Pathologic T3a renal cell carcinoma: a classification in need of further refinement

John M. DiBianco¹, Patrick T. Gomella², Mark W. Ball²

¹Department of Urology, George Washington University Medical School, Washington DC, USA; ²Urologic Oncology Branch, National Cancer Institute, National Institutes of Health, Bethesda, MD, USA

Correspondence to: Mark W. Ball, MD. Urologic Oncology Branch, Center for Cancer Research, National Cancer Institute, Building 10, Room 1-5940, Bethesda, MD 20892, USA. Email: mark.ball@nih.gov.

Provenance: This is an invited Editorial commissioned by Section Editor Xiao Li (Department of Urology, Jiangsu Cancer Hospital & Jiangsu Institute of Cancer Research & Nanjing Medical University Affiliated Cancer Hospital, Nanjing, China).


Submitted Dec 14, 2018. Accepted for publication Dec 18, 2018.
doi: 10.21037/atm.2018.12.51

View this article at: http://dx.doi.org/10.21037/atm.2018.12.51

Kidney cancer is associated with a significant global disease burden (1,2). Although mortality rates for renal cell carcinoma (RCC) in the United States remain stable, the incidence of RCC continues to rise, largely due to increased incidental detection on routine abdominal imaging (1,3). Ultimately, both treatment recommendations and patient prognosis are based on the stage of disease (4,5). Pathologic staging of RCC continues to be refined, especially for the characterization of pathologic T3a (pT3a) disease, which represents locally advanced disease with invasion of the fat or vasculature of the kidney (Figure 1), and is associated with worse oncologic outcomes compared to organ confined disease (6,7).

A recent study, titled “Prognostic evaluation of perinephric fat, renal sinus fat, and renal vein invasion for patients with pathological stage T3a clear-cell renal cell carcinoma” by Shah et al. investigated the prognostic significance of various patterns of extrarenal extension within the categorized T3a for clear-cell renal cell carcinoma (ccRCC) (9). Specifically, they investigated the prognostic significance of invasion of perirenal fat (PF), RSF and RV and/or a combination of these
variables. The authors identified patients who underwent radical nephrectomy between 1970 and 2011 who were diagnosed at pT3a, pNX/0, M0 clear cell RCC utilizing an institutional registry. A total of 563 patients were identified and progression-free survival, cancer specific survival and overall survival were analyzed. They found that while individually tumor PF, RSF, or RV invasion did not differ significantly in oncologic outcomes, tumors found to have a combination of such invasion portended a worse prognosis than those with singular sites. This finding was associated with a 2.66-fold increase in cancer specific mortality. They conclude that their findings may help to refine risk-stratification of non-metastatic T3a RCCs to better inform patient prognosis and clinical trial enrollment (9).

While this manuscript is a valuable contribution to the literature, several points should be considered. First, the median tumor size in this study was 9 cm, meaning that even in the absence of fat or venous invasion, these tumors were at least clinical T2a (cT2a). The significance of pT3a upstaging may differ between cT2a and lower staged tumors. Additionally, these tumors were treated with radical nephrectomy. The significance of these features in the partial nephrectomy population may be different. For these reasons, these results may not be generalizable to patients with either smaller tumors or those who undergo partial nephrectomy. Patients who undergo partial nephrectomy typically have tumors that are smaller, more peripherally located and less complex than those in which radical nephrectomy is undertaken (5).

The decision to perform radical or partial nephrectomy for T2 tumors may be based on multiple patient factors, such as age and comorbidities, as well as tumors factors like
size, RENAL nephrometry score and tumor location (10). Recently, partial nephrectomy has increased in utilization, particularly for larger more complex tumors, although the decision (5,11).

Due to the increased interest in adjuvant systemic therapy, patient specific and disease stratification may be increasingly studied in years to come. Both tumor genetics and staging may be integral to patient follow up strategies, surveillance and possible systemic treatments (12-14). The authors of the current study (Shah et al.) propose their findings may help to refine risk-stratification, however, the recommended follow up surveillance strategy for stage 3 disease is not altered due to substages (12,15). Currently, the follow up strategy only is altered by the upstaging to stage 4 disease.

Several studies have examined the outcomes of partial nephrectomy for T3a tumors with results demonstrating equal to acceptable oncological outcomes when compared to radical nephrectomy (5,16). Gorin et al. examined the clinical outcomes of patients who underwent partial nephrectomy for clinical stage T1 or T2 disease who were then upstaged to T3a on pathological examination (16). They found that approximately 10% of these patients experienced either local or distant disease recurrence, however they did not substratify for the differences between PF, RSF, or RV invasion (16). Ball et al. compared outcomes of segmental RV invasion versus main RV invasion and found that main RV invasion portended worse disease-specific and overall survival (6). Taken together, these studies indicate that the current definition of pT3a kidney cancer may be too broad as there is significant heterogeneity of clinical outcomes in this group. Larger, multi-center studies may be helpful in clarifying how future staging schema may be more predictive of oncologic outcomes.

Acknowledgements

This research was supported (in part) by the Intramural Research Program of the National Cancer Institute, NIH.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

References


Cite this article as: DiBianco JM, Gomella PT, Ball MW. Pathologic T3a renal cell carcinoma: a classification in need of further refinement. Ann Transl Med 2018;6(Suppl 2):S133. doi: 10.21037/atm.2018.12.51