Multilocular cystic clear cell renal cell carcinoma imaging with histopathologic correlation

Emi Marinela Preda1, Ioana Gabriela Lupescu1, Adrian Dijmarescu1, Gelu A. Popa1,3, Mugur C. Grasu1,3, Monica Hortopan2

1Radiology, Medical Imaging and Interventional Radiology Department, Fundeni Clinical Institute, Bucharest, Romania
2Department of Pathologic Anatomy, Fundeni Clinical Institute, Bucharest, Romania
3University of Medicine and Pharmacy «Carol Davila», Bucharest, Romania

ABSTRACT

Renal cell carcinoma (RCC) causes significant morbidity and mortality. Clear cell RCC (ccRCC) is the most common histologic subtype, with worse prognosis compared with other histologic subtypes. The indolent variant of ccRCC is the multilocular cystic RCC (McRCC). Imaging features reflect the various histological findings of each histologic subtypes. Recent advances in imaging technology permit early and more appropriate diagnosis of RCC and facilitate optimal management.

Key words: Multilocular cystic clear cell renal cell carcinoma, renal cell carcinoma, clear cell renal carcinoma, imaging, CT, MRI

INTRODUCTION

Renal cell carcinoma (RCC) is the most common adult renal epithelial cancer, accounting for more than 90% of all renal malignancies and is the most lethal of all urologic cancers (1,2). The most of the cases are diagnosed incidentally during imaging examinations (2-6). In 2004 World Health Organization reviewing renal neoplasms classification, recognizes several distinct histologic subtypes of RCC. Clear cell RCC is the most common subtype and has a less favorable prognosis than papillary RCC and chromophobe RCC. Multilocular cystic RCC is a distinct subtype of clear-cell RCC that appears to have a favorable prognosis. We report a case of McRCC, which is a rare entity, comprising approximately 1 to 2% of all renal tumors and approximately 4% of all ccRCC (although the truth is unknown due to the lack of this subtype before 2004) (3,4,5)

CASE REPORT

Our patient is a men, 65 years old, Known with type II diabetes mellitus in treatment with oral medication, treated hypertension, BRD, prostatic hypertrophy, hospitalized in the Urology Department of Fundeni Clinical Institute,
in order for investigations and treatment for a large right renal cyst diagnosed by ultrasound.

Clinically, the patient had at admission: good general condition, no fever, diuresis present, clear urine and about 10 cm mobile, palpable tumor in right kidney lodge.

Urinary tract ultrasound demonstrated a normal left kidney, half-filled bladder, prostate volume = 24 cc; right kidney with 111 mm cyst of posterior valve, with multiple septa. It is recommended computed tomography examination with iodinated non-ionic contrast.

Computed tomography (CT) revealed a voluminous, well-defined, extrarenal cystic mass (dimensions: 11/8 cm) with predominant fluid component, containing one small expansile dense nodule (about 18/13 mm), with intense enhancement, located just above of some enhanced septa – aspects that are characteristics for a Bosniak cyst type IV. (figure 1). Left kidney with normal CT aspect.

Magnetic Resonance (MR) examination performed in conjunction with CT examination noted expansive mass found in the right kidney with extrarenal development predominantly, having the same dimensions described at CT, with predominant cystic component manifesting water restriction on diffusion sequence and containing a small solid nodule with strong enhancement, located on the anterior wall and several enhancing, irregular septa, which gives a multilocular appearance – features that suggests malignant cystic lesion, requiring surgical resection. MR conclusion was: multilocular right renal cystic tumor. (figure 2)

After clinical evaluation and preliminary lab tests is decided on 04.08.2015, radical nephrectomy through the lumbar approach, with favorable postoperative evolution.

Histopathological parameters were put into evidence after histological preparation with H&E staining: cystic tumor formation with exophytic development in right kidney parenchyma (max. 12/9

Figure 1 - CT exam [nonenhanced - (a) and after intravenous contrast administration in cortico-medullary - (b), nephrographic - (c) and delay, parenchymal phase - (d)]; voluminous, well-defined, extrarenal cystic mass with predominant fluid component (*), containing an expansive small dense nodule (arrow), with intense iodinate enhancement, located just above of some enhanced septa (arrowhead)
Multilocular cystic clear cell renal cell carcinoma imaging with histopathologic correlation - case reports

cm), with a multilocular pattern, polymorph, polychromatic areas with intraseptal solid aspect. The histopathological aspect suggest a renal cell carcinoma - clear cell type, Fuhrman nuclear grade 2, multilocular cystic pattern, richly vascularized, with foci of intratumoral bleeding. The proliferation of renal tumor evolves out of the well defined outline of a conjunctive-collagen pseudo-capsule lamellar without invasive aspects. (figure 3). Renal capsule, vessels, and perinephric fat were free.

The histopathological conclusions where: multilocular cystic clear cell carcinoma clear, Fuhrman nuclear grade 2.

DISCUSSION

With the increasing availability of imaging and methods becoming more efficient, it increased the number of incidentally detected renal tumors in small stages (asymptomatic). Ultrasound (US) is “guilty” for most of incidental “discovered” renal masses.

Computed Tomography (CT) is considered the main method for renal cystic lesions characterization (6), allowing accurate measurement of both fluid attenuation values (0-20HU), but also of fatty, solid or calcified components. Also, CT underpinned the foundations of Bosniak classification, widely embraced by radiologists and urologists. That CT-based renal cyst classification system, proposed in 1986 and modified by Bosniak in 1993, provides a guide for further imaging evaluation or intervention (7).

In its actual form, cysts are classified into four categories, different from the point of view of management: category I (simple renal cysts), category II (benign renal cysts, that are minimally complicated, but not in need of surgery), category III (more complicated cystic lesions, potentially malignant, in need of surgery) and category IV (clearly malignant cystic carcinomas). The classification revised in 1993 include a subset of minimally complicated lesions (category IIF), lesions that
are more complex than a category II cyst but are still thought to be benign and require only serial imaging to confirm stability (8, 9) (table 1).

MR Imaging (MRI) is helpful when renal lesions are detected by others imaging methods but not so well characterized. MRI is offering greater details in complicated renal cysts, with septa and solid nodules or in multilocular cystic lesions (7). Multiplanar MRI acquisitions

<table>
<thead>
<tr>
<th>Category</th>
<th>Criteria and Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>hairline-thin wall; does not contain septa, calcifications, or solid components; water attenuation / fluid signal and does not enhance;</td>
</tr>
<tr>
<td></td>
<td>- no intervention, no follow-up is needed</td>
</tr>
<tr>
<td>II</td>
<td>may contain a few hairline-thin septa in which perceived (not measurable) enhancement may be appreciated;</td>
</tr>
<tr>
<td></td>
<td>- fine calcification may be present in the wall or septa;</td>
</tr>
<tr>
<td></td>
<td>- uniformly high-attenuating lesions that are sharply marginated and do not enhance are included in this group;</td>
</tr>
<tr>
<td></td>
<td>- &lt; 3 cm in diameter</td>
</tr>
<tr>
<td></td>
<td>- no intervention, no follow-up is needed</td>
</tr>
<tr>
<td>III</td>
<td>may contain multiple hairline-thin septa; perceived (not measurable) enhancement of a hairline-thin smooth septum or wall can be identified;</td>
</tr>
<tr>
<td></td>
<td>- there may be minimal thickening of wall or septa, which may contain calcification that may be thick and nodular, but no measurable contrast enhancement is present; there are no enhancing soft-tissue components;</td>
</tr>
<tr>
<td></td>
<td>- totally intrarenal nonenhancing high-attenuating renal lesions (&gt;3 cm) are also included in this category;</td>
</tr>
<tr>
<td></td>
<td>- generally well marginated;</td>
</tr>
<tr>
<td></td>
<td>- need follow-up to prove their benignity by showing stability</td>
</tr>
<tr>
<td>IV</td>
<td>can have all of the criteria of category III but also contain distinct enhancing soft-tissue components independent of the wall or septa;</td>
</tr>
<tr>
<td></td>
<td>- need to be removed</td>
</tr>
</tbody>
</table>

Table 1 - The Bosniak Renal Cyst Classification System

Figure 3 - Photomicrograph of histologic section (H and E X200) shows: (a) typical histologic appearance of clear cell renal cell carcinoma, showing epithelial cells with clear cytoplasm and a distinct cell membrane, separated by a fine branching network of vascular tissue. (b) clear cell epithelial lining with fluid-filled lumen, small polypoid projection into lumen with foci of intratumoral bleeding (arrow); also note dense fibrous component (*)
are very important in cystic renal masses evaluation, because the imaging planes of the sequences may be modified to best depict the mass (10,11).

Multilocular cystic RCC cannot be always differentiated from other complex cystic renal lesions on images. As the name suggests, multilocular cystic RCC is a multiseptated cystic RCC whose septa contain small clusters of clear cells. In these cases, the CT or MR appearances can range from a Bosniak IIF cyst to a Bosniak IV cystic lesion. Multilocular cystic RCCs typically manifest as multilocular cystic tumors, variable-sized, separated from the kidney by a fibrous capsule. Microscopic examination of the surgically resected tissue is often necessary for a diagnosis of certain (3,12,13).

Diffusion-weighted imaging (DWI) is a new technique for renal masses, providing quantification of the Brownian motion of water molecules in tissues, which depends on tissue organization, cellularity, the integrity of cell membranes, and extracellular space tortuosity (14). Qualitative and quantitative information are obtained regarding tissue characterization without the need for Gadolinium administration. Several authors (Goyal, Taouli and Wang) showed that the mean ADC value of clear cell RCC was found to be significantly higher than that of non-clear cell RCC, while Sandrasegaran did not find any significant difference in the ADC values of clear cell RCCs and non-clear cell malignancies (15,16,17,18).

Histopathological analysis demonstrates cysts lined by a monolayer of epithelial cells with clear cytoplasm (13). Multilocular cystic renal cell carcinoma is distinguished from the other subtypes of RCC based on the high incidence of stage I disease at initial presentation (83% to 88%), infrequent metastases, and an extremely high cure rate following surgical resection of the tumor (19,20).

CONCLUSION

Histology of clear cell RCC is reflected in advanced imaging characteristics. Accurate imaging and histologic characterization of RCC and its subtypes (ie multilocular cystic RCC) are very important for the proper management of the patient and its prognostic.

Acknowledgement

This work received financial support through the project entitled „CERO – Career profile: Romanian Researcher”, grant number POSDRU/159/1.5/S/135760, co-financed by the European Social Fund for Sectoral Operational Programme Human Resources Development 2007-2013.

REFERENCES

1. Eble JN, Sauter G et al. Pathology and genetics of urinary systemand male genital organs, Lyon, France, IARC Press, 2004