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Clear Cell Papillary Renal Cell Carcinoma and Renal Angiomyoadenomatous Tumor – Two Variants of a Morphologic, Immunohistochemical and Genetic Distinct Entity of Renal Cell Carcinoma

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Abstract

Clear cell papillary renal cell carcinoma (ccpRCC) and renal angiomyoadenomatous tumor (RAT) share morphologic similarities with clear cell (ccRCC) and papillary renal cell carcinoma (pRCC). It is a matter of controversy whether their morphologic, immunophenotypic and molecular features allow the definition of a separate renal carcinoma entity. The aim of our project was to investigate specific renal immunohistochemical biomarkers involved in the hypoxia-inducible factor pathway and mutations in the *VHL* gene to clarify the relationship between ccpRCC and RAT. We investigated 28 ccpRCC and 9 RAT samples by immunohistochemistry using 25

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Disclosure/conflict of interest

JVB is co-inventor on KIM-1 patents, which have been assigned to Partners Healthcare and licensed by Partners Healthcare to a number of companies. All other authors have no conflict of interest.

markers. *VHL* gene mutations and allele losses were investigated by Sanger sequencing and fluorescence in situ hybridization (FISH). Clinical follow-up data were obtained for a subset of the patients. No tumor recurrence or tumor-related death was observed in any of the patients. Immunohistochemistry and molecular analyses led to the reclassification of three tumors as ccRCC and TFE3 translocation carcinomas. The immunohistochemical profile of ccpRCC and RAT samples was very similar but not identical, differing from both ccRCC and pRCC. Especially, the parafibromin and hKIM-1 expression exhibited differences in ccpRCC/RAT compared with ccRCC and pRCC. Genetic analysis revealed VHL mutations in 2/27 (7%) and 1/7 (14%) ccpRCC and RAT samples, respectively. FISH analysis disclosed a 3p loss in 2/20 (10%) ccpRCC samples. ccpRCC and RAT have a specific morphologic and immunohistochemical profile but they share similarities with the more aggressive renal tumors. Based on our results, we regard ccpRCC/RAT as a distinct entity of renal cell carcinomas.

Keywords

kidney; clear cell papillary renal cell cancer; cytokeratin 7; VHL; renal angiomyoadenomatous tumor; clear cell renal cell carcinoma

Introduction

Clear cell papillary renal cell carcinoma (ccpRCC) has been proposed as a new entity of renal cell cancer by the International Society of Urological Pathology (ISUP) to be included in the next World Health Organization Classification of Renal Tumors (1). It was initially discovered in kidneys with end-stage renal disease in 2006 (2). Since then more than 100 ccpRCC cases have been described and the majority were found in normal functioning kidneys (3–8). They are characterized by tumor cells with clear cytoplasm, linear arrangement of low-grade nuclei located apically distant from the basal membrane and containing varying amounts of tubular, papillary and cystic architecture. Strikingly, the ccpRCC lack mitoses, atypia, pleomorphism, necrosis, hyaline globules, foamy macrophages and vascular invasion. Despite significant morphologic, immunohistochemical and genetic similarities to clear cell renal cell carcinoma (ccRCC) and papillary renal cell carcinoma (pRCC), characteristic genetic differences include *VHL* gene mutations and 3p losses, found in ccRCC. Gain of the chromosomes 7 and 17 or loss of chromosome Y, are absent or extremely rare in ccpRCC cases (4, 9, 10). No disease defining mutation has been identified to date.

The renal angiomyoadenomatous tumor (RAT) was first reported in the kidney of a 93-year-old male by Michal et al. (11). Nine years later, the same group characterized 5 additional tumors (12). Verine pointed out that ccpRCCs are a major differential diagnosis of RAT and emphasized their morphologic, immunohistochemical, molecular and clinical similarities (13). In the literature many terms have been used to probably describe the same entity, including ccRCC with prominent leiomyomatous proliferation and renal cell carcinoma with smooth muscle stroma (12, 14–17).

The epithelial component of ccpRCC and RAT is composed of cells with abundant clear cytoplasm, strong diffuse CK7 activity and low grade nuclei (Fuhrman grade 1 and 2). Due

to their many similarities, several authors regard ccpRCC and RAT to be a variant of the same entity (6, 9, 13, 17, 18).

The aims of our study were to clarify the relationship between ccpRCC and RAT and to identify markers to reliably distinguish ccpRCC and RAT from the biologically more aggressive renal neoplasms.

Materials and Methods

Case Cohort

All tumors were consultation cases from HM and EC and were received from Austria, France, Germany, Italy and Switzerland. Hematoxylin and eosin-stained slides were reviewed for morphologic features of clear cell papillary renal cell carcinoma (ccpRCC) and renal angiomyoadenomatous tumor (RAT) as previously described (3, 5–7, 11, 12, 19). Diagnostic features of ccpRCC include tumor cells with abundant clear cytoplasm, varying papillary, cystic and tubular architecture, low grade nuclei (Fuhrman grade 1 and 2) located apically distant from the basal membrane, and strong diffuse CK7 and CA-IX expression. For diagnosis of RAT, the following criteria were required: cells with clear cytoplasm, low grade nuclei (Fuhrman grade 1 and 2) embedded in a smooth muscle stroma and strong diffuse CK7 staining of the epithelial component. Tumors were staged according to the TNM system (20) and graded according to Fuhrman et al. (21). The morphologic characteristics were scored as previously described (7).

Immunohistochemistry

A total of 25 antibodies were selected as (i) they are involved in the *VHL* signaling pathway, (ii) they are known to be prognostic biomarkers of ccRCC and (iii) they have been reported as markers of ccpRCCs and RATs in a small group of ccpRCCs described in recent USCAP meetings (2011–2014). TMA sections (2.5 μm) were transferred to glass slides and treated using Ventana Benchmark XT, Bond-max (Leica Microsystems) automated systems, as well as manual protocols. TMA construction was not possible in five of the ccpRCC cases due to absence of tissue. The immunohistochemical staining product was described as nuclear, membranous or cytoplasmic (Table 2). The immunohistochemistry results were interpreted as 0 (negative), 1+ (weak staining), 2+ (moderate staining) and 3+ (strong staining). For statistical analysis all 2+ and 3+ stainings were defined as positive, 0 and 1+ as negative. Antibodies and protocols are listed in Table 2.

Fluorescence In Situ Hybridization

FISH, performed to detect *VHL* allele losses, was carried out using the ZytoLight ® SPEC VHL/CEN 3 Dual Color Probe (ZytoVision, Bremerhaven, Germany). Tissue sections were cut from FFPE blocks, deparaffinized and hybridized as previously described (22). Sixty non-overlapping tumor nuclei from three different areas were analyzed and the number of VHL and CEN3 signals was recorded for each nucleus. The total number of VHL and CEN3 signals as well as the VHL/CEN3 ratio and the percentage of tumor cells with less than 2 VHL signals were calculated. Tumors were considered VHL deleted if more than 50% of the tumor nuclei displayed less than 2 VHL signals (23). In two cases TFE3 FISH using SPEC

TFE3 Dual color break apart probe from ZytoVision were done on whole sections as previously described by our group (24).

VHL Sequencing Analysis

Tumor areas displaying >80% tissue in the epithelial portion of the ccpRCC and RAT were marked on the H&E slides. DNA from FFPE tumor tissue samples was obtained by punching 1–2 tissue cylinders (diameter 0.6mm) from each sample. DNA was extracted from the tumor tissue samples according to the Maxwell 16 FFPE Plus DNA Purification protocol (Promega, Fitchburg, USA) for automated DNA purification. DNA concentrations in the samples were measured using the Nanodrop (Thermo Fisher Scientific, Waltham, MA, USA). PCR of the *VHL* gene was performed as previously described (25) using approximately 40 ng of DNA for each amplification. DNA sequencing was performed with the dideoxy chain-termination method using the BigDye® Terminator v1.1 Cycle Sequencing kit (Applied Biosystems, Foster City, USA). The same forward and reverse primers were used for the PCR and sequencing. Cycle sequencing products were analyzed using the AbiPrism 3100 Genetic analyzer (Applied Biosystems). The obtained sequences were compared with the NCBI sequence AF010238 using NCBIs Blast 2 Sequences. All *VHL* point mutations obtained were validated by a second separate PCR and sequencing analysis.

Results

Clinical and Pathologic Findings

The patients with ccpRCC ranged from 29 to 75 years of age (mean age 58 years) and those with RAT from 32 to 68 years of age (mean age 43.3 years) at the time of nephrectomy. Male to female ratio was 1.5:1 in the ccpRCC group (17 men and 11 women) and 3.5:1 in the RAT group (6 men and 1 woman).

Clinical follow-up data was available for 78% (21/27) of the ccpRCC patients and 71% (5/7) of the RAT patients. Mean follow-up time was 29.7 months (range 7 to 84 months) for the ccpRCC patients and 32.3 months (range 25 to 38 months) for the RAT patients. There was no evidence of recurrence or disease-related death in any of the patients. None of the RAT (0/5) patients and 14 % (3/22) of the ccpRCC patients had end-stage renal disease.

In the RAT group, the average diameter of the tumor was 3.1 cm (range 1.8–5.0 cm) compared to 2.6 cm (range 0.5–8 cm) in the ccpRCC group. 67% (4/6) of the RAT patients displayed pathologic stage pT1a and 33% (2/6) stage pT1b. Overall 86% of the tumors (6/7) were Fuhrman nuclear grade 1 and 14% (1/7) were nuclear grade 2. In the ccpRCC cases 77% (20/26) were stage pT1a, 19% (5/26) were pT1b and 4% (1/26) were pT2a. Fuhrman nuclear grade 1 was found in 48% (13/27) and nuclear grade 2 in 52% (14/27) of the tumors. All the ccpRCCs and 6/7 RATs showed at least focal papillary architecture and branched ducts. In contrast to ccpRCC, secretory cells were completely absent in the RAT cases. Both showed variable amounts of cystic areas. All tumors were characterized by absence of mitotic formations, foamy macrophages, calcifications and vascular invasion.

Table 1 summarizes the clinicopathological findings. Morphological characteristics are shown in Table 5.

Immunohistochemical Findings

The immunohistochemical findings are detailed in Table 3. ccpRCC and RAT were strongly positive for CK7, CK19, CA-IX, GLUT-1, E-cadherin, vimentin, β-catenin, parafibromin, PAX-2, PAX-8, p27, p53 and c-MET. Staining for GLUT-1 (p = 0.0572), CD 70 (p = 0.1499) and p16 (p = 0.3702) differed slightly in the RAT samples compared with ccpRCCs, although differences did not show statistical significance. Following the recent results by Cui et al. (26), Aron et al. (53) and Schwartz et al. (56), we tested parafibromin, hKIM-1 (27) and CD133 expression to distinguish ccpRCC/RAT from ccRCC/pRCC. As shown in Table 4, the expression difference reached statistical significance (p < 0.0001). The biomarkers CD70 (28), MET (29) and E-cadherin (30) were able to distinguish between ccpRCC/RAT and ccRCC (p <0.0001). Furthermore, the hKIM-1 and parafibromin were able to distinguish between ccpRCC/RAT and pRCC. All ccpRCC cases exhibited a characteristic CA-IX "cup-like" distribution, sparing the luminal border as it has been described in the literature before (6, 31). In contrast, the RAT tumors and the ccpRCC-like tumor with the VHL mutation showed a circumferential membranous staining pattern. Two RAT samples stained weakly positive for TFE3 and were, therefore, further analyzed by HMB45 and TFEB. Both stainings revealed a negative result. Additionally, TFE3 FISH was performed (see below).

Fluorescence In Situ Hybridization Findings

Three deletions of the short arm of chromosome 3 were identified. All of them occurred in the ccpRCC cases (3/21, 14%) and no deletion was found in the RAT cases (0/7, 0%). The presence of the 3p deletions in the two ccRCC controls were correctly identified. In 9 of the cases FISH was not performed as there was not sufficient tissue after VHL mutation analysis and immunohistochemistry.

TFE3 FISH was performed with the two above-mentioned RAT-like cases that showed weak TFE3 expression. One case showed the typical break apart pattern in >85% of the cells, while the second case was negative. Both cases were reclassified as translocation carcinomas due to immunohistochemical TFE3 positivity.

VHL Gene Mutation Analysis

Three *VHL* mutations were detected in the ccpRCC group (3/27, 11%) in exon 2 (c.351G>C/p.Trp117Cys, c.461C>T/p.Pro154Leu, c.388G>C/p.Val130Leu) and one in the RAT group (1/7, 14%) in exon 1 (c.174_177delGCCG /p.Pro59GlyfsX7). We identified two cases, harboring both a *VHL* mutation and 3p loss. One case showed a 3p loss but no *VHL* mutation and two cases with a *VHL* mutation showed no 3p loss.

Discussion

In the present study we have sequenced the largest number of ccpRCC (27) and RAT (7) cases to date. We found a *VHL* mutation rate of 11% in ccpRCC and 14% in RAT.

Furthermore, we analyzed hypoxia-inducible factor pathway-related proteins to compare these findings with recent findings.

ccpRCC and RAT are currently underrecognized. Recent studies have revealed that they are not rare (7, 32, 33) and that among all RCC the ccpRCC have a prevalence rate between 1.2% and 4.1%, thus representing up to 4,500 new cases of renal cancer in the United States annually (7, 32, 34). Awareness of its morphologic and immunohistochemical features is imperative for a correct classification. In a recent publication Gill et al. underscored the necessity of reclassifying low grade and low stage ccRCC as up to 7% of the cases are in fact ccpRCC (33).

Morphologically, ccpRCC and RAT share many features. Their epithelial component is composed of cells with clear cytoplasm and low grade nuclei. Both tumors have various amounts of smooth muscle stroma, and their epithelial component is characterized by either cystic or papillary architecture. In our cohort the majority of the RAT samples had focal papillary features of the epithelial component, which are typically diffuse CK7 and CA-IX positive. The most relevant differential diagnoses include ccRCC that exhibit papillary features, pRCC exhibiting clear cell characteristics and Xp11 translocation carcinoma. In our cohort, two cases initially classified as RAT had to be re-classified as Xp11 translocation carcinomas after immunohistochemistry and TFE3 FISH analysis. The translocation carcinomas were identified by nuclear TFE3 protein expression. Only one case showed a positive TFE3 FISH result. It is controversial whether TFE3 positivity is sufficient to diagnose TFE3 translocation carcinoma (24, 35), but, from these 2 cases, we concluded that TFE3 translocation cancer falls within the differential diagnostic spectrum of ccpRCC/ RAT. Another differential diagnosis for the case with weak TFE3 staining and negative FISH is TFEB-associated RCC. Those tumors can overlap tremendously with the TFE3 rearranged RCC (36, 37). To rule out this differential diagnosis we did two additional immunohistochemical stainings (HMB45 and TFEB). Both stainings showed a negative result making that differential diagnosis unlikely.

One ccpRCC case was reclassified as ccRCC. That case exhibited typical ccpRCC morphology but was completely negative for CK7 and strongly positive for hKIM-1. This case also revealed a mutation in the *VHL* gene and a 3p loss in the FISH analysis. These findings highlight the importance of molecular testing and should raise awareness of ccpRCC mimicking ccRCC (38).

VHL gene mutations are the genetic hallmark of ccRCC. Initially, it was reported that VHL alterations are absent in ccpRCC. However, three groups have recently identified VHL mutations in ccpRCC at frequencies varying from 15% to 27% (39–41). In concordance with these studies, we also identified VHL gene alterations in ccpRCC, but the prevalence of VHL gene mutations is significantly lower than in ccRCC (42–44). The discrepancy between the number of mutations found in our ccpRCC cases and that reported may be explained by the different detection methods employed, including single nucleotide polymorphism (SNPs) genotyping array, Sanger sequencing, and by the limited number of cases in previous studies. Alternatively, cases with VHL mutations could represent ccRCCs with morphology and immunoprofile which closely mimics that of clear ccpRCC and RAT tumor. Currently,

ccpRCC are diagnosed on the basis of morphology and diffuse strong CK7 expression. The absence of VHL mutations/3p deletions is not diagnostic for ccpRCC. Therefore we suggest to diagnose tumors with diffuse CK7 expression combined with the typical morphology as ccpRCC. In previous studies CCRCC with diffuse CK7 profile have had a completely different prognosis than CCRCC without that CK expression pattern (45). These previous findings justify such an approach. *VHL* inactivation leads to a HIF-dependent CA-IX and GLUT-1 up-regulation. We only found few *VHL* mutations, but in combination with CA-IX and GLUT-1 immunoreactivity in both ccpRCC and RAT. This clearly sets the ccpRCC and RAT apart from ccRCC, which shows *VHL* mutations in up to 80% of the cases (44, 46). Therefore, we believe that the HIF pathway may be activated in a *VHL*-independent manner in most ccpRCCs and RATs, also hypothesized by Rohan et al. (6).

Recently, Lawrie et al. found various mutations in ccpRCC by using Next Generation Sequencing (NGS), including a non-synonymous T992I mutation in the *MET* proto-oncogene (47). This gene was originally described as causing hereditary pRCC (48). Interestingly, Lawrie et al. detected no *VHL* mutation, but found overexpression in all five members of the miR-200 family. The miR-200 family plays an essential role in tumor suppression by inhibiting epithelial-mesenchymal transition (49). To support Lawrie's results we also noted immunoreactivity for E-cadherin and β -catenin. These findings suggest that epithelial-to-mesenchymal transition (EMT) may be incomplete or blocked in ccpRCC contributing to their indolent course (47).

Other genetic alterations characteristic for pRCC include gain of the chromosome 7 and loss of the chromosome Y. However, in ccpRCC, gain of the chromosome 7 have very rarely been reported (4, 5, 9, 10) and no loss of the chromosome Y has been observed to date. Fisher et al. found a unique gene expression profile of ccpRCC when investigating 8 different genes, with only some expression levels comparable with those observed in ccRCC and pRCC (50).

In our FISH analysis, we identified three chromosome 3p deletions in 20 ccpRCC and in 7 RAT samples. All 3p deletions occurred in ccpRCC with a frequency of 14.3 %, but none was detected in RAT. To date, only 4 cases with a 3p loss have been reported in ccpRCC (34, 41). Interestingly, the single case described by Martignoni et al. (41) concurrently harbored a VHL mutation like two of our three cases with a 3p loss. Shi et al. (34) also used FISH and observed monosomy of chromosome 3 in three cases in a series of 11 ccpRCC all lacking mutations in the VHL gene. In 2009, Shannon et al. (14) published a study on 5 ccRCC with smooth muscle stroma and found loss of the entire chromosome 3 in two cases and a 3p loss in one case using FISH. In contrast, Martignoni found no 3p loss in a series of three cases of ccRCC with smooth muscle stroma (17). Given these molecular findings, it has been suggested that RAT and ccRCC with smooth muscle stroma are interchangeable terms (51). However, some of the cases of ccRCC with smooth muscle stroma, particularly those that showed 3p loss, might represent ccRCCs with exuberant, infiltrative smooth muscle, whereas the others might in fact be RAT tumors, particularly the ones that do not show 3p loss (15). Additionally, recent data shows that some tumors with RAT morphology and immunophenotype share a common mutation in the TCEB1 gene which inactivated the VHL pathway and upregulated proteins along the hypoxia-inducible pathway (52). Twenty-

five different antibodies were used to characterize ccpRCC and RAT. We were particularly interested in hypoxia-inducible factor pathway-related proteins as well as other antibodies, which were reportedly used in small series of ccpRCC cases in the 2011 and 2014 USCAP meetings. This gave us the opportunity to compare immunohistochemical findings in ccpRCC and RAT to clarify their interrelationship. Remarkably, there were no statistically significant differences in the staining properties in any of the antibodies in ccpRCC compared to RAT.

Parafibromin and hKIM-1 expression levels differed significantly between ccpRCC/RAT and ccRCC/pRCC. Cui et al. (26) recently demonstrated that parafibromin can be very helpful in differentiating ccpRCC from ccRCC and pRCC. In a study by Aron et al. (53), the difference in the staining positivity rate of ccpRCC and ccRCC was even more striking compared with our study. In addition to parafibromin and hKIM-1 expression, CD70 also proved to be a useful marker in differentiating ccpRCC from ccRCC, since CD70 expression is rare in ccpRCC and very frequent in ccRCC. CD70 was used for immunohistochemistry because we have previously demonstrated that CD70 is a potential biomarker for ccRCC (28, 54). The importance of immunohistochemical stainings in the correct identification of true ccpRCC was also highlighted by Williamson et al. (55). They studied 14 ccpRCC-like tumors, which could not be distinguished from ccpRCC morphologically, but which showed a high 3p deletion frequency (82%) and showed a different immunohistochemical profile, with negative or localized CK7 staining as the most striking feature. These characteristics also led to a reclassification of one of our tumors, primarily diagnosed as ccpRCC.

Recently, Schwartz et al. studied different stem cell markers in renal cancers. They reported a 90 % positivity rate for OCT 3/4 in a series of 10 ccpRCC samples (56). This finding is discrepant to our positivity rate of 8.7%, which may be due to the use of different antibodies or immunohistochemical protocols. However, similarly to Schwartz et al. (56), we also detected a high positivity rate of stem cell marker CD133 (81.8% and 100%, respectively) in ccpRCC. Interestingly, Schwartz et al. reported a CD133 positivity rate of only 14% in ccRCC. It can therefore be concluded that CD133 is an additional tool to distinguish ccRCC from ccpRCC/RAT.

In concordance with Munari et al. (57), we found that about one third of ccpRCC are positive for GATA3, a protein crucial for the regulation of Th2 development and function. However, given that only a moderate staining intensity was seen in no more than 10 % of the tumor cells, we do not consider OCT3/4 and GATA3 as diagnostic tools to differentiate ccpRCC from ccRCC.

No previous studies have reported cancer-related death, vascular invasion or metastasis in ccpRCC (4–7, 51, 58), suggesting that the disease follows an indolent course. Benign biologic behavior was also observed in all RAT cases (12, 14, 59). This is comparable to multilocular cystic RCC, which has an excellent prognosis with no disease recurrence after surgery (7, 12, 59). Specific molecular alterations may account for the indolent course of multilocular cystic RCC. Proposals have been put forward to rename multilocular cystic RCC as multilocular cystic renal cell neoplasm of low malignant potential to underscore this specific biologic behavior (1). Our group has reported that the expression of p27, CA-IX,

CK7 and CK19 is associated with a better prognosis in sporadic RCC (45, 60). Interestingly, our ccpRCC/RAT cases stained strongly positive for all of these markers. Hence, the indolent clinical course of ccpRCC/RAT might in part be due to this specific signaling pathway. However, some of our low grade ccRCC included in our previous publications may in fact be unrecognized ccpRCC (33, 45).

In summary, we have demonstrated that ccpRCC and RAT cannot be distinguished from one another by immunohistochemistry and molecular analyses and both follow a benign clinical course. We regard them as a spectrum of a distinct tumor entity. Precise diagnosis is crucial since it has an excellent prognosis. Given the reliability of TFE3 immunohistochemistry, TFE3 FISH should be performed in cases with equivocal TFE3 immunohistochemistry (35). Taking into account the controversial relevance of the *VHL* mutation analysis in this differential diagnosis, direct *VHL* sequencing is not helpful in separation of ccRCC with prominent smooth muscle stroma from RAT. Our results suggest that a panel of antibodies against CK7, parafibromin and MET are a helpful tool to differentiate most ccpRCC/RAT from other renal tumors (Table 4). In some difficult cases *VHL* mutation testing and TFE3 FISH analysis are helpful tools to distinguish ccRCC and TFE3 translocation carcinoma from ccpRCC/RAT (Figure 5).

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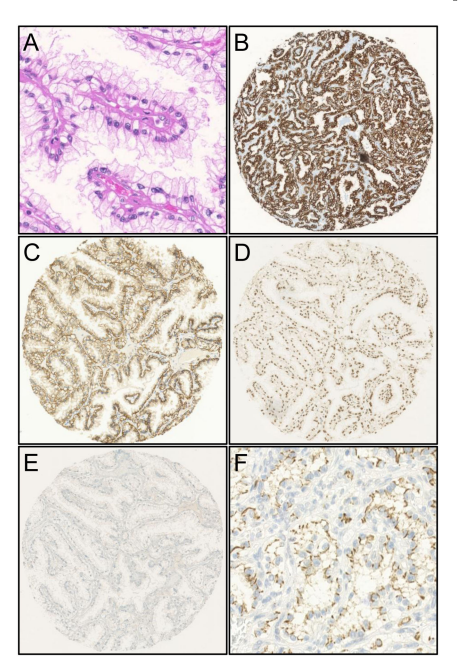


Figure 1. Clear cell papillary renal cell carcinoma. H&E stain (A) and typical immunohistochemical profile with diffuse membranous CK7 positivity (B), membranous "cup-like" CA-IX positivity (C), nuclear parafibromin positivity (D), hKIM-1 negativity (E) and CD133 positivity (F).

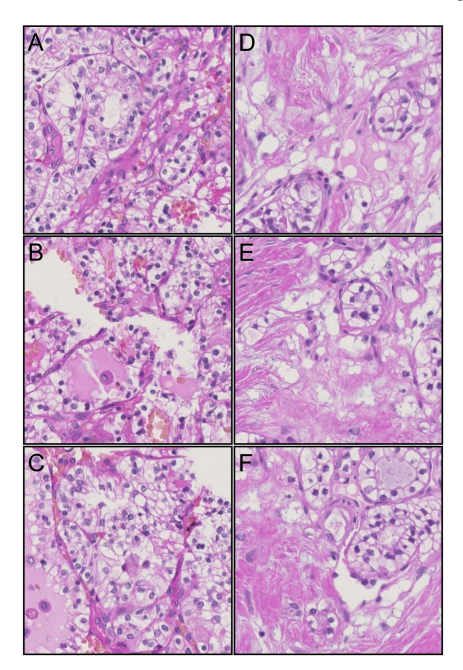


Figure 2. H&E morphology of a ccpRCC-like (A-C) and a RAT-like (D-F) case. Diagnostic features of ccpRCC include tumor cells with abundant clear cytoplasm, varying papillary, cystic and tubular architecture and low grade nuclei (Fuhrman grade 1 and 2) located apically distant from the basal membrane. The epithelial part of RAT tumors is composed of cells with clear cytoplasm, low grade nuclei (Fuhrman grade 1 and 2) embedded in a smooth muscle stroma.

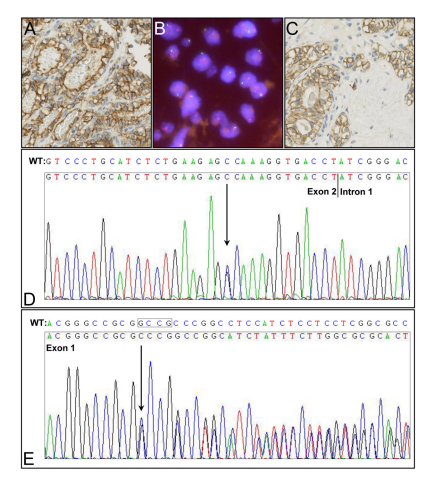


Figure 3.

Molecular features of a ccpRCC-like (A) and a RAT-like (C) case both exhibiting a circumferential CA-IX staining pattern, harboring a *VHL* mutation (D: c.174_177delGCCG/p.Pro59GlyfsX7; E: c.351G>C/p.Trp117Cys) and a 3p deletion detected by fluorescence in situ hybridization (B). The mutation sites are denoted by an arrow. The boundaries between exon and intron are indicated. The upper base pair letter sequence shows the wild type (WT) sequence (D, E). Tumor cells harbor only one VHL (green) signal and two CEN3 copies (orange) (B).

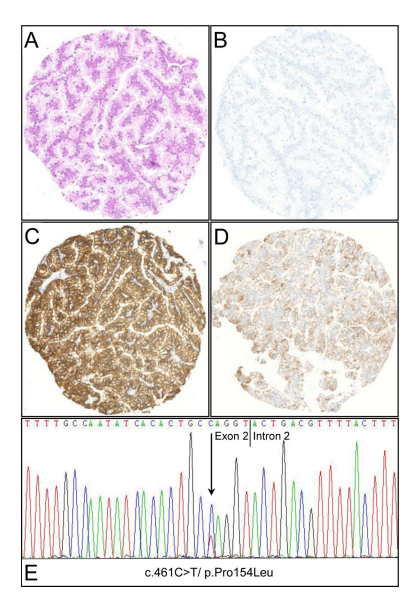


Figure 4. ccpRCC look-alike showing classic ccpRCC morphology on the HE stain (A), however with a typical ccRCC immunohistochemical profile showing CK7 negativity (B), CA-IX positivity (C), hKIM-1 positivity (D) and proof of *VHL* mutation (E).

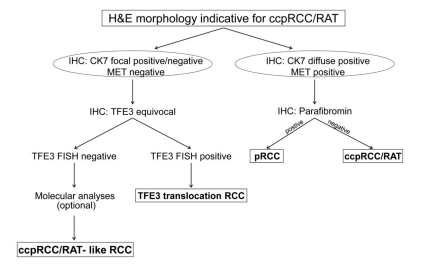


Figure 5.Proposed diagnostic workflow for renal cell carcinoma showing H&E features indicative for ccpRCC/RAT.

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Table 1

Clinicopathological findings

IBD, papillary RCC type 1, kidney adenomas Large cell b-cell lymphoma papillary RCC type 2 Special remarks IgA nephritis Metastasis ou na na na no na ou na ou na na na na ou na ou na ou ou ou ou ou ou ou Tumor Recurrence ou na na na ou na ou na ou na na na na ou na ou na ou ou ou ou ou ou ou Follow-up (months) 19 12 9 na na 35 25 na na 39 25 na na na na 24 na Ξ 84 67 na ∞ 71 Tumor-related death ou ou na ou na ou na ou ou na ou ou ou ou ou ou na na na ou na na na ou ESRD ou na na na ou na ou na ou na na yes na ou na ou na ou ou ou ou ou yes ou pT1bpT1bpT1apT1bpT1a pT1a pT1a pT1apT1bpT1a pT1apT1a pT2a pT1bpT1a pT1b pT1a pT1a pT1a pT1a Stage pT1a pT1a na na Number of Tumors na na Size (cm) 1.8 4.7 1.8 3.5 6.0 4.5 3.8 1.3 5.1 2.2 1.3 2 na S na 7 4 α ∞ Grade (Fuhrman) 7 7 7 Laterality right right right right right right right left right left left left left left left left left na na left na left left na Gender Σ Σ Σ Σ Σ Σ Σ Σ Σ Ľ Σ Σ Σ Σ Σ Σ Σ Σ Σ [I, [I, ſĽ, Ľ, Ľ Age 36 CCPRCC 32 32 34 45 99 89 55 28 57 50 38 63 62 31 55 89 51 75 38 53 51 62 52 RAT 12 13 9 6 9

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F right 2.5 1 pT1a no no 39 no no <th< th=""><th>Age</th><th>- 0</th><th></th><th>Gender Laterality</th><th>Grade (Fuhrman)</th><th>Size (cm)</th><th>Number of Tumors</th><th>Stage</th><th>ESRD</th><th>Tumor- related death</th><th>Follow-up (months)</th><th>Tumor Recurrence</th><th>Metastasis</th><th>Special remarks</th></th<>	Age	- 0		Gender Laterality	Grade (Fuhrman)	Size (cm)	Number of Tumors	Stage	ESRD	Tumor- related death	Follow-up (months)	Tumor Recurrence	Metastasis	Special remarks
M left 1.5 TI pTIa no no 37 no no M na 1 0.5 2 pTIa no 29 no no M right 1 2 1 pTIa no no 22 no no F right 1 2 1 pTIa no no 17 no no F left 1 2.8 1 pTIa no no 15 no no F na 1 0.5 1 pTIa no no 15 no no m left 2 1.8 pTIa no no 14 no no m left 2 1.8 pTIa no no no no no no	71		Ħ	right	2	2.5	1	pT1a	ou	ou	39	no	ou	
M na 1 0.5 2 pTla no no 29 no no M right 1 2 1 pTla no no 22 no no F M right 1 2 1 pTla no no 1 no no F left 1 2.8 1 pTla no no 15 no no m left 2 1.8 1 pTla no no 14 no no m left 2 1.8 1 pTla no no 12 no	57	,	F	left	1	1.5	1	pT1a	no	no	37	no	no	
M left 2 7 pT1b no no 22 no M right 1 2 1 pT1a no no 17 no F right 1 3 pT1a no no 17 no F left 1 2.8 1 pT1a no no 14 no m left 2 1.8 pT1a yes no 12 no m left 2 1 pT1a no no no no	72		M	na	1	0.5	2	pT1a	ou	no	29	ou	ou	
M right 1 2 1 pT1a no no 21 no F right 1 3 pT1a no no 17 no F left 1 2.8 1 pT1a no 15 no m left 2 1.8 1 pT1a yes no no m left 2 2 1 pT1a no no no no	61		F	left	2	2	1	pT1b	ou	no	22	ou	ou	
F right 1 3 pTla no no 17 no F left 1 2.8 1 pTla no 15 no no m left 2 1.8 1 pTla yes no 12 no m left 2 1 pTla no no 10 no no	71		M	right	1	2	1	pT1a	no	no	21	no	no	
F left 1 2.8 1 pT1a no no 15 no no 15 no n	53	3	F	right	1	1	3	pT1a	no	no	17	no	no	
F na 1 0.5 1 pT1a no no 14 no m left 2 1.8 1 pT1a yes no 12 no m left 2 2 1 pT1a no no no no	70		F	left	1	2.8	1	pT1a	ou	no	15	ou	ou	
m left 2 1.8 1 pTia yes no 12 no m left 2 2 1 pTia no no no no	65	2	F	na	1	0.5	1	pT1a	no	no	14	no	no	
m left 2 2 1 pTla no no 10 no	74	+	m	left	2	1.8	1	pT1a	yes	no	12	no	no	
	54	—	ш	left	2	2	1	pT1a	no	no	10	ou	ou	

Abbreviations: ESRD: end-stage renal disease; n. a.: not available; RAT: renal angiomyoadenomatous tumor; IBD: inflammatory bowel disease; ccpRCC: clear cell papillary renal cell carcinoma

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Antibody Overview

Table 2

Antibody	Clone	Species	Vendor	Dilution	Staining Pattern
β-catenin	14/Beta-Catenin	Mouse	BD Biosciences	1:50	membranous
Carbonic anhydrase IX	:	Rabbit	Abcam Limited	1:6000	membranous (partially cytoplasmatic)
c-MET	SP44	Rabbit	Ventana	prediluted	membranous
CD10	SP67	Mouse	Ventana	prediluted	membranous
CD70	301731	Mouse	R&D Systems	1:75	membranous
CD133	:	Rabbit	Abcam Limited	1:500	membranous
Cytokeratin 7	OV-TL 12/30	Mouse	DAKO A/S	1:100	membranous
Cytokeratin 19	RCK108	Mouse	Abcam Limited	1:200	membranous
E-cadherin	EP700Y	Rabbit	Cell Marque Lifescreen Ltd.	1:200	membranous
Estrogen receptor	SPI	Rabbit	Labvision	1:50	nuclear
GATA-3	L50-823	Mouse	Biocare Medical	1:250	nuclear
GLUT-1	-	Rabbit	Millipore Corporation	1:1000	membranous
hKIM-1			Bonventre lab		membranous and cytoplasmatic
Melanosome	HMB-45	Mouse	DAKO A/S	1:50	cytoplasmic
OCT3/4	NINK	Mouse	Novocastra Laboratories Ltd	1:150	nuclear
p16	JC8	Mouse	Santa Cruz Biotechnology, Inc.	1:200	nuclear
p27		Rabbit	Santa Cruz Biotechnology, Inc.	1:30	nuclear
p53	DO-7	Mouse	DAKO A/S	1:80	nuclear
Parafibromin	2H1	Mouse	Santa Cruz Biotechnology, Inc.	1:10	nuclear
PAX-2		Rabbit	ZYMED Laboratories Inc.	1:100	nuclear
PAX-8		Rabbit	Protein Tech Group, Inc	1:200	nuclear
Progesterone receptor	1E2	Rabbit	Ventana	prediluted	nuclear
TFE3	MRQ-37	Rabbit	Cell Marque Lifescreen Ltd.	prediluted	nuclear
TFEB	-	Goat	Santa Cruz Biotechnology, Inc.	1:100	nuclear
Vimentin	Vim 3B4	Mouse	DAKO A/S	1:250	cytoplasmatic

 Table 3

 Results of Immunohistochemistry. Percentage relates to number of interpretable cases.

Antibody	ccpRCC	RAT
β-catenin	95.5	85.7
Carbonic anhydrase IX	95.5	85.7
CD10	31.8	66.7
CD70	22.7	0
CD133	81.8	100
с-МЕТ	91.3	100
Cytokeratin 7	100	100
Cytokeratin 19	88.9	100
E-cadherin	100	100
Estrogen receptor	4.3	0
GATA-3	31.8	42.9
GLUT-1	95.5	85.7
hKIM-1	23.8	0
OCT3/4	8.7	0
p16	18.2	42.9
p27	100	100
p53	72.7	71.4
Parafibromin	95.5	100
PAX-2	63.6	100
PAX-8	95.5	100
Progesterone receptor	0	0
Vimentin	95.5	100

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Table 4

Different expression patterns of hKIM-1, Parafibromin, CD70, CD133, MET and E-cadherin in ccPRCC, ccRCC and pRCC (own data and literature).

Antibody	ccpRCC	ccRCC	Fisher Exact (P) pRCC	pRCC	Fisher Exact (P)
hKIM-1	27.3% (6/22)	74% (54/73)*	<0.0001	93% (28/30)*	<0.0001
Parafibromin	Parafibromin 91.3% (21/23) 7% $(4/61)^{\#}$	7% (4/61)#	<0.0001	19% (7/37)#	<0.0001
CD70	26.1% (6/23)	78% (197/252)§ <0.0001	<0.0001	32% (113/348)§ 0.6475	0.6475
CD133	78.3% (18/23)	78.3% (18/23) $14.3\% (3/21)^{X}$	<0.0001	53% (8/15) ^x	0.16
MET	91.3% (21/23) 0% (0/96) ^y	$\Lambda(96/0) \%0$	<0.0001	90% (18/20) ^y	1
E-cadherin	100% (22/22)	31.9% (22/69) ²	<0.0001	n. a.	n. a.

n. a.: not available

* Lin et al. 2007 (27) #Cui et al. 2012 (26)

\$\text{\$\text{\$\section}\$} \text{Ruf et al. (28)}

^xSchwartz et al. 2012 (56)

^yChoi et al. 2006 (29)

²Cai 2013 (30)

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Table 5
Morphological characteristics of RAT and ccpRCC

#	Papillary Architecture	Branched Ducts	Secretory Cells	% Cystic
RA	Γ			
1	1f	yes	no	15
2	0	no	no	0
3	2f	yes	no	65
4	1f	yes	no	5
5	2f	yes	no	0
6	1f	yes	no	55
7	2f	yes	no	15
ccpl	RCC			
1	1	yes	yes	85
2	2	yes	no	10
3	3	yes	yes	10
4	3	yes	no	10
5	2	yes	no	15
6	3	yes	yes	55
7	1	yes	yes	0
8	3	yes	no	0
9	3	yes	no	0
10	2	yes	no	5
11	3	yes	no	0
12	3	yes	yes	40
13	1	yes	yes	15
14	1	yes	no	5
15	2	yes	no	20
16	2	yes	no	10
17	1	yes	yes	15
18	3	yes	yes	45
19	2	yes	no	30
20	2	yes	yes	20
21	2	yes	no	5
22	1	yes	no	10
23	3	yes	yes	55
24	1	yes	yes	10
25	3	yes	no	35
26	2	yes	yes	5
27	3	yes	yes	30

f = focal

Scored as previously described (7)