ARTICLE

Check for updates

Recurrent *KRAS* mutations are early events in the development of papillary renal neoplasm with reverse polarity

Khaleel I. Al-Obaidy ¹, Rola M. Saleeb^{2,13}, Kiril Trpkov ³, Sean R. Williamson ¹, Ankur R. Sangoi⁴, Mehdi Nassiri^{5,13}, Ondrej Hes⁶, Rodolfo Montironi⁷, Alessia Cimadamore⁷, Andres M. Acosta⁸, Zainab I. Alruwaii ⁹, Ahmad Alkashash⁵, Oudai Hassan¹⁰, Nilesh Gupta¹⁰, Adeboye O. Osunkoya ¹¹, Joyashree D. Sen⁵, Lee Ann Baldrige⁵, Wael A. Sakr¹², Muhammad T. Idrees ², John N. Eble⁵, David J. Grignon^{5,13} and Liang Cheng ⁵

© The Author(s), under exclusive licence to United States & Canadian Academy of Pathology 2022

We evaluated the clinicopathologic and molecular characteristics of mostly incidentally detected, small, papillary renal neoplasms with reverse polarity (PRNRP). The cohort comprised 50 PRNRP from 46 patients, divided into 2 groups. The clinically undetected (<5 mm) neoplasms (n = 34; 68%) had a median size of 1.1 mm (range 0.2–4.3 mm; mean 1.4 mm), and the clinically detected (≥ 5 mm) neoplasms (n = 16; 32%) which had a median size of 13 mm (range 9–30 mm; mean 16 mm). Neoplasms were positive for GATA3 (n = 47; 100%) and L1CAM (n = 34/38; 89%) and were negative for vimentin (n = 0/44; 0%) and, to a lesser extent, AMACR [(n = 12/46; 26%; weak = 9, weak/moderate = 3)]. *KRAS* mutations were found in 44% (n = 15/34) of the clinically undetected PRNRP and 88% of the clinically detected PRNRP (n = 14/16). The two clinically detected PRNRP with wild-type *KRAS* gene were markedly cystic and contained microscopic intracystic tumors. In the clinically undetected PRNRP, the detected *KRAS* mutations rate was higher in those measuring ≥ 1 mm vs <1 mm [n = 14/19 (74%) vs n = 1/15 (7%)]. Overall, the *KRAS* mutations were present in exon 2—codon 12: c.35 G > T (n = 21), c.34 G > T (n = 3), c.35 G > A (n = 2), c.34 G > C (n = 2) resulting in p.Gly12Val, p. Gly12Asp, p.Gly12Cys and p.Gly12Arg, respectively. One PRNRP had a G12A/V/D complex mutation. Twenty-six PRNRP were concurrently present with other tumors of different histologic subtypes in the ipsilateral kidney; molecular testing of 8 of the latter showed wild-type *KRAS* gene despite the presence of *KRAS* mutations in 5 concurrent PRNRP. On follow up, no adverse pathologic events were seen (range 1–160 months; mean 44 months). In conclusion, the presence of *KRAS* mutations in small, clinically undetected PRNRP provides a unique finding to this entity and supports its being an early event in the development of these neoplasms.

Modern Pathology (2022) 35:1279-1286; https://doi.org/10.1038/s41379-022-01018-6

INTRODUCTION

In 2019, we described a distinct subset of epithelial renal neoplasms "papillary renal neoplasm with reverse polarity" (PRNRP) that was characterized by papillary or tubulopapillary architectures, with delicate fibrovascular cores lined by a single layer of eosinophilic cells with finely granular cytoplasm, which had apically located round nuclei with inconspicuous nucleoli. PRNRP was uniformly positive for GATA3 and L1CAM and negative for vimentin and, to a lesser extent, α -methylacyl-CoA-racemase (AMACR/p504s) by immunohistochemistry¹. Furthermore, we demonstrated that these tumors harbor recurrent *KRAS* mutations, which are not known to occur in other types of renal epithelial neoplasms². Genitourinary Pathology Society (GUPS) has considered PRNRP in its recent update on existing renal neoplasms to represent a distinct pattern/subtype within the spectrum of

papillary RCC, indicating that additional studies are needed to further validate it³. Cases with similar or identical morphology to PRNRP have been described in earlier studies as "papillary renal cell carcinoma with oncocytic cells and nonoverlapping low-grade nuclei^{#4} and "papillary renal cell carcinoma type 4/ oncocytic low grade^{"5}, based on their distinct morphology and similar or identical immunoprofile. More recently, several reports have also been published on this entity, reinforcing our morphologic and molecular findings^{6–12}.

In our examination of resected end-stage kidneys, we found small clinically undetected lesions (<5 mm) that were morphologically identical to the PRNRP. These lesions were previously called papillary adenoma type 2, or, more recently, type D^{13,14}. In a prior molecular interrogation of a limited number of samples, we identified recurrent *KRAS* mutations in 3 of these small lesions¹⁵. In the current study, we

Received: 16 November 2021 Revised: 21 January 2022 Accepted: 21 January 2022 Published online: 12 February 2022

¹Robert J Tomsich Pathology and Laboratory Medicine Institute, Cleveland Clinic, Cleveland, OH, USA. ²Department of Laboratory Medicine and Pathobiology, University of Toronto, Toronto, ON, Canada. ³Department of Pathology and Laboratory Medicine, Cumming School of Medicine, University of Calgary, Calgary, AB, Canada. ⁴Department of Pathology, El Camino Hospital, Mountain View, CA, USA. ⁵Department of Pathology and Laboratory Medicine, School of Medicine, Indiana University, Indianapolis, IN, USA. ⁶Department of Pathology, Charles University in Prague, Faculty of Medicine and University Hospital in Plzen, Plzen, Czech Republic. ⁷Section of Pathological Anatomy, School of Medicine, United Hospitals, Marche Polytechnic University, Ancona, Italy. ⁸Women's and Perinatal Pathology, Department of Pathology, Brigham and Women's Hospital, Harvard Medical School, Boston, MA, USA. ⁹Department of Pathology, Regional Laboratory and Blood Bank, Eastern Province, Dammam, Saudi Arabia. ¹⁰Department of Pathology and Laboratory Medicine, Henry Ford Hospital, Detroit, MI, USA. ¹¹Department of Pathology, Emory University School of Medicine, Atlanta, GA, USA. ¹²Department of Pathology, Wayne State University/ Detroit Medical Center, Detroit, MI, USA. ¹³Deceased: David J. Grignon ⁵⁶email: liang_cheng@yahoo.com

1280

analyzed the immunohistochemical characteristics and performed targeted polymerase chain reaction for *KRAS* mutations on an expanded cohort of PRNRP.

MATERIALS AND METHODS Study population and case selection

The files of the pathology departments at the participating institutions were searched for lesions that fulfilled our previously suggested diagnostic criteria for papillary renal neoplasm with reverse polarity (PRNRP)¹. A series of 177 end-stage renal disease specimens from the Indiana University Health archives were reviewed in search of small lesions that resembled PRNRP. Additionally, kidney resection specimens with a diagnosis of papillary adenoma were also re-examined in search of these neoplasms by one of the authors (KIA). A search of additional material from our previous cohort¹ identified sufficient tissue for further analysis in three cases (corresponding to tumors 3, 9, and 16 of the original study). The cases selected for inclusion in the study were divided into small/ clinically undetected [size <5 mm (radiologically and/or grossly undetected –non-incidental)] neoplasms.

Due to the shared positive immunohistochemical reaction for GATA3¹⁶, 10 randomly selected clear cell papillary renal cell carcinomas, were retrieved from the pathology archives at Indiana University.

Histopathologic study

Tissues from all tumors were fixed in 10% neutral buffered formalin and embedded in paraffin. Sections of 4 µm thickness were stained with hematoxylin and eosin. Based on our previous study¹ and other published reports, as recently summarized by Wei et al.¹⁰, a panel of 4 differentiating antibodies between PRNRP and PRCC was utilized. These were directed against GATA3 (L50-829; Biocare Medical, Pacheco, CA), L1CAM (UJ127; Sigma-Aldrich, St. Louis, MO), vimentin (V9; Dako) and α -methylacyl-CoAracemase (AMACR/p504S [13H4; Dako]) and utilized in a Dako automated instrument. Positive and negative controls provided appropriate results for each stain. The staining was recorded as negative (–) or positive (+), the latter indicating a diffuse and strong reaction. For tumors with positive reactions other than strong and diffuse, weak (w) or moderate (m) intensity was recorded.

KRAS mutational analysis

A targeted polymerase chain reaction analysis was performed as previously described¹⁷. Briefly, formalin-fixed, paraffin-embedded tissue blocks were retrieved from each of the 50 neoplasms. DNA extraction from the microdissected tumors areas from 4 um cut slides was performed using the Qiagen QIAamp DNA FFPE Tissue Kit (Qiagen, Valencia, CA, USA). DNA concentration was determined using the NanoDrop Spectrophotometer and adjusted to ~10 ng/ μ l in ddH₂O. Polymerase chain reaction testing was performed according to the recommended procedure using the Qiagen therascreen KRAS RGQ PCR Kit on the Qiagen Rotor-Gene Q MDx instrument. The therascreen KRAS RGQ PCR Kit provides eight separate PCR amplification reactions: seven mutation-specific reactions in codons 12 and 13 of exon 2 of the KRAS oncogene [Gly12Ala (G12A), Gly12Asp (G12D), Gly12Arg (G12R), Gly12Cys (G12C), Gly12Ser (G12S), Gly12Val (G12V), and Gly13Asp (G13D)] and a wild-type control in exon 4. Analysis of crossing thresholds and mutation calls for each PCR amplification reaction was performed by the Rotor-Gene Q therascreen KRAS Assay analysis software once runs were completed. KRAS analysis was also performed on the clear cell papillary renal cell carcinomas, the concurrent ipsilateral renal tumors of different or similar morphologies, and to 4 "mimickers" described in the original series¹.

RESULTS

Clinical and pathologic features

The clinical and pathologic features are tabulated in Tables 1 and 2. The clinically undetected neoplasms constituted 68% (n = 34/50) and ranged from 0.2 to 4.3 mm (mean 1.4 mm, median 1.2 mm). All were diagnosed on nephrectomy (27 total and 7 partial) specimens performed either for end-stage kidney disease (n = 12/34; 35%) or for concurrent ipsilateral tumor (n = 23/34; 68%) with/or without end-stage kidney disease. The clinically detected ones constituted

32% (n = 16/50) and ranged from 9 to 30 mm (mean 16 mm, median 13 mm). All, except one (n = 15/16), were diagnosed on nephrectomy (5 total and 10 partial) specimens performed for radiologically detected neoplasms, with the last case being diagnosed on biopsy. Concurrent ipsilateral tumors were present in 3 cases. All neoplasms were staging category pT1a and showed low nucleolar grade [International Society of Urological Pathology/WHO (ISUP/WHO) grades 1–2]. On follow-up, all patients with available follow-up had a benign neoplasm-related clinical course (range 1–160 months, mean 44 months, median 30 months), although 5 patients died of other causes.

Morphologically, both groups of neoplasms shared similar morphologic features as previously described¹. However, in addition to the commonly described features, cystic expansion with intracystic papillary proliferation was seen in 7 of the clinically detected (≥5 mm) neoplasms. These were filled with eosinophilic proteinaceous material that contained loose macrophages with or without hemosiderin deposits and cellular debris (Fig. 1A). Although the majority of neoplasms were composed of thin cores of arborizing papillary, or less frequently, tubulopapillary, architecture, 3 of the clinically detected neoplasms had edematous and focally fibrotic papillary cores, these frequently contained chronic inflammatory infiltrates (Fig. 1B). Cytologically, the cells were mostly cuboidal cells with eosinophilic finely granular cytoplasm; focal areas of clear cell changes were seen in 3 neoplasms (Fig. 1C). The nuclei were characteristically located at the apical surface opposite to the basement membrane with smooth overlying luminal borders, with hobnail conformation present in 3 neoplasms (Fig. 1D). As we described before, the nuclei were mostly small, non-overlapping with regular contours, frequent nuclear clearing, inconspicuous nucleoli and had only rare areas of wrinkled nuclear contours. Focal areas of conspicuous basophilic nucleoli were seen in the areas of nuclear clearing, however, large prominent and eosinophilic nucleoli (WHO/ISUP nucleolar grade 3) were not seen. Psammoma bodies, intracellular hemosiderin, coagulative-type tumor necrosis, or mitotic figures were absent in all neoplasms.

Immunohistochemically, all evaluated neoplasms showed positive reactions for GATA3 (n = 47/47; 100%) (strong in 46 and weak to moderate in 1). Reactions for L1CAM were positive in 89% of the neoplasms [n = 34/38 (strong in 29 and weak in 5)]. Reactions for vimentin were negative in all neoplasms (n = 0/46; 0%) while reactions for AMACR were mostly negative (n = 34/46; 74%), however, 20% (n = 9/46) showed weak "blush-like" reactions and 3 (n = 3/46; 6%) showed weak to moderate reactions. In these neoplasms, the AMACR reaction had a similar intensity to that seen in the distal convoluted tubules of the kidney (Fig. 2). Interestingly, clear cell papillary renal cell carcinoma tumors showed positive reaction for GATA3 in 80% (n = 8/10); however, in contrast with PRNRP, they were consistently positive for vimentin (n = 10/10; 100%) and negative for L1CAM (n = 0/10; 0%).

KRAS mutational analysis

KRAS mutations were found in 44% (n = 15/34) of the clinically undetected (microscopic) PRNRP. The detected *KRAS* mutation rate was higher [n = 14/19 (74%)] in those measuring 1–5 mm than those measuring <1 mm, where only one PRNRP (measuring 0.6 mm) was positive for *KRAS* mutation [n = 1/15 (7%)] (Fig. 3). Of the clinically detected PRNRP, 88% (n = 14/16) had *KRAS* mutations with the 2 negative neoplasms being markedly cystic and contained small microscopic intracystic tumors. Overall, *KRAS* mutations were in codon 12: c.35 G > T (n = 21), c.34 G > T (n = 3), c.35 G > A (n = 2), c.34 G > C (n = 2) resulting in p.Gly12Val, p. Gly12Asp, p.Gly12Cys and p.Gly12Arg, respectively. One PRNRP had a G12A/V/D complex mutation (Table 3).

No *KRAS* mutations were found either in the clear cell papillary renal cell carcinomas, in 8 of the concurrent ipsilateral tumors of different histology, or the mimickers of PRNRP, included in our previous report¹.

| | VMACR | | | | | d P | ЧЪ | | | | | | | | | > | > | | |
|--------------------------|--|-------------------------------|------------------------|----------------------|--------------------|----------------------|----------------------|---------------------|---------------------|----------------------|------------------------|------------------------|----------------------|------------------------|---|----------------------|---|------------------------|---|
| | | | | | ſ | 2 | 2 | | | | ſ | I | | I | | 2 | 2 | 1 | |
| | VIME | I | I | I | I | ЧN | ЧN | I | I | I | I | I | I | I | 1 | I | I | I | |
| | Istochen L1CAM | + | + | + | I | NP | NP | + | + | 3 | 3 | 3 | I | 3 | + | + | ٩ | ΝΡ | |
| 40 | nmunon ATA-3 | | | | | | 4 | | | | | | | | | | | | |
| - - | hs) G I | + | + | + | + | + | z | + | + | + | + | + | + | + | + | + | + | + | |
| | Follow up durati (Mont | 25 | 84 | 20 | Ш | 160 | 113 | II | Ш | 45 | 134 | 114 | 105 | T | ~ | 35 | 84 | 87 | ç |
| | utsome | A | Q | Q | | G | Х | | | Х | Q | Q | Q | | A | Ģ | Ð | Ģ | ç |
| Ċ | 5 | Ï | Ï | Ï | = (1 |) NE | ă (| | | ă |) NE | Ï | Ï | I | ž | Ï | ž | Ë | 4 |
| this cohort. | Assocaited tumor type (Size; KRAS status) | I | I | PRCC (2 CM) | PRCC (3.5 CN | PRCC (1.5 CM | PRCC (3.8 CN | 11 | 11 | PA (0.5 CM) | CCRCC (2 CN | Oncocytoma (4.2 cm) | CCRCC (7.8 cm) | CCRCC (4.7 cm) | ACD-RCC (2.4 cm; KRAS wild-type); PRNRP (0.8 cm; KRAS mutated) | I | Multiple oncocytomas (KRAS wild- type) | Oncocytoma (1.0 cm) | |
| polarity included in | Multiple tumors | z | z | ۲ | | ۶ | × | | | × | ~ | ۲ | ۲ | ۲ | > | z | > | × | |
| with reverse | KAS status | 12A/V/D omplex lutation | .G12V 35 G > T) | .G12V 35 G > T) | .G12V 35 G > T) | ۵ | .G12V .35 G > T) | .G12C .34 G > T) | .G12V .35 G > T) | | ۵ | .G12V .35 G > T) | .G12D 35 G > A) | .G12C .34 G > T) | ۵ | .G12V 34 G > T) | .G12V 35 G > T) | ۵ | |
| plasms | ¥ Ê | 5 5 E | ч С | άÇ | ч, С | z | ч С | ч С | ч с | z | z | ч, С | άQ | ч С | z | ч С | ч, С | z | |
| enal nec | bize (m | 14 4 | 10 | 1.6 | 1.4 | 0.7 | 1.4 | 1.9 | 28 | 0.9 | 1.4 | 1:1 | 2.1 | - | 0.6 | 10 | 2.1 | 1.2 | 0 |
| apillary | CKD | z | z | ≻ | | ≻ | ≻ | | | ≻ | z | z | z | z | ≻ | ≻ | z | ≻ | ; |
| a of the 50 _F | Laterality | Right | I | Left | Right | Right | Left | 11 | 11 | Right | Right | Left | Right | Left | Right | Right | I | Left | |
| ogic dat | Age | 48 | 45 | 52 | | 42 | 60 | | | 64 | 54 | 63 | 56 | 65 | 75 | 54 | 17 | 70 | ć |
| ed clinicopatholo | Procedure | Partial Nehprectomy | Partial Nehprectomy | Total nephrectomy | | Total nephrectomy | Total nephrectomy | | | Total nephrectomy | Partial Nehprectomy | Total nephrectomy | Total nephrectomy | Partial Nehprectomy | Total nephrectomy | Total nephrectomy | Total nephrectomy | Total nephrectomy | |
| . Detail | nder | | | | | _ | - | | | - | - | | | | | | | | |
| - C | le | | < | | | | | | | | | | < | < | < | | | | |

| Tabl | le 1. conti | nued | | | | | | | | | | | | | |
|------|-------------|------------------------|-----|------------|-------|-----------|------------------------|------------------------|--|---------|----------------------------|---------|-----------|----------|-------|
| | Gender | Procedure | Age | Laterality | ESRD/ | Size (mm) | KRAS status | Multiple tumors | Assocaited | Outsome | Follow | lmmunoh | istochemi | stry | |
| | | | | | ĝ | | | | tumor type (Size; KRAS status) | | up duration (Months) | GATA-3 | L1CAM | VIMENTIN | AMACR |
| 19 | ٤ | Total nephrectomy | 76 | Left | z | 2.8 | p.G12V (c.35 G > T) | ≻ | PRCC (6.5 cm) | NED | 87 | + | NP | I | I |
| 20 | ш | Total nephrectomy | 51 | I | ≻ | 16 | p.G12V (c.35 G > T) | z | I | NED | 60 | + | NP | I | I |
| 21 | ш | Total nephrectomy | 54 | Right | ~ | 0.4 | ND | z | I | NED | 60 | + | + | I | I |
| 22 | Σ | Partial Nehprectomy | 12 | Left | z | 0.6 | QN | ~ | CCRCC (6 CM); CCPRCC (0.6 CM) | DOC | Ŋ | + | 3 | I | 8 |
| 23 | ш | Partial Nehprectomy | 56 | I | z | 11 | p.G12V (c.35 G > T) | z | I | NED | 75 | + | NP | N | NP |
| 24 | ш | Total nephrectomy | 84 | Right | ~ | 0.8 | ND | z | I | NED | 24 | + | I | I | I |
| 25 | ш | Total nephrectomy | 72 | Left | ~ | 1.4 | p.G12V (c.35 G > T) | z | I | DOC | 6 | + | + | I | 1 |
| 26 | ш | Total nephrectomy | 33 | Left | ~ | 0.8 | Ŋ | z | I | NED | 52 | + | + | I | I |
| 27 | ш | Total nephrectomy | 73 | Right | ~ | 0.9 | QN | > | ACD-RCC (3.6 cm; KRAS wild-type); URCC (5.5 cm) | 1 | I | + | + | I | 1 |
| 28 | Σ | Total nephrectomy | 55 | Left | ~ | 0.6 | Ŋ | z | T | NED | 56 | + | I | I | > |
| 29 | ш | Partial Nehprectomy | 68 | Right | z | 10 | ND | z | T | NED | 36 | + | NP | I | I |
| 30 | ш | Partial Nehprectomy | 58 | Left | z | 1:1 | p.G12V (c.35 G > T) | ۶ | ChRCC (3.8 CM; KRAS wild-type) | NED | 45 | + | 3 | 1 | I |
| 31 | ш | Total nephrectomy | 12 | Right | z | 3.9 | QN | ~ | ChRCC (4.5 CM; KRAS wild-type) | I | I | ЧN | ЧN | I | I |
| 32 | ш | Total nephrectomy | 64 | Right | z | 30 | p.G12R (c.34 G > C) | z | PRNRP (1 cm) | I | I | + | + | I | I |
| 33 | T | Partial Nehprectomy | I | I | I | 16 | p.G12V (c.35 G > T) | 1 | I | I | I | + | + | I | 8 |
| 34 | ш | Partial Nehprectomy | 72 | Left | z | 6 | ŊŊ | 1 | I | NED | 24 | ٩N | ЧN | N | NP |
| 35 | ш | Partial Nehprectomy | 62 | Right | z | 0.6 | ŊŊ | | CCRCC (3.2 CM) | NED | 30 | + | + | I | I |
| 36 | ш | Partial Nehprectomy | 63 | Left | z | 1.5 | ND | ≻ | Oncocytoma (2.3 CM) | NED | 7 | + | + | I | > |
| 37 | Σ | Total nenhrectomy | 58 | Left | ≻ | 0.8 | QN | z | T | NED | 22 | + | + | I | I |

| fer | Descedure | Age | | | | | | | | | i di a constante di | • | | |
|-----|------------------------|-----|------------|-------|-----------|------------------------|-----------------|--|---------|----------------------------|---------------------|-----------|----------|-------|
| | ר היפמחו ב | 2 | Laterality | ESRD/ | Size (mm) | KRAS status | Multiple tumors | Assocalted | Outsome | Follow | Inonumu | istochemi | stry | |
| | | | | 3 | | | | tumor type (Size; KRAS status) | | up duration (Months) | GATA-3 | L1CAM | VIMENTIN | AMACR |
| | Total nephrectomy | 72 | Left | ≻ | 6.0 | QN | ~ | Multiple bilateral PAs | NED | 30 | + | ЧN | I | I |
| | Total nephrectomy | 74 | Left | ≻ | 1.8 | p.G12V (c.35 G > T) | z | C/L UC of renal pelvis and PCa | NED | - | + | + | I | 1 |
| | Total nephrectomy | 7 | Left | ≻ | 0.6 | p.G12V (c.35 G > T) | ~ | Right sided ACD-RCC (0.9 CM) | NED | 10 | + | + | I | E |
| | Total nephrectomy | II | Right | | 0.2 | QN | 11 | 1 | II | II | + | + | I | E |
| | Partial Nehprectomy | 79 | Right | z | 16 | p.G12V (c.35 G > T) | z | 1 | NED | 13 | + | + | I | 8 |
| | Total nephrectomy | 68 | Right | ~ | 17 | p.G12V (c.35 G > T) | ~ | ACD-RCC (0.8 CM; KRAS wild-type); CCPRCC (1.5 CM; KRAS wild-type); AML (0.6 CM; KRAS wild- type) | NED | 6 | + | + | 1 | 1 |
| | II | II | | ~ | 6 | p.G12V (c.35 G > T) | II | | 11 | II | + | + | I | I |
| | Total nephrectomy | 71 | Right | z | 1.5 | p.G12V (c.35 G > T) | ~ | URCC (5.5 CM) | NED | - | + | + | I | I |
| | Partial Nehprectomy | 72 | Left | ≻ | 10 | p.G12V (c.35 G > T) | z | I | NED | 12 | + | + | I | > |
| | Total nephrectomy | 73 | Left | z | 4.3 | QN | ~ | PRCC (5.5 CM) | DOC | e | + | + | I | I |
| | Partial Nehprectomy | 58 | Right | z | 6 | p.G12V (c.35 G > T) | ~ | PRCC (4.5 CM) | NED | 15 | + | + | I | m/m |
| | Renal Biopsy | 68 | Left | z | 30 | p.G12D (c.35 G > A) | z | 1 | NED | 4 | + | + | I | 3 |
| | Partial Nehprectomy | 68 | Right | z | 21 | p.G12R (c.34 G > C) | z | 1 | NED | 80 | + | + | I | I |

CINICL CINOMOPDODE RLC, PHNNP papiliary renal neoplasm with reverse polarity, UKLL unclassined renal cell carcinoma, C/L contralateral, UC ur disease, DOC died of other causes, NP not performed, w weak, m moderate. - not available/applicable, = the same patient in the above row.

1284

DISCUSSION

We recently introduced the term "papillary renal neoplasm with reverse polarity (PRNRP)" to describe a subset of renal epithelial

 Table 2.
 A summary of the clinical and pathologic features of the papillary renal neoplasm with reverse polarity cohort.

| Parameter | Clinically undetected | Clinically detected |
|-------------|--|---|
| Age (years) | | |
| Range | 33–84 | 45–79 |
| Mean | 64 | 62 |
| Median | 65 | 64 |
| Gender | | |
| Male | 17/31 (55%) | 4/16 (25%) |
| Female | 14/31 (45%) | 12/16 (75%) |
| Laterality | | |
| Right | 15/33 (45%) | 9/12 (75%) |
| Left | 18/33 (55%) | 3/12 (25%) |
| Size (mm) | | |
| Range | 0.2–4.3 | 9–30 |
| Mean | 1.4 | 16 |
| Median | 1.1 | 13 |
| IHC | | |
| GATA3 | 32/32 [100% (1 = weak)] | 15/15 (100%) |
| L1CAM | 22/26 [85% (6 = weak)] | 12/12 (100%) |
| AMACR | 6/32 [19% (4 = weak; 2 = moderate)] | 6/14 [43% (5 = weak; 1 = weak-moderate)] |
| Vimentin | 0/32 (0%) | 0/12 (0%) |
| | | |

tumors with characteristic features. In the original series, we found distinctive morphologic and immunohistochemical features that set this tumor aside from other described renal cell carcinomas, including papillary renal cell carcinoma (PRCC). Our original series included radiologically detected masses and found alterations of the KRAS molecular pathway in 9 of 10 analyzed cases while none of the control series of PRCCs had these mutations. In the current study, in which we deliberately focused on clinically undetected and incidental neoplasms, we identified recurrent KRAS mutations in neoplasms as small as 0.6 mm, although the mutation detection rate was higher in those measuring ≥ 1 mm. Chang et al⁶ have attributed this to the limited tumor content. However, considering that our clinical assay is routinely used for the detection of mutations in small tumor contents from cytology samples, this is unlikely. However, using more sensitive assays such as digital droplet PCR might be useful to address this tumor content issue.

We used the term "reverse polarity" for this entity to indicate the consistently unusual nuclear localization toward the apical end of the cell as we discussed in detail in our original study¹. Additionally, we intentionally avoided the use of descriptive terminology related to the cytoplasmic staining "eosinophilic or oncocytic" because the acquisition of more knowledge about this entity may identify different cytoplasmic qualities, either focally or non-focally. For instance, "chromophobe" renal cell carcinoma is a term used to this day, which means a "hard to stain"¹⁸, which later on had led to the addition of the "eosinophilic" prefix to some of these tumors. Of note, we identified focal clear cell changes in 3 neoplasms in the current series, a finding described recently by Chang et al as well⁶. Moreover, we did not use such descriptive terminology to avoid lumping it with the prior studies which showed a wide range of morphologic and immunohistochemical features to what has been described as "oncocytic papillary renal cell carcinoma". Therefore, we believe that it is best to maintain



Fig. 1 Unusual features of papillary renal neoplasm with reverse polarity. These include cystic neoplasm filled with eosinophilic proteinaceous material containing loose hemosiderin-laden macrophages and cellular debris (A), edematous and expanded fibrovascular cores containing chronic inflammatory infiltrates (B), focal clear cell changes (C), and hobnail conformation (D).

the "papillary renal neoplasm with reverse polarity" terminology for these distinct neoplasms, and also to avoid terminological differences in future studies.



Fig. 2 Immunohistochemical reaction for AMACR. AMACR staining showing weak to moderate reaction, with similar intensity to that observed in the distal convoluted tubules and collecting ducts and less than the proximal convoluted tubules (inset; arrows are pointing out the distal convoluted tubules).



Fig. 3 Microscopic papillary renal neoplasm with reverse polarity. A representative image of the smallest (0.6 mm) papillary renal neoplasm with reverse polarity with detected *KRAS* mutation in the current series.

Including this series, 158 such neoplasms have been reported with marked consistency in the morphology, despite small variations and some features that were not mentioned in our original series, including focal hobnail conformation and focal clear cell changes. Additionally, a recent paper by Wei et al documented that these neoplasms may be cystic¹⁰. The cysts are usually filled with proteinaceous fluid or blood and may contain floating degenerated cells and filled with hemosiderin. The appearance of thick intracystic proteinaceous fluid or blood may be confused with necrosis, however, true tumor type coagulative necrosis is absent in these neoplasms.

The immunohistochemical profile of PRNRP is consistent. They are consistently positive for GATA3 and keratin 7 and are negative for vimentin. The presence of focal weak positive reaction for vimentin reported in 6% (7/118)^{7,19} is highly atypical; the interpretation of vimentin staining may sometimes be difficult, particularly when the fibrovascular cores are compressed or fibrotic resulting in a false-positive reaction. One PRNRP was reported to show focal reaction with anti-CD117 (KIT) antibody¹⁰ however, as observed by Chang et al, these neoplasms may contain mast cells within the fibrovascular cores, and if prominent. they may give an impression of a positive reaction⁶. Lastly, although these neoplasms typically lack the increased immunoexpression of α-methylacyl-CoA-racemase (AMACR), it is not unusual to see weak cytoplasmic blush or even, in rare instances, moderate reactivity. In these scenarios, however, the expression of AMACR appears to match the expression seen in the distal rather than proximal convoluted tubules, which may serve as internal control when assessing the immunoexpression of AMACR in these neoplasms.

In the recent paper by the Genitourinary Pathology Society (GUPS), it was suggested that the PRNRP represents a subtype of PRCC along with the biphasic, solid, Warthin-like tumors³. Structurally, although both PRCC and PRNRP have predominantly papillary architecture, the presence of papillary structures is not restricted to PRCC and can be prominent in other tumors, including clear cell papillary renal cell carcinoma, hereditary leiomyomatosis and renal cell carcinoma-associated renal cell carcinoma (HLRCC), and MiT-Family translocation renal cell carcinoma. The positive immunohistochemical reactions for GATA3 and L1CAM, the increased mRNA expression of these genes, as well as the negative reactions for vimentin and, to less extent, AMACR, are features that contrast with PRCC including the other suggested subtypes. Additionally, the shared chromosome 7 and/ or 17 trisomy and/ or deletion of Y chromosome in males, that we identified in our original series, is not limited to PRCC and PRNRP, but was also reported in 30–40% of HLRCC and tubulocystic carcinoma^{20–23}. Moreover, the presence of *KRAS* mutations was proven by multiple independent studies to be unique and consistent in PRNRP (mutational detection rate was up to 93% in some studies); therefore, the presence of KRAS mutations in other tumors [0.6% (n = 2/290) of the papillary renal

Table 3. A summary of the KRAS mutation status of the papillary renal neoplasm with reverse polarity cohort.

| Category | | Clinically undetected | | Clinically detected |
|----------------|-----------------------------|-----------------------|---------------------|---------------------|
| Size range (n) | | < 1 mm (n = 15) | ≥ 1-4.3 mm (n = 19) | 9-30 mm (n = 16) |
| KRAS mutation | | 1 (7%) | 14 (74%) | 14 (88%) |
| Variant | p.G12V (c.35 G > T) | 1 | 11 | 9 |
| | p.G12D (c.35 G > A) | 0 | 1 | 1 |
| | p.G12C (c.34 G > T) | 0 | 2 | 1 |
| | p.G12R (c.34 G > C) | 0 | 0 | 2 |
| | p.G12A/V/D complex mutation | 0 | 0 | 1 |

N number of neoplasms.

1285

1286

cell carcinomas "KIRP" TCGA cohort] is likely to represent a random rather than a recurrent event. This finding was also supported by all studies of the PRNRP that included PRCC in their control group, which have shown a negative KRAS mutational status (wild-type). In the same notion, a significant proportion of the low-grade oncocytic tumor, eosinophilic vacuolated tumor, eosinophilic solid and cystic renal cell carcinoma, renal cell carcinoma with leiomyomatous stroma, and acquired cystic disease-associated renal cell carcinoma share TSC/MTOR pathways mutations at a rate that far exceeds the "shared" KRAS mutational rate between PRNRP and any other tumor described thus far, including PRCC; however, all of the aforementioned tumors are considered different entities. Lastly, in our previous study, we performed unsupervised mRNA clustering of 17 tumors obtained from the Cancer Genome Atlas (3 PRNRP, the 2 KRAS mutated PRCCs, and 12 randomly selected PRCCs) and those showed different expression profiles of PRNRP when compared to other tumors

In summary, the presence of *KRAS* mutations in small, clinically undetectable (microscopic) neoplasms suggests a distinct oncogenic pathway for PRNRP as these mutations appear to be an early events in their genesis.

REFERENCES

- Al-Obaidy, K. I. et al. Papillary renal neoplasm with reverse polarity: a morphologic, immunohistochemical, and molecular study. *Am. J. Surg. Pathol.* 43, 1099– 1111 (2019).
- Al-Obaidy, K. I. et al. Recurrent KRAS mutations in papillary renal neoplasm with reverse polarity. Mod. Pathol 33, 1157–1164 (2020).
- Trpkov, K. et al. New developments in existing WHO entities and evolving molecular concepts: the Genitourinary Pathology Society (GUPS) update on renal neoplasia. *Mod Pathol* 34, 1392–1424 (2021).
- Kunju, L. P., Wojno, K., Wolf, J. S. Jr, Cheng, L. & Shah, R. B. Papillary renal cell carcinoma with oncocytic cells and nonoverlapping low grade nuclei: expanding the morphologic spectrum with emphasis on clinicopathologic, immunohistochemical and molecular features. *Hum. Pathol* **39**, 96–101 (2008).
- Saleeb, R. M. et al. Toward biological subtyping of papillary renal cell carcinoma with clinical implications through histologic, immunohistochemical, and molecular analysis. *Am. J. Surg. Pathol.* **41**, 1618–1629 (2017).
- Chang, H. Y. et al. Clinicopathological and molecular characterisation of papillary renal neoplasm with reverse polarity and its renal papillary adenoma analogue. *Histopathology* 78, 1019–1031 (2021).
- Kim, S. S. et al. Recurrent *KRAS* mutations identified in papillary renal neoplasm with reverse polarity-a comparative study with papillary renal cell carcinoma. *Mod Pathol* 33, 690–699 (2020).
- Kiyozawa, D. et al. Morphological, immunohistochemical, and genomic analyses of papillary renal neoplasm with reverse polarity. *Hum Pathol* **112**, 48–58 (2021).
- Lee, H. J. et al. Unilateral synchronous papillary renal neoplasm with reverse polarity and clear cell renal cell carcinoma: a case report with *KRAS* and *PIK3CA* mutations. *Diagn. Pathol* 15, 123 (2020).
- Wei, S. et al. Papillary renal neoplasm with reverse polarity is often cystic: report of 7 cases and review of 93 cases in the literature. *Am. J. Surg. Pathol.* https://doi. org/10.1097/PAS.00000000001773 (2021). Online ahead of print.
- Zhou, L. et al. Papillary renal neoplasm with reverse polarity: a clinicopathologic study of 7 cases. Int. J. Surg. Pathol. 28, 728–734 (2020).
- Tong, K. et al. Frequent KRAS mutations in oncocytic papillary renal neoplasm with inverted nuclei. *Histopathology* 76, 1070–1083 (2020).
- 13. Brunelli, M., Eble, J. N., Zhang, S., Martignoni, G. & Cheng, L. Gains of chromosomes 7, 17, 12, 16, and 20 and loss of Y occur early in the evolution of papillary

renal cell neoplasia: a fluorescent in situ hybridization study. *Mod Pathol* 16, 1053–1059 (2003).

- Calio, A., Warfel, K. A. & Eble, J. N. Papillary adenomas and other small epithelial tumors in the kidney: an autopsy study. Am. J. Surg. Pathol. 43, 277–287 (2019).
- Al-Obaidy, K. I. et al. Recurrent *KRAS* mutation is an early event in the development of papillary renal neoplasm with reverse polarity (abstract 1865). *Mod Pathol* 33, 1002–1163 (2020).
- Mantilla, J. G., Antic, T. & Tretiakova, M. GATA3 as a valuable marker to distinguish clear cell papillary renal cell carcinomas from morphologic mimics. *Hum Pathol* 66, 152–158 (2017).
- Priemer, D. S., Vortmeyer, A. O., Zhang, S., Chang, H. Y., Curless, K. L. & Cheng, L. Activating *KRAS* mutations in arteriovenous malformations of the brain: frequency and clinicopathologic correlation. *Hum Pathol* **89**, 33–39 (2019).
- Thoenes, W., Storkel, S. & Rumpelt, H. J. Human chromophobe cell renal carcinoma. Virchows Arch. B Cell Pathol. Incl. Mol. Pathol. 48, 207–217 (1985).
- Pivovarcikova, K. et al. Renal cell carcinomas with tubulopapillary architecture and oncocytic cells: Molecular analysis of 39 difficult tumors to classify. *Ann. Diagn. Pathol.* 52, 151734 (2021).
- Koski, T. A. et al. Array comparative genomic hybridization identifies a distinct DNA copy number profile in renal cell cancer associated with hereditary leiomyomatosis and renal cell cancer. *Genes Chromosomes Cancer* 48, 544–551 (2009).
- 21. Pivovarcikova, K. et al. Fumarate hydratase deficient renal cell carcinoma and fumarate hydratase deficient-like renal cell carcinoma: Morphologic comparative study of 23 genetically tested cases. *Cesk Patol* **55**, 244–249 (2019).
- Chen, N. et al. Gains of chromosomes 7 and 17 in tubulocystic carcinoma of kidney: two cases with fluorescence in situ hybridisation analysis. *J. Clin. Pathol.* 67, 1006–1009 (2014).
- Sarungbam, J. et al. Tubulocystic renal cell carcinoma: a distinct clinicopathologic entity with a characteristic genomic profile. *Mod. Pathol* 32, 701–709 (2019).

AUTHOR CONTRIBUTIONS

K.I.A., J.N.E., D.J.G., and L.C. performed study concept, methodology, design and writing; R.M.S, T.K., S.R.W., A.R.S, O.H., R.M., A.C., A.M.A, Z.I.A, O.H., N.G., A.O.O., W.A.S. J.N.E., D.J.G., and L.C. provided case material, K.I.A and A.A. provided acquisition of data. K.I.A., R.M.S, T.K., J.N.E., and L.C. participated in manuscript drafting. J.D.S and L.A.B. provided technical and material support. All authors reviewed and approved the manuscript.

COMPETING INTERESTS

The authors declare no competing interests.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

This study was approved by the Institutional Review Board (IRB) at Indiana University, Indianapolis, USA, under protocol number: 1301010350. Ethical approval for this study was waived.

ADDITIONAL INFORMATION

Correspondence and requests for materials should be addressed to Liang Cheng.

Reprints and permission information is available at http://www.nature.com/ reprints

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.