

## Case Report

# Staged, Open, No-Ischemia Nephron-Sparing Surgery for Bilateral-Multiple Kidney Tumors in a Patient with Birt-Hogg-Dubé Syndrome

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Hereditary kidney cancer patients with bilateral multiple kidney tumors represent challenges in the era of rapidly growing minimal invasive treatment techniques. Birt-Hogg-Dubé Syndrome (BHDS) is an autosomal dominant genodermatosis characterized by a triad of benign skin tumors (fibrofolliculomas, trichodiscomas, acrochordons) together with an increased risk of developing malignant renal tumors and pulmonary disease such as pneumothoraces and multiple lung cysts. The morbidity and mortality of the affected patients is determined by the presence of the kidney tumors, which tend to be multifocal and bilateral, as observed in other hereditary kidney cancer syndromes like von Hippel-Lindau disease, familial leiomyomatosis, and hereditary papillary renal cell carcinoma. Herein, a patient with BHDS, presenting with synchronous bilateral multiple kidney tumors, is reported. The report describes the management of kidney tumors with two-stage open nephron-sparing surgery in which the nonvascular clamping technique was utilized.

## 1. Introduction

Birt-Hogg-Dubé Syndrome (BHDS) is an autosomal dominantly inherited genodermatosis caused as a result of the mutations in the FLCN gene encoding the protein “folliculin,” a new protein which hypothetically functions as a tumor suppressor gene [1–4]. The affected patients develop cutaneous follicular tumors (multiple fibrofolliculomas, trichodiscomas, and acrochordons) over the face, neck, and upper trunk and are susceptible to develop renal neoplasms and pulmonary disease (spontaneous pneumothoraces, multiple lung cysts).

Kidney tumors in these patients tend to be multifocal and bilateral, as observed in other hereditary cancer syndromes such as von Hippel-Lindau disease, familial leiomyomatosis, and hereditary papillary renal cell carcinoma [5, 6]. Kidney

tumors in BHDS are commonly and characteristically reported as “hybrid oncocytic renal cell carcinoma,” which contain areas of oncocytomas as well as areas of renal cell carcinoma, chromophobe, or others [4, 7]. Kidney tumors are the major determinants of morbidity and mortality in patients with BHDS; hence the overall renal functions have to be preserved as much as possible while completely removing the tumors within oncological principles [5]. Therefore, this syndrome may represent as an outstanding indication for open-staged nephron-sparing surgery (NSS), in the era of everyday growing laparoscopic/robot-assisted surgery [6–8].

We herein report a patient with BHDS, presenting with typical urological, dermatological, and radiological findings (Figures 1, 2, and 3). Kidney tumors of the patient were managed with two-staged open NSS, and we describe our no-ischemia surgical technique with the kidney under perfusion.

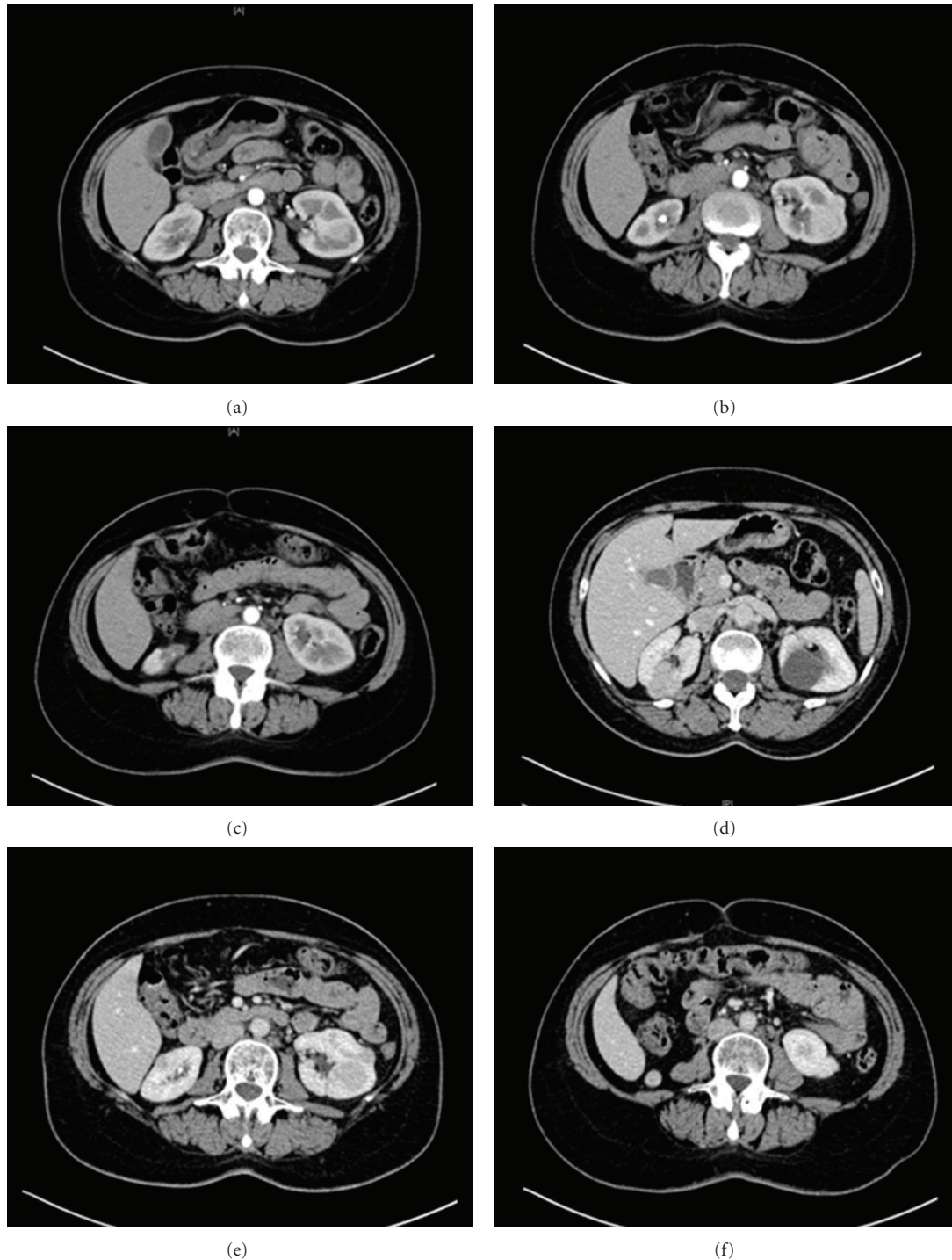


FIGURE 1: Preoperative contrasted CT scans of the patient, showing multiple bilateral kidney tumors, with diameters ranging between 1 and 5 cm. Radiologically, most of the tumors have malignant characteristics due to their hypervascularity and radiopaque enhancement.

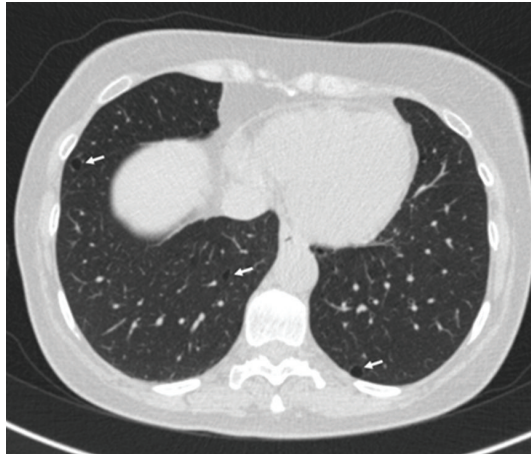
## 2. Case Report

A 56-year-old woman presented with bilateral multiple kidney tumors, diagnosed radiologically during her evaluation for hematuria.

Computed tomography revealed 4 tumors, diameters ranging between 1 and 3 cm in the left kidney, and 4 tumors,

diameters ranging between 1 and 5 cm in the right kidney (Figure 1). Radiologically, most of the tumors had malignant characteristics due to their hypervascularity and radiopaque enhancement.

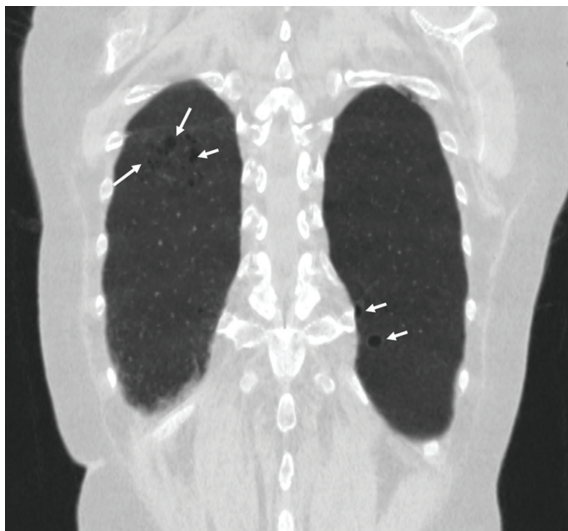
Her chest X-Ray and CT scan of thorax revealed multiple, randomly distributed air-filled small cysts (Figure 2). Intervening lung parenchyma appeared normal. Coronal images,



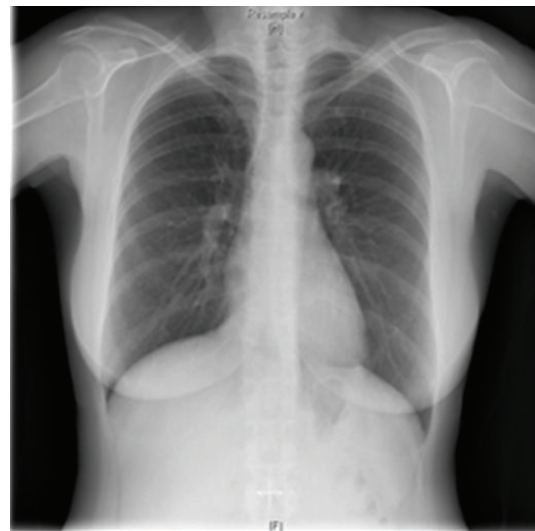
(a) Axial CT images show small cysts randomly distributed



(b) Intervening lung parenchyma appears normal



(c) Coronal image reconstructed with “minimum intensity projection” technique shows small cysts; some are clustered in the right upper lobe. No pneumothorax is identified



(d) X-ray of the chest

FIGURE 2: Preoperative CT scan of the thorax (a–c) and chest X-ray (d).

reconstructed with “minimum intensity projection” technique, showed small cysts; most of them clustered in the right upper lobe. No pneumothorax was identified.

On dermatological examination, she had multiple, asymptomatic, white, 1–3 mm sized, dome-shaped, firm papules over the face, neck, scalp, upper trunk, and back, and multiple skin colored, soft, pedunculated papules on the axillae that were clinically consistent with BHDS (Figure 3). Her dermatological history revealed that her facial papules were present since her early twenties and continued to increase in number.

The clinical diagnosis of the patient was consistent with BHDS.

The patient was scheduled first for a staged approach starting from the right side. Preoperative serum analyses were within normal ranges (serum creatinine: 0.7 mg/dL, BUN: 13 mg/dL, Hb: 13.3 gr/dL), and the patient was healthy otherwise.

Nephron-sparing surgeries were performed using standard intercostals (between 11th–12th ribs) flank incision on each side. The technique consisted of adequate exposure of the kidney. Gerota’s fascia was opened; perinephric fatty tissue was dissected off. Ureter and the vascular pedicle were exposed and marked with vessel loops. After documenting all tumors on the kidney, tumors were “enucleoresected” with the kidney under perfusion, first cauterizing the capsule surrounding the tumor and then by removing the tumor with the help of scissors and brain dissector. First the smaller tumors were enucleated, in order to diminish the risk of bleeding. Frozen sections from tumor beds were done for all tumors. Bleeding from the tumor bed was controlled with 3/0 polyglactin (Vicryl) interrupted sutures and parenchyma was adapted with 2/0 polyglactin (Vicryl) sutures over a surgical bolster. Renal vessels were neither clamped nor compressed during these manipulations.



FIGURE 3: (a) Multiple, asymptomatic, white, dome-shaped, firm papules over the face. (b) Multiple skin colored, soft, pedunculated papules on the axillae (acrochordons).

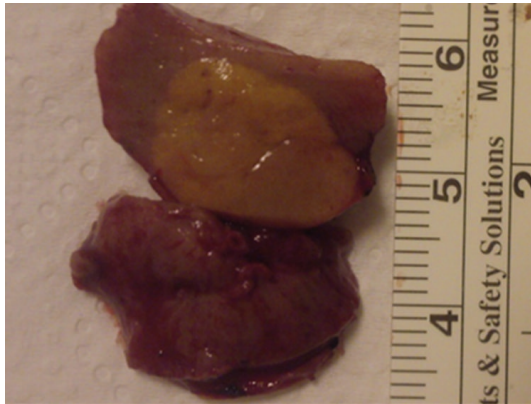


FIGURE 4: Macroscopic view of one of the tumors removed. The cut section appears light brown and glistening. No necrosis or hemorrhage is noted.

A total of 4 tumors were enucleated from the right kidney. Operation time was 105 minutes. Time spent during tumor enucleations and renographies was 45 minutes. Estimated blood loss was 100 mL. Postoperative course was uneventful, with a postoperative serum creatinine of 0.7 mg/dL and serum Hb of 11.9 gr/dL. The patient was discharged 4 days after left NSS.

The left kidney was treated 6 weeks after the first stage. Using the same, open, no-ischemia NSS technique, a total of 4 tumors were enucleated from the left kidney. Operation time of the left side was 120 minutes. Time spent during tumor enucleations and renographies was 55 minutes. Estimated blood loss was 150 mL. Postoperative course was also uneventful on the right side and the patient was discharged 4 days after surgery.

On macroscopic examination (Figure 4), the excised tumors of the right kidney measured 3.7 cm, 1.8 cm, 1.7 cm, and 1.2 cm in maximal diameter, respectively. The cut sections of each tumor appeared light brown and glistening. No

necrosis or hemorrhage was noted. Excised tumors of the left kidney measured 2.7 cm, 2.1 cm, 1.8 cm, and 0.5 cm in maximal diameter, respectively. Cut sections of each tumor were the same with those of right kidney. All of the tumors appeared sharply circumscribed.

Histomorphologically, each tumor revealed similar features (Figure 5). The H&E sections demonstrated histological features of both oncocytoma and conventional variant of renal cell carcinoma (clear cell RCC). Within these hybrid oncocytic tumors were areas classic for oncocytoma (large cells, ill-defined cell borders, finely granular eosinophilic cytoplasm throughout, and large nuclei with homogeneous basophilic chromatin) and other areas consistent with clear cell RCC (well-defined cell borders, clear cytoplasm, round nuclei with centrally located inconspicuous nucleoli). Fuhrman nuclear grade was II in clear cell RCC component. Mitotic features were sparse; no atypical forms were identified. Surgical margins were all negative.

The final serum creatinin level, seen 2 years after the last operation, was 0.9 mg/dL, and her abdominal MR scans (Figure 6) did not show any recurrences or new tumor formation.

### 3. Discussion

Partial nephrectomy, or “NSS,” has become the standard procedure for management of small renal tumors [9, 10]. However, there is a debate going on regarding the technique of applying renal ischemia during NSS to avoid renal damage in the treated kidney [10]. Current literature supports safe ischemia times, within 20 minutes of warm ischemia and up to 2 hours of cold ischemia, to minimize renal ischemic damage. Thus, if ischemia is required, the tumor should be removed within 20 minutes of warm ischemia, regardless of surgical approach, either open or laparoscopic [10].

However, most of the data regarding our current knowledge on this subject is based on large case series, which are composed of “single” small renal masses, and patients with hereditary kidney cancer, such as von Hippel-Lindau disease,

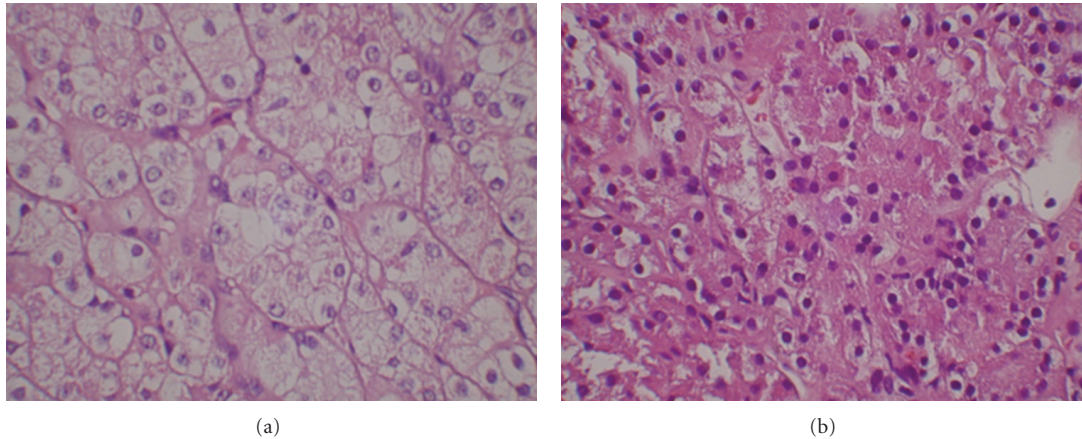


FIGURE 5: Microscopic views of H&E sections, demonstrating histological features of both oncocytoma and conventional variant of renal cell carcinoma.

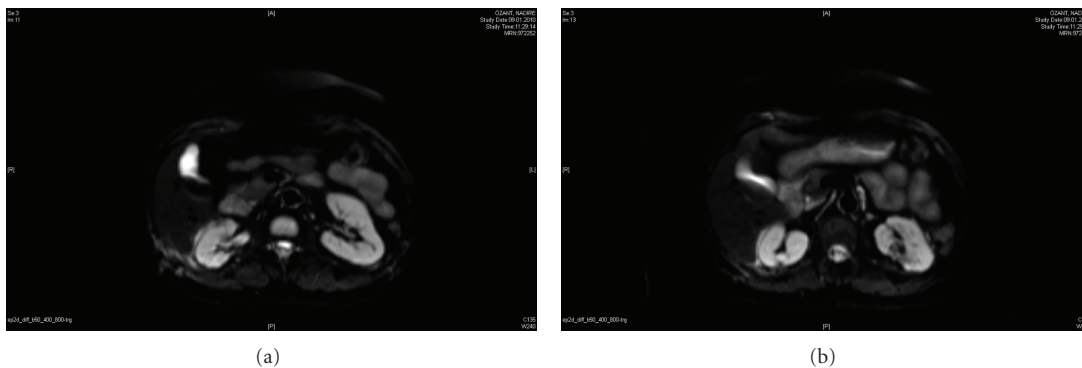


FIGURE 6: MRI scans performed 2 years after surgery, documenting no tumoral lesion.

familial leiomyomatosis, and hereditary papillary renal cell carcinoma usually presenting with bilateral multiple tumors, constitute challenges [9, 11]. Taking into consideration that these patients develop renal tumors at younger ages and that their kidney pathologies are the major determinants of their prognosis, a great care has to be given in their management.

The incidence of multifocality of RCC ranges between 6 and 25% in published series [12, 13]. The dilemma of the management of ipsilateral multiple tumors has recently been investigated by Lin et al. [11]. They compared the operative findings, renal functional outcomes, and intermediate-term cancer-specific survival of patients undergoing either laparoscopic partial nephrectomy or laparoscopic cryoablation and report similar results. However, the number of tumors they treated ranged between 2 and 3 per kidney, and the majority of tumors were <4 cm in size. Furthermore they report a median warm ischemia time of 36 minutes (range: 12–48), although they underline that the postoperative renal function was preserved in all patients in the intermediate term. Interestingly, they managed 2 patients in their series under cold ischemia only without vascular clamping but had to clamp the hilum because of a bleeding during tumor removal [11].

In the present patient, a total of 8 kidney tumors were removed with open NSS from two kidneys and the total surgical manipulation time just for tumor removals and renographies was 45 minutes per kidney, without vascular clamping. This time period is clearly higher than the universally accepted limits under warm ischemia. So even in highly experienced hands, using either open or robot assisted/laparoscopic approach, vascular clamping would increase the risk of renal function loss.

Recently, increasing number of papers are getting published regarding the safety of partial nephrectomy without renal hilar occlusion in selected cases [14]. Robot-assisted surgical techniques did clearly supply manipulative advantages, especially during reconstruction, when compared to pure laparoscopic techniques. However, multiple renal tumor removal for multifocal renal cancer, without renal hilar clamping, has not been reported with robot-assisted surgery yet.

In conclusion, we present the case that the nonvascular clamping open NSS, when used in patients with hereditary kidney cancer syndromes such as BHDS, minimizes the risk of renal function loss.

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