Urothelial cancers: clinical and imaging evaluation

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Abstract: Urothelial cancers are currently diagnosed more commonly parallel to the recent developments in imaging techniques. Early diagnosis of these tumors, together with better treatment options, decreases the morbidity and the mortality significantly. This review aims to summarize the imaging findings of urothelial tumors, both at the pretreatment and posttreatment follow-up stage.

Key words: MRI, urography, hematuria, ureter cancer, urinary tract cancer, computed tomography, diagnosis, staging, ultrasound, urinary tract imaging, urothelial tumors

Introduction

Urothelial tumors originate from kidney collecting systems, ureters, the bladder, and the urethra. Their diagnosis and treatment approaches are quite different from those of parenchymal tumors of the kidney (1–3). Radiologists are expected to depict the presence, location, depth penetration, and local and distant metastatic status of these lesions, which usually present with painless hematuria (1–3). Herein, we aimed to review the radiologic approach, diagnostic methods, and differential diagnosis of urothelial tumors.

Urothelial tumors include pelvic, ureteral, bladder, and urethra tumors, and the majority of them are bladder tumors (3). They are most commonly seen in the 6th and 7th decades of life with a 4:1 male-to-female ratio (1–3). Transitional cell carcinoma (TCC) constitutes 95% of malignant urothelial tumors (3–6). More than 90% of all bladder cancers are TCC, 5%–10% are squamous cell carcinoma (SCC), and the remaining 2%–3% are adenocarcinoma (3,6–10). SCC is associated with chronic inflammation, whereas adenocarcinoma is associated with persistent urachal remnants (3,11). Bladder cancers are commonly located in the trigone and its adjacency (4–10). The second most common location of TCC is the renal pelvis (1,11). It has a tendency to be multifocal and of low grade, similar to in the bladder (1,3). TCC of the urethra commonly involves the distal portion and tends to be superficial, but it behaves more aggressively, unlike in the bladder (4–8). Other rare urinary system tumors are lipoma, fibroma, leiomyoma, hemangioma, sarcoma, lymphoma, mesodermal tumors, carcinoid tumors, and pheochromocytoma, and these should be considered in differential diagnosis (12).

Etiology

Risk factors for TCC include smoking, analgesics (i.e. phenacetin), carcinogens (arsenic), schistosomiasis, familial cancer syndromes, and renal papillary necrosis (4). Smoking increases the risk 4-fold (13). Balkan nephropathy, which is seen in Europe, also increases the risk for TCC (14). TCC tends to be multifocal and highly recurrent (5,15).

Staging

Pathologic staging

Urothelial tumors are grouped into 2 types, based on their morphology, as invasive and superficial. Superficial ones are usually papillary, of low grade,
and limited to the mucosa lamina propria (3). They rarely metastasize; however, recurrence after treatment is common with a good prognosis (5,16). Invasive ones, however, generally appear smooth and develop from high-grade in situ lesions; they are localized to the surface epithelium. They appear as hyperemic mucosa at cystoscopy (3). Carcinoma in situ lesions can manifest with tumor cells in urine cytology and are known as precursors of invasive cancers (17).

Renal pelvis tumors are generally malignant; they involve both kidneys equally and are more commonly seen in males (8). Well-differentiated tumors are Grade I and intermediately differentiated ones are Grade II, whereas poorly differentiated tumors are Grade III (16).

Clinical staging

Clinical staging is done according to tumor–node–metastasis (TNM) classification (3). An analysis of the depth and spread of the tumor and lymph node involvement and a metastases workup are performed (2). Locally, they may involve the bladder orifice and trigone, ureters, uterus, vagina, urethra, vagina, prostate, and rectum (3–8,11). Lymphatic spread occurs through the paravesical, obturator, external iliac, and paraaortic lymph nodes, whereas hematogenous spread includes the liver, lungs, bones, and adrenal glands (2–5,18). Common iliac and paraaortic lymph node involvements are also considered as distant metastases (1).

Tumor staging (2,3)

Tx: Absence of an evaluable primary tumor.
T0: No tumor.
Tis: Carcinoma in situ.
Ta: Papillary tumor in the epithelium (Figure 1a).
T1: Lamina propria is invaded by the tumor.
T2: Muscular layer is invaded by the tumor (Figure 1b).
T2a: Superficial muscular layer is invaded by the tumor.
T2b: Deep muscular layer invaded by the tumor.
T3: Perivesical fat planes are invaded.
T3a: Microscopic invasion of the perivesical fat planes.
T3b: Macroscopic invasion of the perivesical fat planes.
T4a: Adjacent organs (prostate, rectum, and vagina) are invaded by the tumor (Figure 2).
T4b: Abdomen–pelvic wall is invaded by the tumor.

Lymph nodes

Nx: Nodal metastases are unknown.

Figure 1. A patient with papillary bladder cancer (TNM staging: T2b): a) axial T2-weighting MR image demonstrates multiple, projectile, papillary masses in the bladder, and b) axial diffusion-weighted image also localizes these masses with a hyperintense signal pattern.
N1: Single pelvic lymph node less than 2 cm in diameter (Figure 3a).

N2: Pelvic lymph node metastases with a diameter of 2 cm to 5 cm (Figure 3b).

N3: Pelvic lymph node metastases greater than 5 cm in diameter.

Metastases
MX: Distant metastases are unknown.
M0: Absence of distant metastases.
M1: Presence of distant metastases.

Imaging techniques for diagnosis
Conventional cystoscopy is the gold standard of diagnostic methods (1–3). Imaging methods for diagnosis include intravenous pyelography (IVP), ultrasound (US), computed tomography (CT), and magnetic resonance imaging (MRI). The role of the direct urinary system radiograph is limited to depicting the presence of calcification and gas in the collecting system (2). IVP is less sensitive in the detection of smaller lesions (10). US is usually used as a first-line method and is sensitive in the detection of bladder tumors. US can be performed by transabdominal, transrectal, or transurethral approaches. CT is the most commonly used method for evaluation of the primary tumor, lymph nodes, and metastases (1). MRI is more sensitive than CT due to its superior soft tissue contrast resolution and it is useful in the investigation of tumoral depth penetration, distant metastases, bony involvement, and late fibrosis versus recurrence distinction (19–22).

Intravenous pyelography
IVP is the traditional initial imaging method in the imaging workup of hematuria (2). At the initial step of IVP, a direct urinary system radiograph should be

Figure 2. Axial CT images of a bladder cancer patient (T4 disease with seminal vesicle invasion) shows multiple focal wall thickening and a right posterolaterally localized mass (arrows) extruding from the wall into the lumen with contrast material enhancement (arrows) (upper image). The perivesical fat planes and perirectal space are both invaded by the lesion (arrows) (lower image).

Figure 3. Axial CT images of a bladder cancer patient show a right bladder wall mass (star) (upper image) with enhancing perivesical and perirectal lymph nodes (arrows), which correspond to N2 disease (lower image).
obtained prior to the contrast injection in order to visualize any opacity (10). IVP has nephrography, pyelography, and cystography phases (2). Large tumors appear as filling defects at the pyelography and cystography phases (23), whereas small tumors and intradiverticular lesions may not be visualized via this imaging method (10). The main limitation of this method is the radiation exposure. Additionally, as it includes an intravenous iodine-based contrast agent, it is contraindicated in patients with kidney failure and contrast allergy history (2,21,23). IVP has an accuracy range of 26% to 87% in the detection of bladder cancers with a smallest detected lesion size cutoff of 1.5 cm (2,9,10,23).

**Ultrasound**

US is a relatively cheaper and more accessible imaging method and is used as a first-line technique for the imaging workup of hematuria. US can precisely depict bladder and pelvis tumors in experienced hands but is limited in ureteral lesions due to gas superposition (2). Hydronephrosis of the kidneys, stones, cysts, and other filling defects within kidney collecting systems can be easily demonstrated at US, whereas bladder lesions appear as a projectile mass or wall thickening (2) (Figure 4). The disruption of the echogenic line around the bladder wall at US represents a possible invasion in bladder cancer patients (23).

A relatively less invasive nature, lack of any requirement for an intravenous contrast agent, and repeatability are among the main advantages of US; however, operator dependence, difficulty in cases of patient obesity, and abdominal gas superposition are among the main disadvantages of this technique. Bladder lesions that are smaller than 5 mm and localized to the bladder dome or neck can be missed via US (2,23). The overall accuracy of US for bladder cancer detection is around 82%–95% regardless of the lesion size (24,25). US can be limited in visualization of the distal renal pelvis, where IVP and CT have more success, but hydronephrosis secondary to tumors can be detected and graded by US (9,10). Additionally, US can be helpful in the evaluation of patients with sepsis, when IVP is contraindicated. Finally, Kocakoc et al. recently reported that virtual (3-dimensional) ultrasonographic cystoscopy can be a useful alternative for the evaluation of bladder cancers (26).

**Computed tomography**

CT is the most commonly used imaging method for urinary system tumors, especially in staging (9,10). The advent of multidetector technology has increased its use and utility in the evaluation of urinary system tumors. A standard abdomen CT scan includes an intravenous injection of 100–120 mL of nonionic contrast material, at an injection rate of 2.5–3 mL/s and a subsequent image acquisition at the parenchymal phase, with an approximate delay of 60–100 s following the injection (2,27). Kidney parenchyma and tumor enhancement can be evaluated at the nephrographic phase (9,10) (Figure 5). Epithelial tumors can be assessed more accurately at the excretory or pyelographic phase and they appear as filling defects (1,2) (Figure 6). The bladder should be sufficiently full and distended for proper lesion evaluation (Figure 7). Reconstructed images in various planes enable better evaluation of lesions localized to the dome and base of the bladder (27) (Figure 7). In some patients, ureteral lesions can be barely visualized, especially if there is incomplete opacification (9,10). In addition to routine CT scans, several special methods can also be used. These are summarized below.

**CT urography**

CT urography is the cross-sectional analog of IVP. In this technique, image acquisition is performed at 7–15 min after the intravenous contrast injection and
the images are usually evaluated after a maximum intensity projection (MIP) reconstruction algorithm application (9,10). Tumors appear as focal wall thickening or as masses projecting into the lumen (9,10). The usage of multidetector scanners increases both the sensitivity of this technique as well as enables a more accurate depiction of lesions smaller than 4 mm (28).

Antegrade CT pyelography

In this technique, the renal pelvis is imaged following catheterization after a subsequent contrast injection through the catheter. In patients with kidney failure, this technique allows for the determination of the obstruction site without an intravenous contrast injection (2). Moreover, this technique can be considered as an alternative in patients with contrast allergy (Figure 8) (29). However, the invasiveness of this technique is a major drawback.

CT cystoscopy and virtual cystoscopy

This technique enables images to be obtained that are similar to those of conventional cystoscopy. However, its major limitations are the lack of real-time tissue sampling (biopsy), the inability to evaluate mucosa and carcinoma in situ, and the less accurate demonstration of very small lesions (23,30,31). On the other hand, it has some advantages over conventional cystoscopy, such as the fact that it can be easily performed as a noninvasive alternative in patients with benign prostate hyperplasia, bladder infections, urethral strictures, and prostatitis, when the conventional technique is contraindicated (32).

Additionally, virtual cystoscopy is more successful in the evaluation of the bladder neck and in diverticula with a narrow neck, where the conventional technique is limited (32,33). The virtual cystoscopy method allows for a 360° visualization of the bladder on all planes, similar to or even better than the conventional method (32). Through this technique, the location and texture of the lesion can be assessed prior to biopsy or surgery, and this can aid in surgical planning and reduce the procedure time. Moreover, it can be used at the follow-up, as well (30–33) (Figure 9).

In this technique, the bladder is filled directly with air (minimally invasive, through a Foley catheter) or indirectly with intravenous contrast material (noninvasively). The air-filled approach requires 2 CT data sets (prone and supine) and therefore has a higher radiation dose exposure. The indirect method is performed at a supine position with a lower radiation dose exposure (34). However, in the indirect approach, virtual images can be obscured by artifacts secondary to the improper mixture of the contrast material and urine, especially in the increased trabeculation areas of the bladder wall (35).

Magnetic resonance imaging

MR urography

MR urography (MRU) can be used for patients with an iodine-based contrast material allergy or minimal renal insufficiency instead of CT urography.
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Under such circumstances, it is safer to use macrocyclic MRI contrast agents (21). MRU has both static and functional components. In the static part (also called a noncontrast-enhanced or static MRU), heavily T2-weighted sequences are obtained without a contrast material injection, whereas in the functional part (also called a contrast material-enhanced or excretory MRU), the contrast-enhanced 3D gradient echo (GRE) T1-weighted sequences are acquired (20,37–39). Noncontrast-enhanced MRU (NCE-MRU) has less sensitivity for the detection of smaller lesions (37).

In contrast-enhanced MRU (CE-MRU), intravenous furosemide (10–20 mg) can be used to avoid the T2* effect of the contrast dilution and to distend the urinary system sufficiently (40). The most important postcontrast T1-weighted images are those obtained at the nephrographic and pyelographic phases. Smaller lesions appear as enhancing foci at the nephrographic phase, whereas they appear as a filling defect at the delayed excretory phase (36). Ureteral lesions can be visualized in either of these sequences (10).

Papillary tumors are seen as filling defects at MRU (36). Postcontrast T1-weighted images at the nephrographic phase are helpful for diagnosis and such lesions can be differentiated from others by their uniform homogeneous contrast enhancement (41). Flat lesions are challenging to depict due to hyperintense urine on T2-weighted sequences or at NCE-MRU and can be easily missed. CE-MRU is more useful in the diagnosis of such lesions (10,40,41).

MR cystography and virtual MR cystoscopy

About 80% of bladder cancers are polypoid, and MR cystography and virtual MR cystoscopy are important in localizing such lesions (41). On T2-weighted images, since urine appears hyperintense, it is easier to visualize pathologies with different signal characteristics. Virtual cystoscopy is obtained through the postprocessing of the T2-weighted data with multiplanar reconstruction and MIP algorithms. MR cystoscopy has higher sensitivity and specificity for lesions greater than 10 mm, whereas these values diminish for lesions smaller than 10 mm. Virtual cystoscopy performed with multidetector CT has a better spatial resolution than that of MRI, and therefore it is more sensitive to smaller lesions. Mural...
lesions and perivesical infiltration cannot be precisely detected at MR cystography. Moreover, in situ lesions cannot be identified, as there is no information about mucosal color changes (42,43).

The main advantages in using this technique as an alternative in case of a contraindication for the conventional cystoscopy are better localization of the bladder neck and intradiverticular lesions and the easiness of the follow-up of patients with local excision (44). The main disadvantages are operator dependency, inability to obtain real-time biopsy, higher cost, and the requirement of higher-strength magnet systems (42). The common indications for MR cystoscopy can be summarized as the need for bladder evaluation in case of urethral strictures, the investigation of bladder diverticula, and conditions in which conventional cystoscopy is contraindicated (42).

**Diffusion-weighted MRI**

Diffusion refers to the motion of the water molecules in biologic tissues following excitation via heat, which is also known as Brownian motion (45–47). Cancers appear as hyperintense foci on diffusion-weighted MRI (DW MRI) whereas they are hypointense on apparent diffusion coefficient (ADC) maps derived from DW MRI, and this may help to distinguish low-grade lesions from high-grade ones (48). ADC is a reliable quantitative measure and ADC values can help to differentiate cancers from cystitis or wall hypertrophy secondary to outlet obstruction (45–47).

**Differential diagnosis**

Filling defects, which are among the most common imaging features of urothelial tumors, can also be seen in other pathologies that also result in diffuse enhancement, wall thickening, and focal infiltration.
Stone, clot, mucous, debris, fungus ball, concentrated gadolinium, flow-related artifacts, vascular partial volume, inflammation, reflux, peristaltic motion, and jet flow should be considered in the differential diagnosis of urothelial carcinoma (35–37,49–51). Additionally, nephrostomy tubes, instrumentations, urinary stasis, intravesical chemotherapy agents, neobladder and urinary diversions, and urinary stents can mimic diffuse contrast enhancement, as well (1,2).

**Lymph node metastasis**

Spherical lymph nodes greater than 8 mm, oval or ellipsoid nodes greater than 10 mm with rapid enhancement at dynamic scans, and clustered lymph nodes are important findings for metastatic involvement (1–4,52). A few publications have reported that ADC values can be helpful in differentiating benign nodes from malignant ones; however, further research is required to explore this topic (53).

**Surgical treatment of bladder tumors**

Treatment approach varies with patient age and comorbidities. The aim of the treatment in superficial (stage T1) lesions is to prevent recurrence or progression. A bladder-sparing treatment is applied in early-stage tumors, and partial cystectomy or transurethral tumor resections are among such techniques (54). In situ lesions tend to be high grade and they may require cystectomy if there is no response to 2 episodes of bacillus Calmette-Guérin treatment for 6 weeks. T2 and T3 lesions usually require cystectomy as they have infiltrative characteristics in the presence of nodal infiltration, and chemotherapy and radiotherapy should be added to the treatment regimen. Cystectomy is a well-known standard treatment for patients with invasive tumor (54,55). After pelvic surgery, bleeding is a common and significant risk. Techniques of interventional radiology are noninvasive and safe methods for demonstration of the bleeding area and effective treatment (56).

An artificial bladder is reconstructed in cystectomy and this is known as a diversion procedure. Several diversion techniques exist:

**Urinary diversions**

They are mainly grouped into 2 types.

Noncontinent diversions (ureterostomy and conduits): In ureterostomy, ureters are directly attached to the skin without an intestinal loop. This includes 2 stomas and a pouch. This technique is rarely used. In case of an ileal conduit, a reservoir is created from a short segment of the small bowel and the ureters are attached to one side of this loop, whereas the other side is attached to the skin to make up the stoma (Figure 10). An additional pouch is also created in the opening of the stoma. This technique is well established with long-term complication and collecting system dilatation rates of 30% and 20%, respectively (57).

**Conclusion**

Urothelial cancers usually present with painless hematuria and are currently more commonly diagnosed secondary to recent developments in the imaging technology. The application of an accurate imaging algorithm to further investigate painless hematuria will enable prompt and correct diagnosis of urothelial cancers, which will increase the yield of treatment and significantly decrease cancer and treatment-related morbidity and mortality. Moreover, the angiographic techniques can be useful in the pre- and postoperative period management of patients with urothelial cancer.

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References


