungs. The patient provided written informed consent.

The patient underwent partial nephrectomy resection of the left kidney by a left subcostal incision. No postoperative therapy was given to the patient.

Macroscopy: Dissection of the specimen revealed that the tumor was well-circumscribed, solid and grayish yellow in appearance, measuring 4.5x3.5x3.5 cm. No areas of hemorrhage were identified in the tumor. Unlike necrosis zones have been identified.

Figure 1. Contrast enhanced abdominal CT scan. It revealed a tumor on the superior pole of the kidney, and demonstrated the tumor was slightly enhanced on the corticomedullary phase and nephrographic phase.

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Microscopy: The tumor was composed of small, elongated cords or tubules, in a tightly packed arrangement. Myxoid stroma was shown to be interspersed among the tubular cells, and appeared to exhibit slender tubular spindle cell-like structures. Tumor cells were smaller and cubic-shaped or oval, with single small eosinophilic nucleoli and low-grade nuclei. Occasionally, necrosis and foam cell infiltration were identified. The myxoid stroma was stained by acidic mucus (Figure 2).

Discussion

Mucinous tubular and spindle cell carcinoma is a rare, malignant renal epithelial tumor which showed a female predominance and favorable prognosis. Furthermore, it has been recognized as a new entity of Renal Cell Carcinoma (RCC) in 2004, and usually behaved in a low-grade fashion [4]. More than 80 cases have been reported and not much is known about this tumor [5].

MTSCC-K has a wide age distribution, with an age range between 17 and 82 years and a mean age of 53 years. The female incidence is approximately three- or four-fold that of males [6-8].

However, the clinical symptoms and imaging characteristics of the MTSCC-K are similar to the more common renal cell carcinoma (RCC). Therefore, a clear preoperative diagnosis becomes more difficult [11].

The current case underwent abdominal CT scan prior to surgery and no clear enhancement was identified in the arterial phase. However, marginal uneven enhancement was identified in the venous phase. This change is consistent with the previous literature and may provide references for the preoperative clinical diagnosis of MTSCC-K. In addition, it has been reported that fine needle aspiration biopsy may be diagnostic of MTSCC-K, which may aid to improve preoperative diagnosis rates [12].

Therefore, the true origin of MTSCC-K still requires further exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. However, Hes et al. suggested that development of MTSCC-K could be related to a kidney stone, because of the total exploration. In cytogenetic studies, genetic abnormalities found in MTSCC-K cells are monosomies in chromosomes 1, 3, 6, 8 and 13, and total or partial trisomies of chromosomes 7, 11, 16 and 17 [6].

Immunohistochemistry is not helpful in discriminating between papillary RCC and MTSCC-K because the morphological interpretation is sufficient in the distinction between these tumors.

However, Cytokeratin CK7 and CK19, and RCC could be expressed in both MTSCC-K and renal cell carcinoma. Further, CD10 is lowly expressed in MTScottCC, indicating that CD10 may be a useful marker to differentially diagnose these tumors [19,20].

Conclusion

Overall, MTSCC-K exhibits a lower malignant degree and an improved prognosis compared with other types of RCC. In summary, mucinous tubular and spindle cell carcinoma is a rare and only recently described distinctive subtype of renal cell carcinoma. It must be differentiated from papillary renal cell carcinoma, especially with sarcomatoid change, which has a much weaker prognosis.

Ethical standards

I declare that all procedures followed during this research experiment comply with the highest local ethical standards.

Conflict of interest

Youness Dehayni, Mohamed Sinaa, Yassine Elabiad, Bouzid Balla, Abdelghani Ammani, Abdelmoumam Qarro and Mohamed Alami declare that they do not have any conflicts of interest.

References


