Bilateral renal cell carcinoma with leiomyomatous stroma: A rare entity diagnosed synchronously and treated surgically in a staged fashion

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ABSTRACT
Renal cell carcinoma (RCC) accounts for approximately 3% of adult malignancies and 90–95% of kidney neoplasms. Renal cell carcinoma with leiomyomatous stroma (RCCLS) is an extremely rare histopathological entity based on available literature data. Herein, we report a 31-year-old male with incidentally detected synchronous bilateral renal masses who was eventually found to harbor RCCLS after being operated sequentially via nephron-sparing surgery.

Keywords: Leiomyomatous stroma; nephron-sparing surgery; partial nephrectomy; renal cell carcinoma.

Introduction
Renal cell carcinoma (RCC) accounts for approximately 3% of adult malignancies and 90–95% of kidney neoplasms.¹⁻⁴ While the number of patients diagnosed with kidney cancer increases steadily, the incidence of bilateral renal tumors varies between 3-5%.⁵⁻⁷ Description of the RCC subtype has been extremely revised based on morphological and genetic features in the 2016 World Health Organization (WHO) renal tumour classification.⁸ Renal cell carcinoma with leiomyomatous stroma (RCCLS), which was not included in this new classification system. It is an extremely rare histopathological entity based on the available literature data which was firstly defined by Canzonieri et al in 1993.⁹ Thereafter several case reports have been published by various authors.¹⁰⁻¹³

Herein, we report a case with incidentally detected synchronous bilateral renal masses who was eventually found to harbor RCCs with leiomyomatous stroma after being operated sequentially via nephron-sparing surgery (NSS).
the middle zone while the coexistent right-sided counterpart was smaller (2.5 cm in maximal diameter) and situated at the posteromedial part of the lower kidney pole (Figure 1). The R.E.N.A.L. nephrometry scores of the masses were 7p and 9p for the right and left sides, respectively. Both masses demonstrated significant contrast enhancement, which was suggestive of an underlying malignancy. The Hounsfield units of the right and left renal masses were calculated to be 1046±52 HU and 1105±93 HU, respectively after IV administration of contrast medium. Regional lymph node enlargement was not evident on cross-sectional images.

The masses were solid in nature and radiologic characteristics (contrast enhancement, size, heterogeneous architecture etc.) substantially raised the possibility of malignancy. Additionally this was a young patient without any comorbidity, which meant a long- life expectancy. Therefore, renal mass biopsy, histopathological results of which would not change our treatment plan, was not considered for either side.

After thorough patient counseling, we elected to perform a 2-staged nephron-sparing approach with an initial open surgery for the left renal tumor due to its high R.E.N.A.L. nephrometry score and robot-assisted laparoscopic surgery for the right-sided kidney mass concerning its relative morphometric easiness.

**Procedure 1**
A left open partial nephrectomy was performed through 11-12th intercostal incision. After entering the retroperitoneum, left renal artery was identified, dissected and controlled with vessel loop (without creating warm ischemic conditions) through posterior approach. Thereafter, the upper pole and lateral borders of the kidney were mobilized within Gerota’s fascia, without dissecting the perirenal fat overlying the tumor. The mass was enucleoresected with a milimetric rim of normal looking renal parenchyma overlying its pseudocapsule. Frozen section evaluation of the surgical margin revealed benign findings. After repairing collecting system defects with 3/0 polyglactin sutures, the cortical edges were approximated by interrupted 2-0 monofilament sutures. Total operation and warm ischemia times were 97 and 19 minutes, respectively. Estimated blood loss was 150 cc. He was discharged after an uneventful course on the third postoperative day.

**Procedure 2**
One month later, transperitoneal robot-assisted laparoscopic right partial nephrectomy was performed. A total of 5 ports (3 for the robot, 2 for the assistant) were introduced. After mobilizing the colon, Gerota’s fascia was opened and the central renal mass was exposed with its overlying perirenal fatty tissue. Renal pedicle was dissected free and within 15 minutes of warm ischemia time the renal mass was completely enucleoresected and the resultant corticomedullary defect was repaired using the sliding-clip renorrhapy technique.

Histopathological examination of both tumors revealed the same diagnosis, i.e. renal cell carcinoma with leiomyomatous stroma (Figure 2). His postoperative serum creatinine level was 0.93 mg/dL and abdominal CT scan, which was conducted 6 months after the surgery, detected no local recurrence or distant metastasis (Figure 3). Written informed consent of the patient was obtained from the patient for the publication of the procedures as a case report.

**Histopathological examination**
Both left and right renal tumors revealed similar macroscopic features. Well-circumscribed, pale yellow, solid masses with diameters of 3.2 cm and 2.6 cm, were detected, respectively. Microscopically; tumors had a multinodular appearance, due to abundant fibroleiomyomatous stromal component intermingled with epithelial tumor cells. Epithelial component was composed of branching tubules, nests and glands lined by cells with clear cytoplasm and mild nuclear atypia (Fuhrman grade 2). Atypia, mitosis and necrosis were not identified.
The immunohistochemical study was performed using a Ventana BenchMark XT automated stainer (Ventana Medical System, Inc, Tuscon, AZ, USA). Formalin fixed, paraffin embedded tissues were immunostained. CK7 (OV-TL 12/30, monoclonal, RTU; Biocare Medical, CA, USA), pancytokeratin (multiple, monoclonal, RTU; ScyTek, Utah, USA), smooth muscle actin (1A4, monoclonal, RTU; ScyTek, Utah, USA) and desmin (D33, monoclonal, RTU; ScyTek, Utah, USA) were used as primary antibodies. Both tumors had similar immunohistochemical profiles namely epithelial cells showed diffuse and strong immunoreactivity for CK7 and pancytokeratin while the stromal component reacted positively for SMA and desmin.

Discussion

Leiomyomatous stroma may exist in various renal tumors, namely angiomyolipoma, renal angiomyoadenomatous tumor (RAT)/clear cell papillary renal cell carcinoma (CCPRCC) and “renal cell carcinoma with smooth muscle stroma” (RCCSMS).[9-11,13] RCCLS is an uncommon neoplasm, which has an intimate admixture of epithelial and stromal components.[14] Clear epithelial cells represent the epithelial component of this neoplasm and moderate nuclear atypia (mostly Fuhrman grade 2) that is evident in adenomatous structures encompassing a nested or tubular pattern admixed with focal papillary and solid areas constitute the main histopathological alterations observed in RCCLS.[14]

Petersson et al.[15] concluded that the leiomyomatous stroma that may be observed in RCC is not related to the neoplastic process.[15] The smooth muscle cells of this stroma are of polyclonal in origin and represent a reactive hyperplastic process possibly being derived from the muscular layer of large caliber veins located at the capsular periphery or within the tumoral septae. The proliferation of vessel-derived smooth muscle cells may be related to hypoxia induced factors as suggested by Kuhn et al.[10]

These tumors do not have unique genetic alterations.[16,17] On the other hand, some authors have reported molecular changes that resemble those encountered in clear cell RCCs (VHL gene mutation, VHL hypermethylation or loss of heterozygosity of 3p).[8,18] Considering these controversial findings about the relationship between genetic alterations and the development of renal cell carcinoma with leiomyomatous stroma (RCCLS) we did not performe molecular genetic evaluation either for the resected tissue specimen or peripheral blood sample.

Due to the rare incidence of RCCLS, the prognosis of this entity has not been thoroughly evaluated. Nevertheless a good clinical behaviour is likely, given the small sized lesions and relatively lower tumor grades published up to now. Shannon et al.[11] reported that all of their patients (n: 3) were alive with no evidence of disease after 18 months to 5 years of follow-up. In this publication[11], patients’ ages ranged between 48-65 years. One patient was female and the others were male. Despite their small size (2.4-2.6 cm) these solid masses were managed by radical nephrectomy. Our case was a relatively young male patient and had bilateral cT1a masses (2.6 and 3.2 cm), which were treated by NSS. Fuhrman grades of these masses, including the ones reported by Shannon et al.[11], were ≤2. In another study, all 3 patients remained well and alive after 20-52 months of follow-up.[13] Despite the relatively short follow-up period, which is one of the major drawbacks of the present study, our patient still remains disease free.

In conclusion, RCCLS is a rare subtype of RCC. Its histopathological diagnosis needs a meticulous immunohistochemical work-up. Genetic evaluation, which was not done in our patient, can be considered in selected cases, however there is limited data to support its routine application. Nephron-sparing surgery, which was inevitable in our case given the young age of the patient and synchronous bilateral presentation, should be the treatment modality of choice whenever its technically feasible. Despite limited data about its natural hist-
tory, this tumor portends a favorable prognosis once excised completely with surgically negative margins.

**Informed Consent:** Written informed consent of the procedures and publication as a ‘case report’ was obtained from the patient.

**Peer-review:** Externally peer-reviewed.


**Conflict of Interest:** No conflict of interest was declared by the authors.

**Financial Disclosure:** The authors declared that this study has received no financial support.

**References**

cinomas is polyclonal and not part of the neoplastic process. Virchows Archiv 2014;465:89-96. [CrossRef]

