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Diagnostic Accuracy of Multiparametric Magnetic Resonance Imaging to Identify Clear Cell Renal Cell Carcinoma in cT1a Renal Masses

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Abstract

Purpose

Detection of small renal masses is increasing with the use of cross-sectional imaging, although many incidental lesions have negligible metastatic potential. Among malignant masses, clear cell renal cell carcinoma is the most prevalent and aggressive subtype, and a method to identify such histology would aid in risk stratification. Our goal was to evaluate a likelihood scale for multiparametric magnetic resonance imaging in the diagnosis of clear cell histology.

Methods

Patients with cT1a masses who underwent MRI and partial or radical nephrectomy from December 2011 to July 2015 were retrospectively reviewed. Seven radiologists with different levels of experience and blinded to final pathology independently reviewed studies based on a predefined algorithm, and applied a clear cell likelihood score: 1) definitely not, 2) probably not, 3) equivocal, 4) probably, and 5) definitely. Binary classification determined the accuracy of clear cell versus 'all other' histologies, and inter-observer agreement was calculated with a weighted κ statistic.

Results

In total, 110 patients with 121 masses were identified. Mean tumor size was 2.4 cm and 50% were clear cell. Defining clear cell as scores ≥4 demonstrated sensitivity and specificity of 78% and 80%, respectively, while scores ≥3 were 95% and 58%, respectively. Inter-observer agreement was moderate to good, with a mean κ of 0.53.
Conclusions

A clear cell likelihood score with MRI can reasonably identify clear cell histology in small renal masses, and may reduce the number of diagnostic renal mass biopsies. Standardization of imaging protocols and reporting criteria are needed to improve inter-observer reliability.
Introduction

Approximately one quarter of small renal masses (SRM) less than 4 cm are histologically benign. However, the role of percutaneous biopsy in the management of SRM remains controversial. Recent evidence indicates that while renal mass biopsy (RMB) is highly accurate in renal cell carcinoma (RCC) diagnosis, multiple studies demonstrate underutilization. Potential explanations include that RMB is an invasive procedure with a 1.4%-4.7% complication rate, fails to provide diagnostic information for malignancy in 14% of cases, and is unreliable in tumor grade determination.

Based on excellent soft-tissue contrast and a combination of qualitative, semi-quantitative, and quantitative characteristics, multiparametric magnetic resonance imaging (mpMRI) can histologically subtype renal cell carcinoma (RCC). As an alternative to computed tomography, mpMRI could potentially provide both histologic and anatomic information prior to therapeutic intervention, obviating the need for biopsy or additional imaging. However, literature pertaining to the clinical applicability of mpMRI to define higher-risk RCC in SRM is sparse. Furthermore, to our knowledge, the inter-observer reproducibility between more than three reviewers in mpMRI evaluation of RCC has not been reported.

Clear cell RCC (ccRCC) is the most common RCC variant and notably potentially aggressive. On MRI it is characterized by heterogeneous high signal intensity on T2-weighted imaging, presence of microscopic fat, and avid enhancement equal to or greater than renal cortex. Given that the probability of ccRCC diagnosis is proportional to the added effect
of individual parameters, a likelihood score based on those MRI parameters might provide a useful tool for patient management, as has been done with prostate cancer.  

Our goal was to evaluate the diagnostic accuracy of a subjective likelihood scale for mpMRI in the diagnosis of ccRCC in cT1a lesions and to assess the inter-reader reproducibility of such scale.
Materials and Methods

Patients

This retrospective series was performed under Institutional Review Board approval. Patients with cT1a renal masses who underwent either partial or radical nephrectomy and pre-surgical mpMRI between December 2011 and July 2015 were identified. Patients with poor quality or limited MRI examinations (n=3) (i.e. not including the sequences described below) were excluded from analysis. Patient demographics and clinical findings including age, sex, tumor size, and final pathology were extracted from chart review. Histologic analysis was performed by genitourinary pathologists according to the World Health Organization classification of renal neoplasms.²²

Image Acquisition

mpMRI studies were performed on 1.5T and 3.0T scanners from multiple institutions with diverse parameters, as many patients underwent mpMRI elsewhere and were referred to the authors’ institution for treatment. In total, 54 patients (49%) had imaging at our institution, while 56 patients (51%) had imaging obtained elsewhere. An MRI fellowship-trained radiologist (blinded), who was not involved in imaging interpretation, reviewed all mpMRI exams of patients in this study to ensure they were of sufficient image quality and basic sequences were included. At the authors’ institution mpMRI includes the following sequences: T2-weighted images in coronal and axial (with fat-suppression) planes (4-8mm slice thickness, 256x154 to
348x280 matrix, 70-120ms echo time (TE), and 964-4820ms repetition time (TR)), axial chemical shift T1-weighted images (5-8mm thickness, 193x168 to 500x286 matrix, 2.38/4.87ms (1.5T) or 1.1/2.3ms (3.0T) TE, and 100-216ms TR), and fat-suppressed dynamic contrast-enhanced (DCE) T1-weighted imaging, including corticomedullary and late nephrographic and/or excretory phases in either axial or coronal planes (3-6mm slice thickness, 176x149 to 320x259 matrix size, 1.6-2.4ms TE, and 3.4-5.4ms TR). Diffusion weighted images (DWI) were routinely acquired at the authors’ institution but inconsistently present in mpMRI studies performed elsewhere so these were not evaluated. Furthermore, given the controversial role of DWI in characterizing renal masses and the importance of acquisition protocol standardization, these images were not included in the analysis.

Image Analysis

Seven radiologists with fellowship training in body MRI from a single institution with a busy MRI service (approximately 7,000 abdominopelvic MRIs/year), varying levels of experience (IP and JRL 15 years, DC 12 years, TY 10 years, GK 7 years, DC 5 years, ADL 1 year), and blinded to final pathology, independently reviewed each study on a PACS workstation (iSite, Philips Healthcare, Best, Netherlands). A Likert scale was used to convey the subjective radiologic impression of the likelihood of encountering ccRCC at surgery. The clear cell likelihood score (ccLS) was defined as: 1) definitely not ccRCC, 2) probably not ccRCC, 3) equivocal for ccRCC, 4) probably ccRCC, and 5) definitely ccRCC. “Cannot be assessed” was also allowed if perceived inadequacy of the MRI protocol. Prior to image analysis, all radiologists received a refresher
training session with a slide presentation including examples of the main imaging features associated with ccRCC histology and other common histologic diagnoses in renal masses. A detailed review of such features is published elsewhere (figure 1). Each reader was allowed to measure signal intensity using region of interest (ROI) analysis, and quantify the corticomedullary enhancement and arterial-delayed enhancement ratio per individual judgment. For statistical purposes, a composite ccLS was calculated as a mean of the seven readers when appropriate.

Statistical Analysis

Scoring each renal mass for ccRCC diagnosis was compared to the reference standard, which was determined by histology (ccRCC vs. ‘all other’ histologies). Receiver operating characteristic (ROC) analysis was performed to calculate the area under the curve (AUC) and corresponding 95% confidence interval for each reader. Tumors that could not be assessed were excluded from ROC analysis. Multiple comparisons in mean AUCs among readers were performed with a Bonferroni correction. Pairwise inter-observer agreement was assessed by weighted κ statistics. A p-value of 0.05 or less was considered statistically significant. All statistical analysis was performed with SAS version 9.4 (SAS Institute Inc., Cary, NC, USA).
Results

In total, 110 patients with 121 cT1a renal masses were identified. Patient demographics and final pathologic findings are shown in table 1. Mean tumor size was 2.4 cm (range 0.5–4.0 cm), and histology was ccRCC in 50% (n=61), non-clear cell RCC in 33% (n=40), and benign lesions in 17% (n=20).

Histologic distribution of the entire cohort by ccRCC versus ‘all other’ histologies and assigned ccLS is shown in figure 2, while overall comparison of ccLS score to final histology is shown in table 2. Overall, a mean of 2.8 tumors (2.3%) could not be assessed by reviewers. Tumors with ccLS 4-5 were categorized as mpMRI-identified ccRCC (MR-ccRCC). Combining all seven readers gave a mean accuracy of 79%, sensitivity of 78%, specificity of 80%, positive predictive value (PPV) of 80%, and negative predictive value (NPV) of 80% (figure 3). Augmenting the MR-ccRCC definition to include ccLS 3 tumors changed the mean accuracy to 77%, sensitivity to 95%, specificity to 58%, PPV to 70%, and NPV to 93%, which inversely gave ccLS 1-2 a specificity of 95% and PPV of 93% for ‘all other’ histologies.

Among all readers, a mean of 12.2 tumors (10.1%) were false positive ccLS 4-5 lesions. The most common false positive histology was oncocytoma (mean 2.7 tumors, 22.1%) followed by papillary RCC (mean 1.3 tumors, 10.7%), chromophobe RCC (mean 1 tumor, 8.2%) and lipid-poor angiomyolipoma (AML) (mean 0.9 tumors, 7.4%). A mean of 2.7 tumors (2.2%) per reader were false negative ccRCC tumors graded as ccLS 1-2; 50% without high signal intensity on T2-weighted imaging, 79% without high contrast avidity, 100% without intravoxel fat, and 84.2% without central scar, which suggested against higher scoring.
ROC curves for the seven readers are shown in figure 4, with AUC ranging from 0.82-0.92. Comparisons of the highest rated reader (Reader 7, AUC 0.92) compared to all other readers showed a statistically significant difference between all readers except reader #4, and years of experience did not correlate with improved diagnostic accuracy (p=0.31). Inter-reader variability was moderate to good, with a mean weighted κ of 0.53 (range 0.38-0.64).
Discussion

Multiparametric MRI is an appealing alternative to RMB, in that it can provide both pathologic information to direct management and anatomic information required for surgical planning. To our knowledge, our series is the first to specifically evaluate the diagnostic accuracy of mpMRI for ccRCC in cT1a lesions. The proposed ccRCC likelihood scoring system demonstrated a high specificity and PPV for ccRCC in ccLS 4-5 lesions, and a high specificity (95%) and PPV (93%) for non-ccRCC in ccLS 1-2 lesions, in addition to moderate inter-observer reliability. Therefore, ccLS could support an algorithm where all ccLS 4-5 are encouraged to undergo curative interventions, all ccLS 1-2 are placed on active surveillance (especially if they are less than 3 cm), and only ccLS 3 undergo RMB. This proposal would result in a biopsy rate in this series of only 20%, unnecessary treatment of oncocytoma and lipid-poor AML in 4.5% and 1.7% of the surgical cohort, respectively, and 4.4% of ccRCC placed on active surveillance. It should be noted that ccLS was not a good predictor of overall malignancy. This is expected since it was developed for ccRCC diagnosis. Accordingly, while most ccRCCs received a high score, most non-clear cell RCCs were assigned a low score.

In one of the first studies to evaluate a comprehensive classification of renal masses with mpMRI, Pedrosa et al. defined eight distinct qualitative mpMRI patterns of malignant lesions among T2-, chemical shift T1-, and contrast enhanced T1-weighted sequences. They reported that two radiologists demonstrated a slightly higher sensitivity and specificity of 92% and 83%, respectively, compared to our series for diagnosing ccRCC. Of these 48 ccRCC tumors mean tumor size was much larger than the present series (6.0 cm), and they excluded benign
lesions, which may explain the slightly higher accuracy. Cornelis and colleagues compared mpMRI of 100 renal masses to pathologic findings, and noted high specificity for diagnosing papillary RCC and oncocytomas. Differing from our study, they included significant parameters quantified by two radiologists, and all RCC subtypes were mean >4.0 cm. Recently, Hötker and colleagues published their evaluation of 124 renal masses by MRI compared to pathologic findings. Although the median tumor size of their ccRCC cohort (n=81) was 8.0 cm, they noted that the apparent diffusion coefficient, peak enhancement, and downslope of DCE sequence significantly correlated with ccRCC histology. In addition, after two independent radiologist’s review they noted significant interreader agreement (r= 0.82-0.99).

Despite other investigators use of mpMRI in large (>4.0 cm) renal tumors, in the context of clinical practice we feel the most relevant use of mpMRI is in classification of SRM (<4.0 cm). In a database of over 2700 patients, Frank and colleagues noted that 23% of tumors <4.0 cm were pathologically benign, compared to only 8% of tumors ≥4.0 cm. Therefore, the current role of RMB is in identifying SRM histology to plan surgical intervention or active surveillance. Recent meta-analysis of RMB have demonstrated low false positive and false negative rates of 4.0% and 3.1%, respectively, in addition to low complication rates (4.9% hematoma and 1.2% pain). However, as previously mentioned, this is in the context of a 14% non-diagnostic rate. In addition, histologic grade concordance between biopsy and final pathology was 52-76%. In comparison, our mpMRI results demonstrated false-positive and false negative rates of 6.2% and 4.4%, respectively, no complications, and a 2.3% non-diagnostic rate. As for tumor grade, we did not analyze grade concordance in this initial series. Prior work has looked at
differentiating high grade and low grade ccRCC in large tumors with good success. The ability to do so in smaller tumors is unknown, but likely more difficult.

Additionally, many RMB studies carried selection bias as percutaneous biopsy may not be easily performed and/or may carry worse outcomes (e.g. lower diagnostic yield, increased complications) in those patients with SRMs in difficult anatomic locations (e.g. perihilar, anterior), particularly at centers without expertise in performing these procedures. Alternatively, mpMRI would potentially avoid additional procedures, and the lack of invasiveness might encourage provider utilization.

The issue of false-positives was most notable with oncocytomas, emphasizing the challenge in differentiating it from ccRCC. Similarly, Cornelis and colleagues noted difficulty in their retrospective evaluation as two radiologists blinded to final pathology had a diagnostic specificity of 94%, but only 19% sensitivity, in distinguishing ccRCC from oncocytoma. Similarly, Rosenkrantz et al. found no MRI features that would allow differentiation of oncoytoma from chromophobe RCC using a standard clinical MRI protocol similar to the mpMRI protocols in our study. Further studies, specifically with arterial spin labeling (ASL), may have the potential to separate oncocytomas and ccRCC. Although not currently commercially available, this technique labels the spin of arterial water allowing, in effect, contrasted imaging without intravenous tracers. Prior work has shown significant differences in mean ASL perfusion levels of oncocytoma and ccRCC. Adding this technique to the mpMRI algorithm may further improve diagnostic accuracy.
As highlighted above, most studies regarding mpMRI involve at most a few reviewing radiologists.\textsuperscript{7,12} To our knowledge, we are the first to evaluate inter-reader variability among a large cohort of radiologists. The seven readers in our series had varying levels experience, ranging from in-fellowship training to senior faculty (1-15 years) and we found that radiologist level of experience did not appear to correlate with ccLS accuracy. The reason for this is speculative, but might be a reflection of more recent emphasis on mpMRI exposure in residency and fellowship programs. Additionally, the structured algorithm might mitigate some effects of experience. As for inter-reader variability, we demonstrated good consistency (mean weighted $\kappa$ 0.53) among our radiologists when utilizing the algorithm and ccLS scoring. However, this also highlights opportunity for improvement. Given the multitude of adjustable parameters with mpMRI, inter-center standardization is needed to specify imaging protocols that will best characterize SRM.

Our study has limitations that deserve mention. First is the retrospective design. While our radiologists were blinded to final pathology, this structure has inherent bias. Not all patients with SRM undergo mpMRI, so there is selection bias that could affect our estimates of RMB (ccLS 3, 20%) or unnecessary ccLS 4-5 surgery (6.2%) in this series. In addition, not all patients who had mpMRI underwent surgery. This selection bias limits our ability to capture benign appearing lesions, which could also affect our accuracy. Therefore, the strength of this modality will ultimately depend on a prospective analysis to verify that ccLS can accurately direct SRM management. Second is the single-institution series. As stated previously, all mpMRI sequences are not equal, and standardization is required to improve inter-center reproducibility. However, the reasonable diagnostic accuracy of the ccLS considering the
multiple mpMRI studies performed at different institutions with different imaging protocols is encouraging. Future studies should be multi-institutional to ensure stability of results between various centers. Third, our study is small with a cohort of only 121 SRM, of which 61 were ccRCC. However, in comparison to other discussed series,\textsuperscript{7,12,13} we do report the largest SRM series to date. Larger series are ultimately needed to define the accuracy of this work.

These limitations notwithstanding, we have demonstrated that ccLS can accurately predict ccRCC in cT1a renal masses. In addition to providing anatomic information for surgical planning, this could streamline SRM decision-making and obviate the need for additional procedures prior to definitive management. Moreover, our reported technique has good inter-reader variability. With future prospective and multi-institutional studies we hope to verify these initial results and the generalizability of this technique.
References


Figure Legends


Figure 2. Histologic distribution of the entire cohort by clear cell likelihood score. Tumors that could not be assessed are not shown.

Figure 3. Binary classification of the seven readers defining clear cell RCC as ccLS 4-5. RCC- renal cell carcinoma. ccLS- clear cell likelihood score.

Figure 4. Receiver operating characteristic curves of the seven readers defining ccRCC and ccLS 4-5. RCC- renal cell carcinoma. ccLS- clear cell likelihood score.
Table 1. Patient demographics and final pathology. SD= standard deviation, RCC= renal cell carcinoma.

<table>
<thead>
<tr>
<th></th>
<th>n= 110 patients (121 tumors)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y (mean ± SD)</td>
<td>57 ± 14</td>
</tr>
<tr>
<td>Male/Female (%)</td>
<td>61/39%</td>
</tr>
<tr>
<td>Tumor size, cm (mean ± SD)</td>
<td>2.4 ± 0.8</td>
</tr>
<tr>
<td>Histology (%)</td>
<td></td>
</tr>
<tr>
<td>Clear cell RCC</td>
<td>61 (50%)</td>
</tr>
<tr>
<td>Papillary RCC</td>
<td>25 (21%)</td>
</tr>
<tr>
<td>Chromophobe RCC</td>
<td>9 (7%)</td>
</tr>
<tr>
<td>Other RCC</td>
<td>6 (5%)</td>
</tr>
<tr>
<td>Oncocytoma</td>
<td>7 (6%)</td>
</tr>
<tr>
<td>Angiomyolipoma</td>
<td>6 (5%)</td>
</tr>
<tr>
<td>Other benign</td>
<td>7 (6%)</td>
</tr>
<tr>
<td>Final pathology</td>
<td></td>
</tr>
<tr>
<td>pT1a</td>
<td>97 (96%)</td>
</tr>
<tr>
<td>pT3a</td>
<td>4 (4%)</td>
</tr>
<tr>
<td>Fuhrman Grade</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>6 (6%)</td>
</tr>
<tr>
<td>2</td>
<td>72 (71%)</td>
</tr>
<tr>
<td>3</td>
<td>15 (15%)</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Unknown</td>
<td>8 (8%)</td>
</tr>
</tbody>
</table>
Table 2. Mean percentage according to histologic subtype versus clear cell likelihood score. Values in each cell represent column percentage within each histology, while numbers in parenthesis indicate the mean number of renal masses across all readers. ccLS- clear cell likelihood score, RCC- renal cell carcinoma, AML- angiomyolipoma.

<table>
<thead>
<tr>
<th>ccLS</th>
<th>Clear cell RCC</th>
<th>Papillary RCC</th>
<th>Chromophobe RCC</th>
<th>Oncocytoma</th>
<th>Lipid-poor AML</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>1- Definitely not</td>
<td>0.7% (0.4)</td>
<td>66.1% (16.6)</td>
<td>20.9% (1.9)</td>
<td>0.0%</td>
<td>29.7% (1.9)</td>
<td>13.0% (1.7)</td>
</tr>
<tr>
<td>2- Probably not</td>
<td>3.8% (2.3)</td>
<td>20.3% (5.1)</td>
<td>15.4% (1.4)</td>
<td>27.1% (1.9)</td>
<td>15.7% (1.0)</td>
<td>12.2% (1.6)</td>
</tr>
<tr>
<td>3- Equivocal</td>
<td>17.3% (10.6)</td>
<td>5.6% (1.4)</td>
<td>48.4% (4.4)</td>
<td>34.3% (2.4)</td>
<td>25.0% (1.6)</td>
<td>25.2% (3.3)</td>
</tr>
<tr>
<td>4- Probably</td>
<td>38.8% (23.7)</td>
<td>4.0% (1.0)</td>
<td>11.0% (1.0)</td>
<td>24.3% (1.7)</td>
<td>9.4% (0.6)</td>
<td>27.5% (3.6)</td>
</tr>
<tr>
<td>5- Definitely</td>
<td>38.6% (23.6)</td>
<td>1.2% (0.3)</td>
<td>0.0%</td>
<td>14.3% (1.0)</td>
<td>4.7% (0.3)</td>
<td>20.6% (2.7)</td>
</tr>
<tr>
<td>Cannot be assessed</td>
<td>0.8% (0.5)</td>
<td>2.8% (0.7)</td>
<td>4.4% (0.4)</td>
<td>0.0%</td>
<td>15.6% (1.0)</td>
<td>1.5% (0.2)</td>
</tr>
</tbody>
</table>
Signal Intensity on T2WI (compared to kidney)

Enhancement

- Central non-enhancing area and/or Intravoxel fat
- +/- Central scar

Hyperintense
- Intense
- Moderate

Isointense
- Intense
- Moderate
- Mild

Hypointense
- Intense
- Mild

ONCO
chrRCC
ccRCC

pRCC

AML

Other (e.g. adenoma leiomyoma, lymphoma)

CORTICO-MEDULAR PHASE

INTENSE =/= than renal cortex
MODERATE 50% of renal cortex
MILD 25-30% of renal cortex

Arterial-Delayed Enhancement Ratio

ADER = SI_{art} - SI_{pre}/SI_{del} - SI_{pre}
>1.5 AML with minimal fat
<1.5 ccRCC
Abbreviations and Acronyms

(mp)MRI = (multiparametric) magnetic resonance imaging
SRM = small renal mass
RMB = renal mass biopsy
(cc)RCC = (clear cell) renal cell carcinoma
TE = echo time
TR = repetition time
DWI = diffusion weighted imaging
ccLS = clear cell likelihood score
ROC = receiver operating characteristic
AUC = area under the curve
PPV = positive predictive value
NPV = negative predictive value
AML = angiomyolipoma
ASL = arterial spin labeling