Urinary Tumor Markers For Bladder Cancer

Policy Number: **APEA – G2125** – Urinary Tumor Markers For Bladder Cancer

Initial Presentation Date: 7/01/2020
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Policy Description

Bladder cancer is defined as a malignancy that develops from the tissues of the bladder. It is the most common cancer of the urinary system. The cancer typically arises from the urothelium, although it may originate in other locations such as the ureter or urethra (Lerner, 2018).

Tumor biomarkers are proteins detected in the blood, urine, or other body fluids that are produced by the tumor itself or in response to it. Urinary tumor markers may be used to help detect, diagnose, and manage some types of cancer including bladder cancer (Hottinger & Hormigo, 2011).

Related Policies

<table>
<thead>
<tr>
<th>Policy Number</th>
<th>Policy Title</th>
</tr>
</thead>
<tbody>
<tr>
<td>AHS-G2124</td>
<td>Serum Tumor Markers for Malignancies</td>
</tr>
<tr>
<td>APEA-G2054</td>
<td>Liquid Biopsy</td>
</tr>
<tr>
<td>AHS-M2109</td>
<td>Molecular Panel Testing of Cancers to Identify Targeted Therapy</td>
</tr>
</tbody>
</table>

Indications and/or Limitations of Coverage

Application of coverage criteria is dependent upon an individual’s benefit coverage at the time of the request.

1. Urinary biomarkers (bladder tumor antigen (BTA) test, nuclear matrix protein (NMP22) test, or fluorescence in situ hybridization (FISH) UroVysion Bladder Cancer test) **MEET COVERAGE CRITERIA**:

   a. An adjunct in the diagnostic exclusion of bladder cancer for patients who have an atypical or equivocal cytology
   
   b. As an adjunct in the monitoring of high-risk, non-muscle invasive bladder cancer

2. The use of fluorescence immunocytotherapy (ImmunoCyt/uCyt) **MEETS COVERAGE CRITERIA** as an adjunct to cystoscopy or cytology in the monitoring of persons with bladder cancer.
The following does not meet coverage criteria due to a lack of available published scientific literature confirming that the test(s) is/are required and beneficial for the diagnosis and treatment of a patient’s illness.

3. Urinary biomarkers (bladder tumor antigen (BTA) test, nuclear matrix protein (NMP22) test, or fluorescence in situ hybridization (FISH) UroVysion Bladder Cancer test) **DO NOT MEET COVERAGE CRITERIA** for screening of bladder cancer, evaluation of hematuria, diagnosing bladder cancer in symptomatic individuals, and all other indications.

4. The use of fluorescence immunocytology (ImmunoCyt/uCyt) **DOES NOT MEET COVERAGE CRITERIA** in the evaluation of hematuria, diagnosing bladder cancer, or for screening for bladder cancer in asymptomatic persons and all other indications.

5. Any other urinary tumor markers for bladder cancer not mentioned above **DO NOT MEET COVERAGE CRITERIA**.

**Scientific Background**

In 2015, there were approximately 541,000 cases of bladder cancer and 188,000 deaths related to bladder cancer worldwide (Fitzmaurice et al., 2017). In 2019 in the United States, there were more than 80,000 cases and 17,000 deaths due to bladder cancer (NCCN, 2019; Siegel, Miller, & Jemal, 2017). Bladder cancer is the sixth most common cancer in the United States, affects men four times more frequently than women, and is typically diagnosed in individuals above the age of 40, with 73 the median age at diagnosis (DeGeorge, Holt, & Hodges, 2017; NCCN, 2019). Bladder cancer risk factors include smoking, a family history of the disease, pelvic radiation, obesity, diabetes, and chronic infection of the urinary tract.

Bladder cancer commonly presents as painless hematuria (blood in urine) and may be gross (visible) or microscopic. Gross hematuria tends to increase the likelihood of bladder cancer, but hematuria as a whole may be transient or due to non-cancer related causes (Kurtz, 2018). Other common symptoms of bladder cancer include pain or irritative and obstructive voiding symptoms such as urge incontinence, dysuria, straining, or nocturia. These symptoms are often mistaken for another condition such as kidney stones, can be temporary, and are not necessarily specific for bladder cancer (Y. Lotan, Choueiri, Toni, 2017). In fact, hematuria is the most common symptom of bladder cancer, but a study reported a 13% prevalence rate of bladder cancer out of 6728 patients with hematuria (DeGeorge et al., 2017; Sutton et al., 2018). Approximately 70%-75% of patients present with superficial tumors (50 – 70% of which can recur but are usually not life threatening), and 25%-30% present as invasive tumors with a high risk of metastasis (Chou & Dana, 2010; Kaufman, Shipley, & Feldman, 2009).

Cystoscopy (white light) is the gold standard for a diagnosis of bladder cancer. This procedure involves a bladder examination and urine sample for cytology. Any lesions are observed and recorded. Cystoscopy does not detect all malignancies or visualize the upper urinary tract. Furthermore, although cystoscopy is minimally invasive, it may be uncomfortable and promote anxiety, which can lead to suboptimal compliance with management recommendations. Fluorescent cystoscopy is somewhat more efficient at detecting tumors than white light cystoscopy; although, it comes with its own set of issues such as higher false-positive rates and costs (Y. Lotan, Choueiri, Toni, 2017; Mitra, Birkman, & Penson, 2017). Cytology, or the analysis of cells in urine, is often completed in addition to cystoscopy analysis.

Although cystoscopy has long been the gold standard for a diagnosis of bladder cancer, its high cost and unpleasant burden has led to the search for a non-invasive test that can match the high specificities and sensitivities set by cystoscopy. Urinary biomarkers including “Cell-free proteins and peptides, exosomes, cell-free DNA, methylated DNA and DNA mutations, circulating tumor cells, miRNA, IncRNA, rtRNA and mRNAs” have now been identified for bladder cancer diagnostic purposes (Lopez-Beltran et al., 2019). Urine is exposed to urothelial tissue in
many different locations, and therefore has the potential to contain several biomarkers associated with cancer. Validation of these biomarkers could lessen the use of cystoscopy as well as increase the overall sensitivity for bladder cancer identification (D'Costa, Goldsmith, Wilson, Bryan, & Ward, 2016). However, because of the lower disease prevalence in a screening population, even in those at increased risk, the use of biomarkers for screening is not cost effective or recommended (Y. Lotan et al., 2009). Despite the promise of urine biomarkers, cystoscopy remains the procedure of choice both for initial diagnosis and for surveillance in previously treated patients.

The two most studied urinary biomarkers are bladder tumor antigen (BTA) and nuclear matrix protein 22 (NMP22). The BTA test is designed to detect complement factor H-related protein (hCFHrp) which is elevated in cancer cells. This test is available in both a quantitative and qualitative version, and its manufacturer-recommended cut-off is 14U/mL (Mahnert et al., 1999; Mitra et al., 2017). Similarly, the NMP22 test is designed to detect a protein that is more highly available in cancer cells than normal cells. In this case, cancer cells release more NMP22 into the urine following apoptosis than normal cells do. The NMP22 tests are also available in a quantitative and qualitative version, and its FDA-approved cut-off is 10U/mL (Grossman, Messing, Soloway, & et al., 2005; Mitra et al., 2017; Zuiverloon, de Jong, & Theodorescu, 2017).

The FDA has approved two additional tests for urinary biomarkers. One is UroVysion, which is designed to detect chromosomal alterations that are distinctive of bladder cancer. This test is a fluorescent in situ hybridization (FISH) assay that uses DNA probes to detect alterations (such as aneuploidies) on chromosomes 3, 7, and 17 or loss of the 9p21 locus. The second test is known as ImmunoCyt (or uCyt+) that uses a similar fluorescent technique to detect certain glycoproteins that are expressed solely on cancerous cells (Mitra et al., 2017).

Epigenetic changes may also play an important role in bladder cancer tumorigenesis. These changes are becoming more prevalent as identification rates increase due to improvements in high-throughput DNA sequencing technologies. Epigenetic changes can “regulate [the] gene expression outcome without changing the underlying DNA sequence” with alterations based on DNA methylation, nucleosome positioning, microRNA regulation and histone medications (Li, Duymich, Weisenberger, & Liang, 2016). All of these epigenetic-based changes are distorted in each human cancer type. “A substantial portion (76%) of all primary bladder tumors displays mutations in at least one chromatin regulatory gene. These mutations cause epigenetic dysregulation in bladder cancers (Li et al., 2016).”

Recently, Pangea Laboratory has created a laboratory developed test termed Bladder CARE™ which measures the methylation status of specific DNA biomarkers in urine for the detection of bladder cancer via an at-home collection kit. This non-invasive test has not been approved by the FDA, is purported to be more cost-effective, and uses an epigenetic-based detection approach. Specifically, the methylation of bladder cancer DNA biomarkers are measured (Pangea, 2019a). As little as 5 ng of urine DNA from a 100 mL urine sample is required, and it has a limit detection of 0.1% leading to the identification of a single cancerous cell in a sample of 1,000 normal cells (Pangea, 2019a). The authors claim that Bladder CARE™ has a sensitivity of 94% and specificity of 86%, allowing for the identification of 88% of low-grade bladder cancer cases; these results are based on a study completed by Pangea Laboratory and Zymo Research which analyzes urine samples from 182 patients (97 with bladder cancer and 85 healthy controls) (Pangea, 2019b).

Another test, termed the Bladder EpiCheck test, has been developed by the Israeli company Nucleix. This non-invasive epigenetic urine test helps to detect bladder cancer with a panel of 15 DNA methylation biomarkers. Nucleix reports a sensitivity of 92%, a specificity of 88% and a negative predictive value of 99% for the Bladder EpiCheck test; these results are based on a multi-center clinical study with 353 bladder cancer patients (Nucleix, 2015). Similar results have been reported by D'Andrea et al. (2019).
Recently, Piao et al. (2019) have developed a way to differentiate patients with bladder cancer from patients with a nonmalignant hematuria without bladder cancer by measuring urinary cell-free microRNA expression. This study shows that the non-invasive measurement of urinary microRNA-6124 and microRNA-4511 can be used as a diagnostic tool with a sensitivity of >90% (Piao et al., 2019). This testing method will help to reduce the number of unnecessary cystoscopies in patients with hematuria that are being evaluated for bladder cancer.

The performance of an epigenetic-based bladder cancer detection tool has been evaluated by Fantony et al. (2017); the urine-based TWIST1/NID2 methylation assay has been analyzed for the detection of urothelial carcinoma via the addition of urine cytology. This multi-institutional study analyzed data from 172 patients. The authors note that “The AUC [area under the curve] for cytology alone with equivocal cytologies positive was 0.704, and improved to 0.773 with the addition of the DNA methylation assay (p < 0.001) (Fantony et al., 2017).” The authors conclude by stating that this TWIST1/NID2 methylation assay is a sensitive diagnostic tool that adds value to urine cytology for the detection of urothelial carcinoma, which is the most common type of bladder cancer.

Soubra and Risk (2015) found the sensitivity of fluorescent cystoscopy to be 0.92 and the sensitivity of white light cystoscopy to be 0.71; the specificity of fluorescent cystoscopy was lower at 0.57, and the specificity of white light cystoscopy was identified at 0.72. Furthermore, fluorescent cystoscopy’s sensitivity for carcinoma in situ (which is difficult to visualize) was measured at 0.924, while white light cystoscopy’s sensitivity for carcinoma in situ was much lower at 0.605, but these differences tended to decrease on higher grade lesions (Soubra & Risk, 2015). Cytology is also a common analytic technique in addition to cystoscopy. Its overall sensitivity is low at 0.34 and its sensitivity for grade 1 and 2 tumors is even lower at 0.12 and 0.26, respectively (Yair Lotan & Roehrborn, 2003).

Breen et al. (2015) compared the sensitivity and specificity values of four diagnostic tests (cytology, NMP22, UroVysion, and CxBladder); CxBladder was found to have the highest sensitivity at 74% and cytology was identified with the highest specificity at 95%. The authors report comparable sensitivity values for cytology, NMP22, and UroVysion at 46%, 45.9% and 47.7% respectively (Breen et al., 2015). It is important to note that even though CxBladder is reported to have the highest sensitivity, the specificity (81.7%) is the lowest; the other tests were reported to have superior specificities with NMP22 at 88%, and UroVysion at 87.7% (Breen et al., 2015).

Sathianathen, Butaney, Weight, Kumar, and Konety (2018) published a study focusing on biomarkers in patients presenting with hematuria. This study encompassed BTA, NMP22, FISH, and uCyt+, as well as a fifth biomarker known as AssureMDx. Sensitivities ranged from 0.67 (BTA) to 0.95 (AssureMDx, second highest was uCyt+ at 0.83) while specificities ranged from 0.68 (BTA) to 0.93 (quantitative NMP22). However, this data is consistent with the previously published meta-analysis that covered all settings, not just hematuria (Chou et al., 2015). Cytology was also found to have superior specificity to all studied biomarkers; although, biomarkers tended to have better sensitivity. The authors concluded that, due to the high heterogeneity and small sample size, more studies were needed to validate biomarkers to replace diagnostic evaluation of hematuria (Sathianathen et al., 2018).

Although many studies emphasize the high validity of biomarkers such as NMP22 and BTA, these studies often have a large proportion of high-grade tumors which inflate the specificity and sensitivity; hence, the problem of identifying low-grade cancers remains. There may be changes at the genetic level in a low-grade cancer, but the proteins tested in the urine may still be relatively normal (D’Costa et al., 2016). Another issue is the conflicting results for the validity of the biomarkers. For example, the sensitivity of the quantitative NMP22 test has been found to range from as low as 0.26 to 1.00 with its specificity ranging from 0.49 to 0.98. Similarly, the BTA STAT test’s sensitivity and specificity have been found to range from 0.29 to
0.91 and from 0.54 to 0.86 respectively (Zuiverloon et al., 2017). For comparison, a study found the sensitivity and specificity of flexible cystoscopy (out of 778 hematuria patients) to be 0.98 and 0.938, respectively (Sutton et al., 2018).

Dudley et al. (2019) have developed a novel high-throughput sequencing method that uses urine derived tumor DNA (utDNA) known as utDNA CAPP-Seq (uCAPP-Seq) to detect bladder cancer. This technique was used to analyze samples from 118 patients with early stage bladder cancer and 67 healthy adults. “We detected utDNA pretreatment in 93% of cases using a tumor mutation-informed approach and in 84% when blinded to tumor mutation status, with 96% to 100% specificity (Dudley et al., 2019).” These results show that utDNA can be used to diagnose early-stage bladder cancer with high sensitivity and specificity.

Clinical Validity and Utility

A meta-analysis of 57 studies detailed the accuracy of several biomarkers for the diagnosis and surveillance of bladder cancer. These included the six FDA-approved tests (quantitative and qualitative NMP22, quantitative and qualitative BTA, FISH, and uCyt+) as well as a laboratory developed test that does not require FDA approval termed CxBladder. Sensitivities ranged from 0.57 (qualitative NMP22) to 0.82 (CxBladder); however, the CxBladder cohort was only comprised of one study. The specificities ranged from 0.74 (quantitative BTA) to 0.88 (qualitative NMP22). Sensitivity increased as a tumor progressed (higher grade or stage) with low accuracy for lower stage or grade tumors. A cytologic evaluation performed with a biomarker assessment increased sensitivity as well but missed about 10% of cases. Ultimately, the authors concluded that urinary biomarkers reported many false-positive results and failed to identify a large percentage of patients with bladder cancer (Choue et al., 2015). The authors also noted that this was the first study which focused on the measurement of clinical outcomes based on urinary biomarkers.

The ideal marker will be “easier, better, faster, and cheaper” (Schmitz-Dräger et al., 2015). Overall, although there have been numerous promising studies for the clinical utility of these urinary biomarkers, the biomarkers do not yet measure up to the standards set by cystoscopy as the primary method of diagnosis. Most of the biomarkers are yet to be well-validated and the ones that are, such as NMP22 and BTA, fall short of cystoscopy’s standards (D’Costa et al., 2016). Furthermore, because of the lower disease prevalence in a screening population, even in those at increased risk, the use of biomarkers for screening is not cost effective or recommended (Y. Lotan et al., 2009). Although the cost of tests is non-clinical, it is still a crucial issue; the BTA and NMP22 tests are relatively inexpensive at $25 but ImmunoCyt costs around $80 and the CxBladder and UroVysion cost $325 and $800, respectively (Zuiverloon et al., 2017). For comparison, a cystoscopy cost around $210 in 2016, and a cystoscopy with a biopsy cost about $370 (Halpern, Chughtai, & Ghomrawi, 2017). These biomarkers to date have not been highly recommended within any clinical guidelines. Therefore, the authors concluded that biomarkers have not had significant effect on clinical decision-making (Schmitz-Dräger et al., 2015). The majority of studies performed on these biomarkers did not focus on their ability to predict the course of cancer (D’Costa et al., 2016) but some biomarkers may play a role in the diagnosis or surveillance of bladder cancer in the future (Schmitz-Dräger et al., 2015). Even this may be a difficult barrier to cross; Meleth et al. (2014) prepared an assessment for the Agency for Healthcare Research and Quality that stated “although UroVysion is marketed as a diagnostic rather than a prognostic test, limited evidence from two small studies (total n=168) supported associations between test result and prognosis for risk of recurrence (Meleth et al., 2014).” The authors went on to note that no studies that established clinical utility were found.

D’Andrea et al. (2019) analyzed 357 urine samples from patients at five different centers under surveillance for non-muscle-invasive bladder cancer to investigate the clinical utility of the Bladder EpiCheck™ non-invasive urine test. A specificity of 88% was identified with this test, a negative predictive value of 94.4% for the detection of any cancer, and a negative predictive value of 99.3% for the detection of high grade cancer; the use of the Bladder EpiCheck™ test
helped to improve the cancer recurrence predictive value by a difference of 16-22% (D'Andrea et al., 2019). This high-performing diagnostic test may help in the surveillance of non-muscle-invasive bladder cancer.

Tan et al. (2018) completed a systematic review to identify the diagnostic sensitivity and specificity of urinary biomarkers for the diagnosis of bladder cancer. The authors report that multi-target biomarker panels were more accurate than single biomarker targets, and that both the sensitivity and specificity of urinary biomarkers were higher in primary diagnostic scenarios compared to patients under surveillance (Tan et al., 2018). The authors note that “few biomarkers achieve a high sensitivity and negative predictive value,” with single biomarkers reporting a sensitivity of 2-94% and specificity of 46-100%, and multi-target biomarkers reporting a sensitivity of 24-100% and specificity of 48-100% (Tan et al., 2018).

Guidelines and Recommendations

National Comprehensive Cancer Network (NCCN) (NCCN, 2018, 2019, 2020)

The NCCN has stated that “Urine molecular tests for urothelial tumor markers are now available. Many of these tests have a better sensitivity for detecting bladder cancer than urinary cytology, but specificity is lower. Considering this, evaluation of urinary urothelial tumor markers may be considered during surveillance of high-risk non-muscle-invasive bladder cancer. However, it remains unclear whether these tests offer additional information that is useful for detection and management of non-muscle-invasive bladder tumors. Therefore, the panel considers this to be a category 2B recommendation (NCCN, 2018, 2019, 2020).”

The NCCN previously stated that an FDA-approved urinary biomarker test such as fluorescence in situ hybridization (FISH) or nuclear matrix protein 22 may be considered in monitoring for recurrence (NCCN, 2018). However, updated NCCN (2019, 2020) guidelines no longer address these biomarker tests.

National Academy of Clinical Biochemistry Laboratory Medicine (NACB) (NACB, 2010)

The NACB Laboratory Medicine Practice Guidelines do not recommend use of any FDA‒approved urinary tumor marker tests for the diagnosis of bladder tumors or for monitoring bladder cancer patients. The guideline states that “There are no prospective clinical trial data that establish the utility of any of the FDA cleared markers or the proposed markers for increasing survival time, decreasing the cost of treatment or improving the quality of life of bladder cancer patients (NACB, 2010).” The NACB is now known as the AACC, or American Association for Clinical Chemistry, and have not since released any further updates on this topic.

American Urological Association (AUA) (Chang et al., 2017; Chang et al., 2016; Davis et al., 2012)

The AUA’s guidelines on the diagnosis, evaluation and follow-up of asymptomatic microhematuria (AMH) in adults do not recommend use of urine markers (NMP22, BTA-stat, UroVysion) as part of routine evaluation (Davis et al., 2012).

The AUA and Society of Urologic Oncology (SUO) joint guidelines on Diagnosis and Treatment of Non-Muscle Invasive Bladder Cancer (NMIBC) do not recommend using urinary biomarkers to replace cystoscopy when monitoring NMIBC (grade B), although a clinician can use biomarkers to evaluate a patient’s response to Bacillus Calmette-Guerin (BCG) therapy or a separate cytology such as FISH. However, a urinary biomarker should not be used for monitoring a patient with a normal cystoscopy or a history of low-risk cancer (Chang et al., 2016).

Similarly, the joint guidelines between the AUA, the SUO, the American Society of Clinical Oncology (ASCO), and the American Society for Radiation Oncology (ASTRO) regarding non-
metastatic muscle-invasive bladder cancer note that molecular biomarkers may be important for staging cancer and deciding a course of treatment soon. Nevertheless, at this time the biomarkers have not been properly validated (Chang et al., 2017).

**U.S. Preventive Services Task Force (USPSTF) (Moyer, 2011; USPSTF, 2019)**

The USPSTF concluded in 2011 that there was insufficient evidence to evaluate screening for bladder cancer in asymptomatic adults, assigning a grade I to this recommendation. Since then, there have been no further guidelines published on this topic by the USPSTF (Moyer, 2011).

In 2019, the USPSTF published the following statement regarding bladder cancer screening in adults: "Literature scans conducted in April 2019 in the MEDLINE and PubMed databases and the Cochrane Library showed a lack of new evidence to support an updated systematic review on the topic at this time (USPSTF, 2019)."

**3rd International Consultation on Urological Diseases & Société Internationale d’Urologie (ICUD-SIU) (Monteiro et al., 2018)**

With a level of evidence of 3 and a grade of “B”, the ICUD-SIU recommends, "examination of urine cytology must be a part of the expectant management or active surveillance protocol." Concerning the surveillance strategies for NMIBC, "Surveillance strategies following a negative 3 months surveillance cystoscopy should be: (1) for low risk disease, cystoscopy 6–9 months later and annually thereafter; consider cessation following five recurrence-free years. No upper tract imaging necessary unless hematuria present; (2) for intermediate risk, cystoscopy with cytology every 3–6 months for 2 years; then every 6–12 months during years 3 and 4; then annually for lifetime. Upper tract imaging every 1–2 years; (3) for high risk, cystoscopy with cytology every 3 months for 2 years; then every 6 months during years 3 and 4; then annually for lifetime [Level of evidence: 3; Grade C] (Monteiro et al., 2018)."

**National Cancer Institute (NCI) (NCI, 2018)**

In the 2018 update to the NCI’s *Bladder and Other Urothelial Cancers Screening (PDQ®)—Health Professional Version*, the NCI states that “There is inadequate evidence to determine whether screening for bladder and other urothelial cancers has an impact on mortality... Based on fair evidence, screening for bladder and other urothelial cancers would result in unnecessary diagnostic procedures with attendant morbidity (NCI, 2018).”

**European Association of Urology (EAU) (Babjuk et al., 2017)**

The EAU has published guidelines on non-muscle-invasive bladder cancer. Regarding urinary molecular marker tests, the EAU has stated that “Driven by the low sensitivity and low negative predictive value of urine cytology, numerous urinary tests have been developed. None of these markers have been accepted for diagnosis or follow-up in routine practice or clinical guidelines (Babjuk et al., 2017).” Further, as an exploratory measure after hematuria or after other bladder cancer symptoms have been identified, the EAU states that “It is generally accepted that none of the currently available tests can replace cystoscopy. However, urinary cytology or biomarkers can be used as an adjunct to cystoscopy to detect missed tumours, particularly CIS [carcinoma in situ]. In this setting, sensitivity for high-grade tumours and specificity are particularly important (Babjuk et al., 2017).” Finally, the EAU states that currently, there is no urinary marker with the ability to replace cystoscopy.

**State and Federal Regulations, as applicable**

On April 16, 1997, the FDA approved the *Bard BTA stat™ Test*, created by Bard Diagnostic Sciences Inc. From the FDA site: “the BTA stat test is an *in vitro* diagnostic immunoassay indicated for the qualitative detection of bladder tumor associated antigen in urine of persons
diagnosed with bladder cancer. This test is indicated for use as an aid in the management of bladder cancer patients in conjunction with cystoscopy.”

On April 15, 1998, the FDA approved the *BTA TRAK™ Test*, created by Bard Diagnostic Sciences Inc. From the FDA site: “the BTA TRAK test is an *in vitro* diagnostic immunoassay indicated for the quantitative detection of bladder tumor associated antigen in human urine. This test is indicated for use as an aid in the management of bladder cancer patients in conjunction with cystoscopy.”

On July 2, 1996, the FDA approved the *MATRITECH NMP22™ TEST KIT*, created by Alere Scarborough Inc. From the FDA site: “The Matritech NMP22 Test Kit is an enzyme immunoassay (EIA) for the *in vitro* quantitative determination of nuclear matrix protein NMP22 in stabilized voided urine.”

On July 30, 2002, the FDA approved the *NMP22 BladderChek*, created by Matritech Inc. From the FDA site: “The Matritech NMP22 BladderChek Test is indicated for professional and prescription home use as an aid in monitoring bladder cancer patients, in conjunction with standard diagnostic procedures.” This assay is qualitative.

On January 24, 2005, the FDA approved the *UROVYSION BLADDER CANCER KIT*. From the FDA site: “The UroVysion Bladder Cancer Kit (UroVysion Kit) is designed to detect aneuploidy for chromosomes 3, 7, 17, and loss of the 9p21 locus via fluorescence in situ hybridization (FISH) in urine specimens from persons with hematuria suspected of having bladder cancer.”

On February 23, 2000, the FDA approved the *ImmunoCyt*, created by Diagnocure Inc. From the FDA site: “ImmunoCyt is a qualitative direct immunofluorescence assay intended for use in conjunction with cytology to increase overall sensitivity for the detection of tumor cells exfoliated in the urine of patients previously diagnosed with bladder cancer. ImmunoCyt is indicated for use as an aid in the management of bladder cancer in conjunction with urinary cytology and cystoscopy (FDA, 2018).”

All of the FDA-approved tests apart from ImmunoCyt are approved for both diagnosis and surveillance of bladder cancer whereas ImmunoCyt is only approved for surveillance (Darwiche, Parekh, & Gonzalgo, 2015).

Additionally, many labs have developed specific tests that they must validate and perform in house. These laboratory-developed tests (LDTs) are regulated by the Centers for Medicare and Medicaid (CMS) as high-complexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA ’88). As an LDT, the U. S. Food and Drug Administration has not approved or cleared this test; however, FDA clearance or approval is not currently required for clinical use.

### Applicable CPT/HCPCS Procedure Codes

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<th>Code Number</th>
<th>Code Description</th>
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<td>86294</td>
<td>Immunoassay for tumor antigen, qualitative or semiquantitative (e.g., bladder tumor antigen)</td>
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<td>86316</td>
<td>Immunoassay for tumor antigen; other antigen, quantitative, each</td>
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<td>86386</td>
<td>Nuclear matrix protein 22 (nmp22), qualitative</td>
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<td>88120</td>
<td>Cytopathology, in situ hybridization (eg. FISH), urinary tract specimen with morphometric analysis, 3-5 molecular probes, each specimen, manual</td>
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<td>Code Number</td>
<td>Code Description</td>
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<td>88121</td>
<td>using computer-assisted technology (for morphometric in situ hybridization on cytologic specimens other than urinary tract, see 88367, 88368) (for more than 5 probes, use 88399)</td>
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<td>0013M</td>
<td>Oncology (urothelial), mRNA, gene expression profiling by real-time quantitative PCR of five genes (MDK, HOXA13, CDC2 [CDK1], IGFBP5, and CXCR2), utilizing urine, algorithm reported as a risk score for having recurrent urothelial carcinoma</td>
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</tbody>
</table>


Procedure codes appearing in Medical Policy documents are included only as a general reference tool for each policy. They may not be all-inclusive.

Evidence-based Scientific References


FDA. (2018). Devices@FDA.


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